Hitachi Real-time Tissue Elastography:

Publications & International Communications

Clinical Abstracts

HI-RTE
Hitachi Real-time Tissue Elastography
Hitachi Real-time Tissue Elastography for applications using Endoscopic Ultrasound
ACCURACY OF SECOND-GENERATION ENDOSCOPIC ULTRASOUND (EUS) ELASTOGRAPHY FOR THE DIFFERENTIAL DIAGNOSIS OF SOLID PANCREATIC MASSES: A QUANTITATIVE ANALYSIS OF TISSUE STIFFNESS
Julio Iglesias-Garcia, Jose Larino, Enrique Dominguez-Munoz

Background: EUS-elastography (EE) allows analyzing tissue stiffness during a standard endoscopic ultrasound examination. Contrarily to the subjective qualitative analysis associated to the first-generation elastography equipments, second-generation elastography allows a quantitative analysis of the tissue stiffness. Aim of the study was to evaluate the accuracy of second generation EUS-elastography for the differential diagnosis of solid pancreatic masses.

Methods: 57 consecutive patients (mean age 60 years, 32-84 years, 41 male), who underwent EUS for the evaluation of solid pancreatic masses were prospectively included in the study. EUS-elastography was performed under conscious sedation with the linear Pentax EUS (EG 3830 UT) and the Hitachi EUB 900. Two different areas (A and B) from the region of interest were selected for the quantitative elastographic analysis: Area A is a representative area of the mass and B refers to a soft reference area. The quotient B/A (strain ratio) is considered as the result of the elastographic evaluation. EUS-FNA was performed in all cases for cytological diagnosis. Data are shown as median and 95%CI, and compared by the Student-t test. Diagnostic accuracy of EUS-elastography for detecting malignancy was calculated after drawing the corresponding ROC curves.

Results: Size of solid pancreatic masses was 29.7±10.1 mm (mean±SD). Tumors were located in the head of the pancreas in 44 patients, in the body in 10 patients and in the tail in 3 patients. Final diagnosis was pancreatic cancer (PC) in 32 patients and inflammatory mass (IM) in 25 patients. Strain ratio was 16.62 (95%CI 7.26-30.64) in PC and 3.46 (95%CI 1.3-12.6) in IM (p<0.001). Sensitivity and specificity of EUS-elastography (strain ratio) for detecting pancreatic malignancy was 100% and 92%, respectively (cut-off point 5.85). The elastographic value of the mass (area A) was 0.02 (95%CI 0.01-0.05) in PC and 0.22 (95%CI 0.03-0.49) in IM (p<0.001). The area under the ROC curve for detecting malignancy by the analysis of the elastographic value of A (AUC=0.944) was slightly higher to that obtained by the analysis of the strain ratio (quotient B/A) (AUC=0.910).

Conclusion: Second-generation EUS-elastography is a very useful tool for the differential diagnosis of solid pancreatic masses. It allows a quantitative and objective evaluation of tissue stiffness, and thus provides additional information for the detection of malignancy.

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QUANTITATIVE ANALYSIS OF EUS-ELASTOGRAPHY IN PANCREATIC CANCER AND HEALTHY CONTROLS
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Introduction: Endoscopic ultrasound (EUS) elastography is a promising new technology for the visualization of tissue elasticity in EUS examinations. Recent data show a high sensitivity in detecting malignant pancreatic lesions, but there are limitations in the technical procedure and evaluation is still subjective. The aim of our study was to develop a quantitative analysis of EUS-elastography in patients with malignant pancreatic masses and controls.

Patients and methods: 86 patients (42m, 44w) with pancreatic cancer and 28 healthy controls (11m, 17w) were investigated by EUS-elastography. In all patients diagnosis of malignancy was confirmed by histology or cytology. Over 80% in the patient group suffered from ductal adenocarcinoma. Quantification of elastography data was performed by analysis of EUS videos evaluating histogram values for the colours blue, green and red in a 10s video. A region of interest (tumour or normal pancreas) was defined in each video, and the mean-values for the different colours were measured three times during the video. Data from patients and controls were compared by using an unpaired t-test.

Results: Figure 1 shows the video histogram analysis for the colours blue, green and red comparing the cancer and the control group. In pancreatic cancer the colour blue was dominant with a sensitivity of 100%. For normal pancreas the green colour was dominant with a similarly high sensitivity. The analysis of the red colour showed a significant difference for cancer and controls with a higher red colour in normal pancreas, but no absolute separation between the groups.

06-07-09
Conclusions: Quantitative analysis of EUS-Elastography allows a highly sensitive discrimination between pancreatic cancer tissue and normal pancreas, especially by evaluating the blue colour. However, since this quantitative analysis is based on the subjective definition of ROI's, investigator with sufficient experience are still required. Thus, quantitative elastography may provide a potent new tool to identify malignant lesions in patients with pancreatic masses.

Figure 1

Digestive Disease Week, May 30th – June 4th, 2009, Chicago, USA, M1430

REAL-TIME TISSUE ELASTOGRAPHY DETERMINES OPTIMAL TREATMENT AND PREDICTS THE PROGNOSIS OF ULCERATIVE COLITIS

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Background/Aims: Ulcerative colitis (UC) is a chronic inflammatory disease with frequent remissions and relapses. Although colonoscopic examination is important in determining treatment for patients with UC, it is an invasive procedure and can lead to complications. Non-invasive methods to facilitate evaluation of the large bowel have therefore been sought. Realtime tissue elastography is a new technique that visualizes the differences in tissue strain produced by freehand compression during routine ultrasonography. The aim of this study was to evaluate the usefulness of real-time tissue elastography in patients with UC.

Methods: Real-time tissue elastography was performed before colonoscopy in 41 patients with UC. Findings were classified into four types (normal, homogeneous, random, hard) on the basis of color arrangement. Endoscopy findings were classified into four types as follows A: normal mucosa, B: mucosal edema and erosion, C: punched-out ulcer, D: extensive ulcer. We then compared the relationship between real-time tissue elastography and colonoscopy findings, and also investigated whether elastography could reflect clinical stage, therapeutic response, and prognosis of UC patients.

Results: On elastography, 13 cases were classified as normal, 15 as homogeneous, 6 as random, and 7 as hard; while on endoscopy, 13 were classified as type A, 18 as type B, 8 as type C, and 2 as type D. We found significant associations between normal type and type A, homogeneous type and type C, random type and type C, and hard type and type D (p<0.001). Twelve patients (75%) with the normal type and six (66%) with the homogeneous type were in remission, while six (100%) with the random type and five (72%) with the hard type were in the active phase. All patients with the normal and homogeneous types responded to treatment with prednisolone and leukocytapheresis and were induced into remission. In contrast, no patients with the random or hard type responded to this treatment. Following the addition of tacrolimus, cyclosporine A and/or ganciclovir, five (38%) with the random type were induced into remission, whereas remission was achieved in only one (20%) with the hard type and three (50%) required colectomy. Of the 21 patients induced into remission and followed for more than a year, remission was maintained in 13 (76%) of 17 with the normal and homogeneous types and 1 (25%) of 4 with the random and hard types.

06-07-09
Conclusion: Findings of real-time tissue elastography reflected colonoscopic findings. We consider that real-time tissue elastography may be a useful tool in determining the optimal treatment for and predicting the prognosis of UC patients.

Digestive Disease Week, May 30th – June 4th, 2009, Chicago, USA, T1430

ANALYSIS OF EUS ELASTOGRAPHY TO DIFFERENTIATE MASS FORMING PANCREATITIS AND PANCREATIC CANCER BY USING STRAIN RATIO
Fumihide Itokawa, Takao Itoi, Atsushi Sofuni, Takayoshi Tsuchiya, Toshio Kurihara, Fuminori Moriyasu

INTRODUCTION: Recently, EUS elastography has been reported to supplemental information which can be applied for the diagnosis of pancreatic diseases. However, there is limit in evaluation only for color, and quantification by numerical value is required.

AIM: The aim of our study was to evaluate the ability of EUS elastography and quantification by using strain ratio(non mass area/mass area:SR) in order to distinguish mass forming pancreatitis(MFP) from pancreatic cancer(PC).

PATIENTS AND METHODS: The subjects were 105 patients performed an endoscopic ultrasound(EUS) for pancreas in our hospital till September 2006 to November 2008. The disease were 6 with mass forming pancreatitis(MFP), 5 with chronic pancreatitis(CP), 61 with pancreatic cancer (PC), 6 with neuroendocrine carcinoma (PNET), 4 with auto immune pancreatitis (AIP), 5 with SCN, 2 with SPN, 1 with Schwanoma, 1 with GIST, 1 with renal cell carcinoma pancreatic metastasis, 7 with IPMN, 1 with malignant lymphoma and 5 with normal control. A histological diagnosis by surgery or endoscopic ultrasonography fine needle aspiration (EUS-FNA) was performed except normal control. The ultrasound was used the HITACHI HI VISION900, and EUS scope was PENTAX EG-3630UR, EG-3670URK and EG-3870UTK. Strain ratio was subsequently performed to choose a mass area and a non-mass area, and the ratio was measured by calculating in real time.

RESULTS: Elastography for all PC showed intense blue coloration, which indicated that the mass lesions had malignant aspects. While MFP presented the coloration pattern of mixed green, yellow and low intensity of blue. Normal control was an even application of green to red. The mean SR of MFP and PC were each 23.08±12.65 and 39.08±20.54, respectively, which was significant difference (p<0.05)

CONCLUSION: EUS elastography is potentially capable of further defining the tissue characteristics of benign and malignant lesions. This study suggested that it was useful for the quantification by using strain ratio to characterize the tissue hardness of pancreatic disease and distinguish MFP from PC.

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USEFULNESS OF ENDOSCOPIC ULTRASOUND (EUS) ELASTOGRAPHY FOR THE DETECTION OF MALIGNANT INFILTRATION OF MEDIASTINAL AND ABDOMINAL LYMPH NODES
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Background: EUS-elastography allows analyzing tissue stiffness during a standard EUS examination, which may be of help in the differential diagnosis of solid lesions. Detection of malignant infiltration of LN is highly relevant to define the optimal therapeutic strategy for different tumors. We hypothesized that EUS-elastography may provide with additional information to the conventional EUS B-mode for the detection of malignancy in mediastinal and abdominal lymph nodes (LN). The aim of the study was to evaluate the usefulness of EUS-elastography in this setting.

Methods: 57 consecutive patients (mean age 64 years, 19-82, 43 males) who underwent EUS for the evaluation of LN were prospectively included in the study. EUS and elastography were performed under conscious sedation by using the linear Pentax EUS (EG 3830 UT) together with the Hitachi EUB 8500 and 900. EUS-guided fine needle aspiration was performed in all cases. Histology of
surgical specimens was considered as the reference method in operated cases. Positive cytology for malignancy together with compatible EUS, PET and CT imaging were considered as the reference method to define malignancy in non-operated cases. EUS, PET, CT imaging, clinical presentation and a minimum follow-up of six months were required for final diagnosis of benign disease in cases of benign cytology. Elastographic pattern of the different LN are described. Probability of malignancy according to the EUS-elastographic pattern was calculated.

**Results:** A total of 63 LN were evaluated. Size of LN was 17.4mm as a mean (range 4-59mm). 54 LN were located in the mediastinum, and 9 in the abdomen. Malignancy was confirmed by reference methods in 31 cases, whereas the remaining 32 LN were finally considered as benign. Three different elastographic patterns were identified: 1) a heterogeneous blue-predominant pattern (n=26 LN), 2) a heterogeneous green-predominant pattern (n=23 LN), and 3) a heterogeneous mixed green-blue pattern with geographical appearance and no color predominance (n=14 LN). Malignant LN showed either a blue-predominant pattern (n=24 LN) or a mixed green-blue pattern (n=7 LN). On the contrary, most benign LN (72%) showed a green-predominant pattern. The probability of malignancy in a LN showing a green-predominant pattern was of 0%, and of 92% in case of a blue-predominant pattern. Finally, the probability of malignancy in a LN showing a mixed green-blue pattern was of 50%.

**Conclusions:** EUS elastography is a very useful tool for the differential diagnosis of mediastinal and abdominal LN. It provides with specific colour patterns supporting the malignant or benign nature of the LN.

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**TRANSRECTAL ENDOSCOPIC ULTRASOUND (TRUS) ELASTOGRAPHY IN INFLAMMATORY BOWEL DISEASE (IBD)**

Nadan Rustemovic, Silvija Cukovic-Cavka, Davor Radic, Milorad Opacic, Zeljko Krznaric, Irena Hrstic, Roland Pulanic, Boris Vucelic

**BACKGROUND AND AIMS:** Establishing the diagnosis of Crohn's disease (MC) or ulcerative colitis (UC) sometimes is very difficult. When IBD is confined to the colon, there is a lack of diagnostic tools for distinction between Crohn's colitis and ulcerative colitis, which is especially important in the definitive phenotyping before surgical decision. The aim of this study was to assess the potential role of the TRUS elastography in distinction between MC and UC. The idea is based upon the fact that MC is transmural disease, and UC is limited to the mucosa and submucosa. These tissue characteristics are reflected in differences of the elasticity in rectal and perirectal tissue. Changes in the tissue elasticity can be obtained qualitatively by elastography with different colours (from red-soft tissue to blue-hard tissue) or quantitatively using strain ratio score.

**METHODS AND RESULTS:** Rectal wall thickness and elastomode of patients were measured by TRUS elastography. Endoscopist was blind for patient diagnosis. SPSS ver. 17 was used for statistical analysis. In pilot study we included 31 patients; 16 patients (52%) with MC and 15 patients (48%) with UC. Average thickness of rectal wall in all study patients was 6.43 mm (± 0.47 SE). In MC group mean rectal thickness was 7.28 mm (± 0.76 SE) compared to 5.52 mm (± 0.44 SE) in UC group. There was no statistical significant difference between MC and UC groups in perirectal thickness (t=1.97, df=29, p=0.058). Perirectal elastomode showed statistically significant difference between this two groups (.2=.18.6, df=2, P<0.001). Twelve (75%) patients in MC group had hard elastomode compared to none of patients in UC group, meaning that hard elastogram had positive predictive value of 100% for patients with MC. We also evaluated strain ratio (SR) of rectal tissue. Strain ratio is ratio of strain between two regions of interest (ROI) in the same image. Mucosal tissue was used as first ROI and perirectal tissue as second. SR was measured 3 times and middle value was used in statistical analysis. Sixteen patients were included in our study; 7 (44%) with MC and 9 (56%) with UC. Mean value of SR was higher in MC group (1.07 ± 0.26) then in UC group (0.26 ±0.21). This difference was statistically significant (t=6.85, df=14, p<0.001).

**CONCLUSION:** TRUS elastography provides a valuable information regarding the stiffness of the rectal and perirectal tissue, and can help to differentiate MC from UC. This is a promising new diagnostic tool in the field of IBD. Our study is ongoing and we expect improvement of the method, with increasing number of patients.
ENDOSCOPIC ULTRASONOGRAPHY (EUS) STRAIN RATIO (SR-EUS) VS. CONTRAST-ENHANCED EUS (CE-EUS) FOR THE DIAGNOSIS OF FOCAL PANCREATIC SOLID LESIONS
Fatima A. Figueiredo, Marc Giovannini, Erwan Bories, Christian Pesenti, Fabrice Caillol, Genevieve Monges, Jean Robert Delpero

BACKGROUND: There are many difficulties in the differential diagnosis between pancreatic cancer and chronic pancreatitis despite recent progress. As a means of trying to overcome this in the endosonography field, the measurement of the tissue stiffness (elastography) by new image processors and the use of contrast agent to enhance the color Doppler pattern have been introduced.
AIMS: Our aim is to compare prospectively the ability of the SR-EUS and CE-EUS to differentiate between benign and malignant focal pancreatic lesions.

METHODS: Thirty-eight patients with a focal pancreatic lesion were included to date. EUS procedures were performed with linear-array echoendoscopes (FG36X or EG38UT, Pentax), an ultrasound platform (Hitachi 7500 or 8500) with an integrated elastography module. In order to obtain the SR-EUS, a circular area is adjusted to the focal lesion and a second one is adjusted to the surrounding tissue. The SR-EUS (focal lesion area strain % / surrounding tissue area strain %) is calculated. After elastography, SonoVue® 2.4 mL (Bracco) was injected intravenously at a rate of 1 ml/sec, following a flash of 10mL saline solution. The pattern of enhancement (hypo or hypervascular) was defined using power Doppler mode. EUS-FNA was performed by using a 22-gauge FNA needle (Echotip, Cook Endoscopy,). The final diagnosis was based on the histological assessment of the EUS-FNA samples and/or surgical specimens when available. A positive cytological diagnosis was taken as a final proof of malignancy. For negative cytological specimens, the diagnosis was confirmed by surgery or follow-up of at least six months.

RESULTS: The study population comprised 21 (55%) men, mean age 62±16 years, with 12 (32%) benign and 26 (68%) malignant pancreatic lesions. Univariate analysis determined that the variables associated with malignancy included lesion size, SR-EUS and CE-EUS (Table). Logistic regression analysis determined that hypovascular lesions at CE-EUS (odds ratio 6.2 [95% CI, 1.2-32.1], p=0.03) was the only variable independently predictive of malignant pancreatic lesion. CONCLUSION: In this small group, CE-EUS was superior to SR-EUS for the differentiation between benign and malignant focal pancreatic lesions.

Univariate analysis

<table>
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<th>Characteristics</th>
<th>Benign lesions (n=12)</th>
<th>Malign lesions (n=26)</th>
<th>p</th>
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</thead>
<tbody>
<tr>
<td>Age (yr) mean±SD</td>
<td>58±6</td>
<td>65±14</td>
<td>0.06</td>
</tr>
<tr>
<td>Sex Male n(%)</td>
<td>6 (50%)</td>
<td>14 (54%)</td>
<td>0.60</td>
</tr>
<tr>
<td>Size mm mean±SD</td>
<td>23±14</td>
<td>35±12</td>
<td>0.03</td>
</tr>
<tr>
<td>SR-EUS (yr) mean±SD</td>
<td>15±21</td>
<td>34±47</td>
<td>0.05</td>
</tr>
<tr>
<td>CE-EUS hypovascular pattern</td>
<td>3 (25%)</td>
<td>20 (77%)</td>
<td>0.003</td>
</tr>
</tbody>
</table>

SR = strain ratio CE = contrast enhanced

DIGESTIVE DISEASE WEEK, May 30th – June 4th, 2009, Chicago, USA, W1258
(Background) Real-time tissue elastography™(EG) (Hitachi, Japan) provides a new tissue characterization of degree of hardness in real time. We have performed this procedure over 600 cases for GI diseases using EUS with the function of EG (EUS-EG), and here we will report the diagnostic capability of the involvement of hepatoduodenal ligament (HDL) and lymphnodes (LN) that substantially influence the therapeutic strategy of biliary malignancies.

(Patients and Methods) Seventy-five patients with biliary malignancies were enrolled in this study (40 with bile duct cancer and 35 with gallbladder cancer). There were 55 cases with enlarged LN (more than 5mm in diameter) and 20 cases who were estimated whether to have HDL involvement using EUS-EG. As a preliminary study, normal HDL images on EUS-EG were estimated for 10 patients. Diagnostic criteria of the HDL involvement and LN involvement were as follows; Involved HDL images were defined as inhomogeneous hardness based on the normal HDL images which were determined as homogeneous hardness in a preliminary study. Involved LN images were defined as the harder mass than the surrounding connective tissue or as inhomogeneous hardness mass. In this system, both B-mode and EUS-EG images are represented on dual screen at the same time, we can compare both images of same scan plane precisely. EUB-8500 and HV-900 (Hitachi, Japan) as an ultrasound diagnostic machine and EG-3630UR, EG-3670URK and EG-3780UTK (Pentax, Japan) as endosonoscopes were used in this study.

(Results) The sensitivity and specificity diagnosing HDL involvement were 89%, 75%, respectively. The extension of inflammation was erroneously depicted as cancer involvement in some cases. The sensitivity and specificity diagnosing LN involvement were 96%, 89%, respectively. The granulomatous change in the enlarged LN was misdiagnosed in some cases.

(Conclusion) EUS-EG may add the new information and be a useful method in the diagnosis of involvement of hepatoduodenal ligament and lymphnodes in biliary malignancies.

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EVALUATION OF LIVER FIBROSIS IN DIFFUSE LIVER DISEASE USING REAL-TIME TISSUE ELASTOGRAPHY
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[Objective] Real-time Tissue Elastography (RTE) is developed for visualizing the tissue hardness/softness by using ultrasound. We have been investigating its ability of evaluating fibrosis in diffuse liver disease. Recently, newly developed low frequency probe (EUP-L52) has been applied to RTE, and a patient who had difficulty of visualizing RTE image for the reason of low penetration such as obesity has been improved. In this study, multiple linear regression analysis was performed using several features of RTE image to estimate the RTE fibrosis value, and compared with the fibrosis stage to evaluate the clinical usefulness of RTE.

[Material and Method] 26 patients with chronic hepatitis C or liver cirrhosis diagnosed by liver biopsy, and 6 healthy volunteers were examined in this study. The indicated stages of fibrosis were F0 in 2 patients, F1 in 6 patients, F2 in 8 patients, F3 in 6 patients, and F4 in 4 patients. RTE were performed with HITACHI HI VISION 900 and EUP-L52 linear probe(3-7MHz). Scan was performed through the right intercostal space to observe right lobe. Probe was slightly held to detect the strain by heartbeat. All RTE images were transferred to an external PC, and analyzed with prototype image analysis software. Color data inside the ROI were converted to relative strain value, and features of RTE image such as mean of relative strain value (MEAN), standard deviation of relative strain value (STD), area of blue region (AREA), and complexity of blue region (COM) were calculated. Then, multiple regression analysis was performed with features of RTE image and fibrosis stage.

[Results] Features of RTE image were highly correlated with fibrosis stage. Correlation coefficient of MEAN, STD, AREA, and COM were r=-0.604, 0.593, 0.592, and 0.578. With these 4 parameters, multiple regression analysis was performed and derived the regression equation, which significantly fit with the data. RTE fibrosis value was calculated from this equation and had high correlation with fibrosis stage(r=0.729).

[Conclusion] As a result of having analyzed RTE quantitatively, the quantity of characteristic reflected staging well. RTE is particularly useful as the modality that can grasp improvement of the fibrosis by a hepatitis diagnosis and the treatment non-invasively.

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TISSUE REAL_TIME ELASTOGRAPHY IN AUTOIMMUNE PANCREATITIS
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Introduction: Endoscopic ultrasound (EUS) elastography is a recent imaging procedure that distinguishes tissues due to their specific consistency and has been recently evaluated in patients with histologically proven pancreatic tumours. The aim of this prospective evaluation was to investigate the role of this new technique for characterisation of mass lesions caused by autoimmune pancreatitis.

Summary: Prospective single centre evaluation enrolling 5 patients with mass lesions finally diagnosed as autoimmune pancreatitis. Elastographic recordings were compared to histology as gold standard in all 5 patients. Adequate and reproducible elastographic recordings of the pancreatic lesions could be obtained in all 5 patients with the final diagnosis of autoimmune pancreatitis. Patients with autoimmune pancreatitis presented with a characteristic stiff elastographic pattern not only of the mass lesion but also of the complementary pancreatic tissue which is unique and could not been found in patients with ductal adenocarcinoma and healthy subjects. Final diagnosis was achieved by transcutaneous biopsy in all patients. None of the patients with mass lesions and autoimmune pancreatitis have been operated.

Conclusions: EUS elastography of the pancreas shows a typical and unique finding with homogenous stiffness of the whole organ discriminating this disease from the circumscribed mass lesions in ductal adenocarcinoma and other neoplasia. Knowledge of this finding helps to avoid unnecessary operations.

EuroEUS, April 30 – May 2nd, 2009, Berlin, Germany

REAL-TIME SONOELASTOGRAPHY - A NEW APPLICATION IN THE FIELD OF LIVER DISEASE
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Ultrasound elastography is a new imaging technique that allows a noninvasive estimation and imaging of tissue elasticity distribution within biological tissues using conventional real-time ultrasound equipment with modified software. Elastography has been reported to be useful for differentiation and characterization of various malignant tumors, such as breast, prostate, thyroid, pancreas, lymph nodes, gastrointestinal stromal tumors, hepatocellular carcinoma and liver metastasis. Transient and, more recently, real-time elastography has been proved to be useful for noninvasive assessment of liver fibrosis in patients with diffuse liver diseases. Elasticity imaging promises to make an important contribution to ultrasound practice.

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ENDOSCOPIC ULTRASOUND ELASTOGRAPHY FOR EVALUATION OF LYMPH NODES AND PANCREATIC MASSES: A MULTICENTER STUDY

06-07-09

AIM: To evaluate the ability of endoscopic ultrasound (EUS) elastography to distinguish benign from malignant pancreatic masses and lymph nodes.

METHODS: A multicenter study was conducted and included 222 patients who underwent EUS examination with assessment of a pancreatic mass (n = 121) or lymph node (n = 101). The classification as benign or malignant, based on the real time elastography pattern, was compared with the classification based on the B-mode EUS images and with the final diagnosis obtained by EUS-guided fine needle aspiration (EUS-FNA) and/or by surgical pathology. An interobserver study was performed.

RESULTS: The sensitivity and specificity of EUS elastography to differentiate benign from malignant pancreatic lesions are 92.3% and 80.0%, respectively, compared to 92.3% and 68.9%, respectively, for the conventional B-mode images. The sensitivity and specificity of EUS elastography to differentiate benign from malignant lymph nodes was 91.8% and 82.5%, respectively, compared to 78.6% and 50.0%, respectively, for the B-mode images. The kappa coefficient was 0.785 for the pancreatic masses and 0.657 for the lymph nodes.

CONCLUSION: EUS elastography is superior compared to conventional B-mode imaging and appears to be able to distinguish benign from malignant pancreatic masses and lymph nodes with a high sensitivity, specificity and accuracy. It might be reserved as a second line examination to help characterise pancreatic masses after negative EUS-FNA and might increase the yield of EUS-FNA for lymph nodes.

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FEASIBILITY OF TISSUE ELASTOGRAPHY USING TRANSCUTANEOUS ULTRASONOGRAPHY FOR THE DIAGNOSIS OF PANCREATIC DISEASES


Objectives: We investigated the feasibility of using real-time tissue elastography (EG) with transcutaneous ultrasonography (EG-US) for pancreatic diseases.

Methods: A preliminary study (phase I) and a prospective (phase II) study were conducted. Phase I: subjects were 10 volunteers, 5 with cancer, 2 with endocrine tumor, 5 with chronic pancreatitis, 14 with intraductal papillary-mucinous neoplasm. To determine the characteristic EG images (diagnostic criteria for phase II), B-mode images were compared with EG images and histopathologic findings. Phase II: 53 consecutive patients were enrolled. The visualization rate by EG-US in lesions visualized by B mode was assessed and the correct diagnosis rate by B-mode alone (B-diagnosis) or in combination with EG-US was evaluated.

Results: Phase I: normal parenchyma was a homogeneous color. In cancer, EG-US showed a markedly hard area with soft spots inside. Endocrine tumor was uniform and soft comparable to parenchyma. Chronic pancreatitis showed a mixture of various colors. Phase II: we identified 77.4% (41/53) of the lesions and observed 60.0% (15/25) of the cancers, 100% (3/3) of the endocrine tumor, 92.0% (23/25) of the cases of chronic pancreatitis cases on EG-US. The B-diagnosis rates ranged from about 70% - 80%. The diagnosis rates of the combination were more than 90% of lesions of each type.

Conclusions: The EG-US is feasible in the diagnosis of pancreatic diseases.


NEURAL NETWORK ANALYSIS OF DYNAMIC SEQUENCES OF EUS ELASTOGRAPHY

06-07-09
BACKGROUND: EUS elastography is a newly developed imaging procedure that characterizes the differences of hardness and strain between diseased and normal tissue.

OBJECTIVE: To assess the accuracy of real-time EUS elastography in pancreatic lesions.

DESIGN: Cross-sectional feasibility study.

PATIENTS: The study group included, in total, 68 patients with normal pancreas (N = 22), chronic pancreatitis (N = 11), pancreatic adenocarcinoma (N = 32), and pancreatic neuroendocrine tumors (N = 3). A subgroup analysis of 43 cases with focal pancreatic masses was also performed.

INTERVENTIONS: A postprocessing software analysis was used to examine the EUS elastography movies by calculating hue histograms of each individual image, data that were further subjected to an extended neural network analysis to differentiate benign from malignant patterns.

Main Outcome Measurements: To differentiate normal pancreas, chronic pancreatitis, pancreatic cancer, and neuroendocrine tumors.

RESULTS: Based on a cutoff of 175 for the mean hue histogram values recorded on the region of interest, the sensitivity, specificy, and accuracy of differentiation of benign and malignant masses were 91.4%, 87.9%, and 89.7%, respectively. The positive and negative predictive values were 88.9% and 90.6%, respectively. Multilayer perceptron neural networks with both one and two hidden layers (3-layer perceptron and 4-layer perceptron) were trained to learn how to classify cases as benign or malignant, and yielded an excellent testing performance of 95% on average, together with a high training performance that equaled 97% on average.

LIMITATION: A lack of the surgical standard in all cases.

CONCLUSIONS: EUS elastography is a promising method that allows characterization and differentiation of normal pancreas, chronic pancreatitis, pancreatic cancer, and neuroendocrine tumors. The currently developed methodology, based on artificial neural network processing of EUS elastography digitalized movies, enabled an optimal prediction of the types of pancreatic lesions. Future multicentric, randomized studies with adequate power will have to establish the clinical impact of this procedure for the differential diagnosis of focal pancreatic masses.


EUS ELASTOGRAPHY FOR PANCREATIC MASS LESIONS: BETWEEN IMAGE AND FNA?

Chronic pancreatitis and pancreatic cancer are major causes of pancreatic disease--related morbidity and mortality. Because these diseases often occur concurrently, detection of focal cancer in a background of inflammation can be especially difficult. The diagnosis and management of both chronic pancreatitis and pancreatic cancer rely heavily upon endoscopic and radiologic imaging. The main techniques include CT; magnetic resonance imaging (MRI); ERCP; and EUS, especially EUS-guided sampling methods (EUS-guided FNA [EUS-FNA] or EUS-guided Trucut biopsy [EUS-TCB]). Continued improvements in the technology and use of uniform diagnostic criteria have increased our ability to obtain accurate staging, diagnosis, and prognosis. Among these techniques, EUS and EUS-FNA and/or EUS-TCB have proven to be very powerful techniques to accurately diagnose chronic pancreatitis and pancreatic cancer. For EUS evaluation of pancreatic cancer, EUS-FNA is considered a safe, accurate, and efficient procedure; however, multiple biopsies (on average 5-7) are usually required, because of the inability to specifically target tumor cells that are admixed with inflammation and/or fibrosis.1 When compared with other imaging modalities, the results of EUS-FNA of pancreatic masses are excellent and achieve a sensitivity of 85% to 90% and a specificity of virtually 100%.2-4 However, lower (<75%) sensitivity is reported when there is a coexistent presence of chronic pancreatitis or “pseudotumoral” pancreatitis. 5,6 This is an important issue, because approximately 20% to 35% of patients who undergo EUS-FNA of pancreatic lesions have features of underlying chronic pancreatitis. 7,8 Given these limitations, new technologies, eg, EUS elastography, which measures tissue stiffness, are being investigated. Elastography uses sonic and US waves to compress
tissue. Tissue that is fibrotic and stiff will compress less than softer, healthy tissue. Because the malignant and chronic inflammatory stiff tissue is usually harder than the adjacent normal tissue, by measuring the tissue strain induced by compression, we can estimate tissue hardness, which may be useful to distinguish pancreatic cancer and chronic pancreatitis compared with normal tissue. In this month’s issue of Gastrointestinal Endoscopy, Saffo et al describe the accuracy of real-time EUS elastography in a group of 68 patients, hoping to differentiate between normal pancreas (n = 22), chronic pancreatitis (n=11), pancreatic cancer (n=32), and neuroendocrine tumors (n=3). In their study, they used clinical history and imaging criteria to define “normal pancreas,” and cytologic diagnosis, surgery, or 6-month clinical follow-up to distinguish malignancy from “pseudotumoral” pancreatitis. The investigators then performed a postprocessing analysis, based on neural networks, to analyze the EUS elastography video sequences. The degree of tissue stiffness was indicated by the elastography and was graphically represented by a color-hue scale. By using a cutoff of 175 for the mean hue histogram values recorded on the region of interest, differentiation of benign and malignant masses was achieved, with a high sensitivity (91.4%), specificity (88.9%), and accuracy (90.6%). Furthermore, in a subgroup analysis, good sensitivity (93.8%) and overall accuracy (86%) was reported to differentiate pancreatic cancer from “pseudotumoral” pancreatitis. However, the specificity was low (63.6%). When using a more stringent cutoff, of 190, specificity increased to 90.9%, and a positive predictive value of 94.7% was demonstrated but at the sacrifice of sensitivity. These results suggest that this technique can be complementary to EUS in its ability to improve the diagnostic accuracy of pancreatic mass lesions of chronic pancreatitis versus pancreatic cancer, which is a major limitation of EUS-FNA. EUS elastography has previously been shown to distinguish benign and malignant pancreatic lesions and lymph nodes. However, a study by Janssen et al reported that EUS elastography has limited accuracy (60%) for diagnosing chronic pancreatitis. This is likely because these studies used qualitative electrophysiological analyses (predominant color, distribution of colors, and constancy of pattern), which are subject to interobserver variation. In addition, chronic pancreatitis may also have some degree of tissue stiffness similar to pancreatic cancer, which can make the differentiation difficult. Safto et al were able to overcome this limitation by developing a quantitative, thus objective, image interpretation algorithm by using a sophisticated statistical method called artificial neural network analysis. Artificial neural network analyses are computational analytical tools that are inspired by the biological nervous system. These consist of networks of highly interconnected computer processors called “neurons,” which are capable of performing parallel computations for data processing and knowledge representation. Just the way the human brain processes information, a neural network learns through repeated adjustments of the weighted interconnections and can infer complex data. There are several issues that should be considered about this study. Perhaps the most significant issue is of placing this technology in the context of our current clinical algorithm. Most patients with pancreatic cancer are treated with chemotherapy or surgical resection. For chemotherapy, a tissue diagnosis is a nearly absolute requirement. It is unlikely that any imaging methods, particularly one with a specificity of <95%, will replace biopsy. For potentially surgical patients, the main issue is to exclude “nonsurgical” diseases, such as autoimmune pancreatitis, metastatic cancers to the pancreas, or chronic pancreatitis. In this regard, the current study is unlikely to change current clinical practice. The main value may be to guide biopsy in cases where there is diffuse inflammatory change, to possibly reduce the number of biopsies needed, and to further exclude disease in patients with a low probability of cancer, such as those with an equivocal CT or MRI (eg, “prominent head of pancreas” in the absence of a defined mass). Another value, although not specifically evaluated in this study, may be to distinguish normal pancreas from chronic pancreatitis. Although EUS has generally been shown to be accurate, there is only modest interobserver agreement among expert endosonographers (k=0.45). An objective measure, such as elastography, may provide some needed standardization. Other efforts to standardize EUS criteria for chronic pancreatitis include the Rosemont classification, which gives different weights to each EUS criteria in predicting the diagnosis of chronic pancreatitis. Future studies should compare traditional methods, weighted qualitative methods, and new quantitative methods, such as EUS elastography. Other key technical issues to resolve with elastography are how to integrate it into real-time EUS. In the current study, all images were analyzed off-line under more ideal circumstances. As with other image interpretation methods, the technology must provide the information at the time a decision must be rendered, namely, at the time of the procedure. Also, it would be very helpful to consider applying elastography to other methods of imaging the pancreas, such as MRI. Recent advances in MRI-elastography, particularly for detection of fibrosis and/or cirrhosis of the liver, are highly promising and have the potential to offer an even less-invasive diagnosis of chronic pancreatitis. In summary, the use of EUS with different complementary invasive and noninvasive techniques is evolving and aims to provide accurate...
differentiation of pancreatic masses. The use of neural network analysis of dynamic sequences of EUS-elastography is a promising new technique that may provide important additional information to distinguish benign and malignant pancreatic lesions. The optimal role of this technique is likely to guide EUS-FNA to regions most likely to harbor cancer and to exclude cancer in patients with a low probability. Further investigation will be required to determine these outcomes.

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Abbreviations: EUS-FNA, EUS-guided FNA; EUS-TCB, EUS-guided trucut biopsy; MRI, magnetic resonance imaging.

REFERENCES
INDICATIONS AND LIMITATIONS FOR ENDOSONOGRAPHY WITH ELASTOGRAPHY IN PATIENTS WITH PANCREATIC DISEASE

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Objectives: Recently, endoscopic ultrasound has been using elastography to provide an image-guided method for distinguishing tissue with regard to its specific consistency. The objective of this prospective study was to examine to what extent this new technology could be used to differentiate and characterise normal pancreatic tissue and focal pancreatic diseases.

Patients and methods: Prospective study, 108 patients with unclassified pancreatic lesions and 10 patients with a healthy pancreas. The elastographic results were compared with histology as the gold standard.

Results: It was only possible to document adequate elastographic examinations for 30% of the study population. The fundamental limitation was the restricted penetration depth, which did not allow for reproducible elastographic depiction for lesions with diameters above 30 mm (58% of the study population). The second most common reason was artefacts (15%), primarily in patients with calcified chronic pancreatitis and cystic lesions. By contrast, it was possible to carry out adequate and reproducible elastograms on all healthy organs (soft to intermediate specimens) and on the majority of small pancreatic masses < 30 mm (hard specimens). The clinical benefit of differential diagnosis is slight as all pancreatic masses, which were harder than the surrounding pancreatic tissue, were represented irrespective of their nature. The only exception was patients with autoimmune pancreatitis for whom a consistently harder specimen was detected, which enabled these patients to be distinguished from patients with other entities.

Conclusion: Endosonography with elastography provides additional and complementary information for improved tissue characterisation in patients with pancreatic disease. The clinical benefit remains unclear.

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FREEHAND REAL-TIME ELASTOGRAPHY: IMPACT OF SCANNING PARAMETERS ON IMAGE QUALITY AND IN VITRO INTRA- AND INTEROBSERVER VALIDATIONS.

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Real-time elastography is a method for visualization of the elastic properties of soft tissue and may potentially enable differentiation between malignant and benign pathologic lesions. Our aim was to validate the method on a tissue-mimicking (TM) phantom and to evaluate the influence of different scanning parameters and investigator variability. A TM-phantom containing eight spherical inclusions with known storage modulus was examined using two different transducers on an ultrasound (US) scanner equipped with software for real-time elasticity imaging. The ultrasound transducers were moved vertically in a repetitive manner to induce strain. Two investigators performed series of standardized elastography scans applying a 0-4 categorical quality scale to evaluate the influence of
seven parameters: dynamic range of elasticity, region-of-interest, frequency of transducer movement, rejection of elastogram noise, frame rate, persistence and smoothing. Subsequently, repeated examinations of four selected inclusions were performed using a visual analog scale (VAS) where investigators marked a 100 mm horizontal line representing the span in image quality based on experience from the first examination. The hardest and softest inclusions were imaged more clearly than the inclusions with elasticity more similar to the background material. Intraobserver agreement on elastogram quality was good (kappa: 0.67 - 0.75) and interobserver agreement average (kappa: 0.55 - 0.56) when using the categorical scale. The subsequent VAS evaluation gave intraclass-correlation coefficients for the two observers of 0.98 and 0.93, respectively, and an interclass-correlation coefficient of 0.93. Real-time elastography adequately visualized isoechoic inclusions with different elastic properties in a TM-phantom with acceptable intra- and interobserver agreement. Dynamic range of elasticity was the parameter with most impact on the elastographic visualization of inclusions.


ENDOSCOPIC ULTRASOUND ELASTOGRAPHY IN THE DIFFERENTIAL DIAGNOSIS OF PANCREATIC SOLID MASSES: TOWARDS THE VIRTUAL BIOPSY
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Endoscopic ultrasound elastography (EUS-E) allows analyzing tissue stiffness during a standard endoscopic ultrasound (EUS) examination, thus providing with additional information about the features of solid lesions. The aim of the study was to evaluate the elastographic patterns of solid pancreatic masses and the diagnostic accuracy of EUS-E in this setting.

METHODS: 80 consecutive patients (mean age 62 years, 24-84 y., 54 males) who underwent EUS-E for the evaluation of solid pancreatic masses, and 10 controls with normal pancreas (mean age 59 years, 20-83 y., 5 males), were prospectively included in the study. EUS-E was performed under conscious sedation by the linear Pentax EG 3830 UT and Hitachi EUB 8500. EUS-FNA was performed for cytological diagnosis after elastographic examination. The different elastographic patterns of solid pancreatic masses are described. Diagnostic sensitivity, specificity, PPV, NPV and overall accuracy of EUS-E for malignancy were calculated compared to final diagnosis based on cytology, histology of surgical specimens and clinical follow-up.

RESULTS: Mean size of pancreatic masses was 31.7 ± 13.6 mm. Tumors were located in the head of the pancreas in 61 patients, in the body in 16 and in the tail in 3. Final diagnosis was pancreatic cancer (PC) in 48 cases, inflammatory mass (IM) in 23 patients, neuroendocrine tumor (NT) in 8 cases and metastatic tumor in 1 patient. 4 different patterns could be defined: 1) Homogeneous green, 2) heterogeneous green-predominant with yellow and red lines, 3) heterogeneous blue-predominant with a geographic appearance and green and red lines, and 4) homogeneous blue. A homogeneous green pattern was only observed in normal pancreas. 18 masses showed a heterogeneous green-predominant pattern, and all of them were IM. Thus, the probability of IM is 100% in the presence of this pattern. A heterogeneous, geographically distributed, blue-predominant pattern was found in 54 cases. This pattern was observed in all cases of malignant tumor (either PC or metastasis) and in 5 cases of IM. Thus, the probability of malignancy in the presence of this pattern is of 90.1%. A homogeneous blue pattern was exclusively seen in the 8 patients with NT. Thus, the probability of NT is 100% in the presence of this pattern. Sensitivity, specificity, PPV, NPV and overall accuracy for malignancy were 100%, 78.3%, 91.9%, 100% and 93.7% respectively.

CONCLUSION: EUS-E is a very useful tool for the differential diagnosis of solid pancreatic masses. This technique adds important information to EUS evaluation by providing with highly specific atterns, which strongly support the benign or malignant nature of the disease.

Digestive Disease Week, May 17th – 22nd, 2008, San Diego, USA (366)
NEURAL NETWORK ANALYSIS OF DYNAMIC SEQUENCES OF EUS ELASTOGRAPHY USED FOR THE DIFFERENTIAL DIAGNOSIS OF CHRONIC PANCREATITIS AND PANCREATIC CANCER
Adrian Saftoiu, Peter Vilmann, Florin Gorunescu, Dan Ionut Gheonea, Marina Gorunescu, Tudorel Ciurea, Gabriel Lucian Popescu, Alexandru Iordache, Sevastita Iordache

BACKGROUND Endoscopic ultrasound (EUS) elastography is a newly developed imaging procedure that characterizes the differences of hardness and strain between diseased tissue and normal tissue.
AIM The aim of our study was to prospectively assess the accuracy of EUS elastography to differentiate between normal pancreas (N=22), chronic pancreatitis (N=11), pancreatic adenocarcinoma (N=32) and pancreatic neuroendocrine tumors (N=3). A post-processing analysis based on specially designed software was used to analyze the EUS elastography movies by calculating hue histograms of each individual image. Furthermore, a neural network analysis based on average hue histograms of the EUS elastography movies was tested in order to differentiate benign versus malignant EUS elastography patterns.
RESULTS Based on the study group of 68 patients, the sensitivity, specificity and accuracy of differentiation of benign and malignant cases were 91.4%, 87.9% and 89.7%, based on a cut-off of 175 for the mean hue histogram values recorded inside the region of interest. The positive predictive value and negative predictive value were 88.9% and 90.6%, respectively. A multi-layer perceptron (MLP) neural network was trained to learn how to classify cases as benign or malignant, and yielded a very good testing performance of 95% in average, together with a high training performance equaling 97% in average.
CONCLUSIONS EUS elastography is a promising method that allows characterization and differentiation of normal pancreas, chronic pancreatitis and pancreatic cancer. The method might prove useful for the differentiation of pancreatic tumors, as well as to target the EUS-FNA biopsy into harder regions suspicious of malignancy. The currently developed methodology based on artificial neural network processing of the EUS elastography digitalized movies, enabled an optimal prediction of the pancreatic lesions type. Future multicentric, randomized studies with adequate power will have to establish the clinical impact of this procedure for the differential diagnosis of focal pancreatic masses.

Digestive Disease Week, May 17th – 22nd, 2008, San Diego, USA (671)

ENDOSCOPIC ELASTOSONOGRAPHY IS HIGHLY PREDICTIVE OF DEFINITE PATHOLOGY
Jan-Werner Poley, E. J. Kuipers

BACKGROUND: Elastosonography is a technique that allows for real-time assessment of elasticity or stiffness of tissue with the aid of conventional ultrasound instruments. In recent years this technique has been developed for endoscopic ultrasound (EUS). Preliminary data show that it might be a useful adjunct to standard morphological criteria to define a lesion without FNA sampling, or to select the optimal location for FNA to increase the yield of tissue sampling. To further elucidate this hypothesis we evaluated the performance of elastosonography in series of patients referred for EUS.
METHODS: patients were examined by an experienced endosonographer with a linear echoendoscope (Pentax 3830-UT) combined with a Hitachi EUB-8500 system to allow for realtime processing of elastonography images. Based on these images lesions were classified as either benign, malignant or indeterminate based on previously published data. A definite diagnosis was made on results of either EUS-FNA, surgical pathology or clinical follow-up (at least six months).
RESULTS: 41 procedures were performed in the same number of patients (M/F 17/24, age range 26 to 74 yrs): 25 procedures were done for mediastinal pathology, 13 for pancreatic masses, 1 each for rectal, adrenal and submucosal lesions. Elastosonography images were interpreted as benign in 10 cases (24%), indeterminate in 7 cases (17%), and malignant in 24 cases (59%). All patients with a benign result on elastosonography had benign disease during follow-up and EUS-FNA. Of the patients with lesions that were judged to be indeterminate on elastosonography (n=7; 17%) 5 had benign disease based on clinical follow-up and EUS-FNA whereas 2 had malignant disease based on EUS-FNA. In the group with malignant elastosonography (n=24) one patient had benign disease. A comparison of the elastography classes benign plus indeterminate versus malignant yielded a
sensitivity of elastosonography of 92%, a specificity of 94%, and negative and positive predictive values of respectively 96% and 88%.

CONCLUSION: Elastosonography is highly sensitive and specific in determining the nature of both mediastinal and pancreatic lesions with excellent positive and negative predictive values and can therefore be helpful in guiding EUS-FNA and patient management.

*Digestive Disease Week, May 17th – 22nd, 2008, San Diego, USA, (760)*

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**ENDOCOPIC ULTRASOUND ELASTOGRAPHY FOR THE UPPER GASTROINTESTINAL TRACT DISEASE**

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BACKGROUND & AIM: It is known that some diseases, such as cancer, lead to changes in the hardness of tissue. Endoscopic Ultrasound Elastography (EUE) is a new technique to assess the elasticity of tissue. It is expected to improve the ability for diagnosis. The aim of this study was to evaluate EUE to differentiate between benign and malignant gastrointestinal masses and lymph nodes.

PATIENT & METHOD: From July 2006 to September 2007, 35 patients underwent EUE examinations. Nine patients underwent evaluation of esophageal cancer, 7 of gastric cancer, 14 of submucosal tumor, 2 of mediastinal lymph nodes, which all cases were diagnosed histopathologically after examinations.

RESULT: All cases could be drawn images of EUE, which can be classified into three patterns. Type 1 is a common finding among the normal walls of digestive tract, which showed about the same hardness as nearby fatty tissues and was demonstrated from red to green on the images. Type 2 and 3 were image findings of tumors or inflammatory diseases. Both were demonstrated to be harder than nearby adipose tissues. We classified the cases demonstrated mosaic-like into type 2 and ones demonstrated uniformly into type 3. In the stomach or esophageal cancer cases, a neoplasm part was drawn as type 2 or type 3 and change of hardness was closely similar to the loupe image of pathology. About the lymph node including stomach or esophageal cancer cases, the malignant lymph node was drawn as type 3, but the inflammatory lymph node the view of type 2. In the submucosal tumor, GIST was shown as type 3 and aberrant pancreas was shown as type 1. No complication occurred with EUE examinations.

CONCLUSION: EUE could make it much easier to diagnose benign or malignancy of extrinsic lymph nodes. In the field of diagnosis of invasion depth of stomach or esophageal tumor, we have impression that the border in deep parts becomes clearer.

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**USEFULNESS OF THE CHARACTERIZATION OF TISSUE HARDNESS OF PANCREATIC MASS USING ELASTOGRAPHY ENDOSCOPIC ULTRASOUND. -FIRST TRIAL OF THE QUANTIFICATION USING STRAIN RATIO-**

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INTRODUCTION: Cancer is starting the change of tissue hardness by its own fibrosis from early stage. The reconstruction of tissue elasticity provides the sonographer with important additional information which can be applied for the diagnosis of these diseases.

AIM: The aim of our study was to evaluate the ability of endoscopic ultrasound elastography to differentiate between benign and malignant pancreatic masses.

PATIENTS AND METHODS: The subjects were 53 patients performed an endoscopic ultrasound(EUS) for pancreas in our hospital till September 2006 to October 2007. The disease were 5 with tumor forming pancreatitis(TFP), 32 with pancreatic cancer(PCA), 1 with GIST, 1 with renal cell carcinoma pancreatic metastases, 1 with IPMA and 13 with normal control. A histological diagnosis by
surgery or endoscopic ultrasonography fine needle aspiration (EUS-FNA) was performed for all subjects. The ultrasound was used the HITACHI HI VISION900, and EUS scope was PENTAX EG-3630UR and EG-3670URK. The calculation of tissue elasticity distribution is performed in realtime and the results are represented in color over the radial B-mode image. Malignant tissue appeared in blue color, fibrosis in blue to green, and normal tissue in green to red. In addition, we performed the quantification by using strain ratio (non mass area/mass area:SR) in order to evaluate the objective hardness as numerical value between mass to non mass area especially to distinguish TFP from PC.

RESULTS: Elastography for all PCs showed intense blue coloration, which indicated that the mass lesions had malignant aspects. While TFP presented the coloration pattern of mixed green, yellow and low intensity of blue. GIST and renal cell carcinoma pancreatic metastasis presented green and yellow pattern. Normal control was an even application of green to red. The mean SR of TFP and PC were each 24.99±12.54 and 42.08±21.57, respectively, which was significant difference (p<0.05)

CONCLUSION: EUS elastography is potentially capable of further defining the tissue characteristics of benign and malignant lesions. This study suggested that it was useful for the quantification by using strain ratio to characterize the tissue hardness of pancreatic disease and distinguish TFP from PC.

Digestive Disease Week, May 17th – 22nd, 2008, San Diego, USA (M1443)

INDICATIONS AND LIMITATIONS OF TISSUE REAL-TIME ELASTOGRAPHY USING ENDOSCOPIC ULTRASOUND IN PANCREATIC DISEASE

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Aim: Sonoelastography is an imaging procedure that distinguishes tissues due to their specific consistency. The aim of this prospective study was to investigate the role of this new technique for characterisation and differentiation of normal pancreatic parenchyma and pancreatic tissues affected by various focal diseases. Patients and methods: Prospective single center study, enrolling 108 patients with unclassified focal lesions of the pancreas and ten controls. Elastographic recordings were compared to histology as gold standard in all patients.

Results: Adequate elastographic recordings of the pancreatic lesions could be obtained in 30% of our study population. This was mainly due to limited depth penetration (58%), artifacts (15%), that were particularly prominent in patients with calcified chronic pancreatitis and cystic lesions. In contrast, adequately and reproducible elastographic recordings were obtained from all healthy pancreatic organs (soft/intermediate pattern) and the majority of lesions smaller than 30 mm independent of their dignity (hard pattern). Patients with autoimmune pancreatitis presented with a unique stiff elastographic pattern that allowed for discrimination from all other disease entities.

Conclusion: EUS elastography of the pancreas has the potential to achieve complementary information for improved tissue characterization especially in autoimmune pancreatitis.

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ENDOSONOGRAPHIC ELASTOGRAPHY IN THE DIAGNOSIS OF MEDIASTINAL LYMPH NODES

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Background and study aims: Ultrasonographic elastography is a new technique for describing the mechanical properties of tissue during real–time ultrasonography. The aim of this study was to test the feasibility of this method in endosonography (EUS) of the dorsal mediastinum, and to compare the elastographic patterns of lymph nodes with results from EUS–guided fine–needle aspiration biopsy (FNAB).
Patients and methods: 50 consecutive patients undergoing EUS-guided FNAB of at least one paraesophageal lymph node were included. Each of these targeted lymph nodes was examined also elastographically. The elastographic patterns were described and compared with the histologic results by a first examiner. The elastographic classification was subsequently further tested by two blinded reviewers.

Results: In total, 66 lymph nodes were examined; 37 lymph nodes revealed benign and 29 malignant tissue at the histologic evaluation. Good elastographic records were obtained for all lymph nodes. Of the 37 benign lymph nodes, 31 showed a homogeneous pattern of intermediate elasticity, whereas a dominance of hard tissue with variable patterns was found in 23 of 29 malignant lymph nodes. Applying these criteria, the accuracy range among the three examiners was between 81.8% and 87.9% for benign lymph nodes and between 84.6% and 86.4% for malignant ones. The interobserver agreement was excellent (kappa = 0.84).

Conclusion: EUS elastography of mediastinal lymph nodes can be performed reliably. The results are good for a noninvasive technique, but they remain inferior to the success rate of EUS-guided FNAB. The method might occasionally be useful for targeting the most suitable lymph nodes for FNAB.

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DYNAMIC ANALYSIS OF ENDOSCOPIC ULTRASOUND ELASTOGRAPHY USED FOR THE DIFFERENTIATION OF CHRONIC PANCREATITIS AND PANCREATIC CANCER
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INTRODUCTION: Endoscopic ultrasound elastography is a new imaging procedure that allows the reconstruction of elasticity distribution by characterizing the difference of hardness between diseased tissue and normal tissue.

AIMS & METHODS: The aim of our study was to describe EUS characteristics encountered in 28 patients prospectively included with normal pancreas (N = 3), chronic pancreatitis (N ~ 5), pancreatic adenocarcinoma (N = 17) and pancreatic neuroendocrine tumors (N = 3). The pattern of real-time EUS elastography images was compared with the conventional EUS aspects, while the final diagnosis was obtained by EUS-FNA cytology analysis, by surgical pathology or minimum 6 months followup.

RESULTS: EUS elastography patterns were compared in patients with pancreatic cancer, neuroendocrine tumors, chronic pancreatitis and normal pancreas. Post-processing analysis based on hue histograms of the EUS elastography average images was also done, yielding a semi-quantitative method to quantify the elasticity of the pancreatic tissue. The movies were analysed and filtered by computer-enhanced dynamic analysis using a public domain Java-based image processing tool (ImageJ) developed at the National Institutes of Health. The sensitivity, specificity and accuracy of differentiation of pancreatic adenocarcinoma and neuroendocrine tumors were 80.0%, 91.7% and 84.4%, based on a cut-off of 175 for the mean hue histogram values recorded on the region of interest. The positive predictive value and negative predictive value were 94.1% and 73.3%.

CONCLUSION: EUS elastography is a promising method that allows characterization and differentiation of normal pancreas, chronic pancreatitis and pancreatic cancer. The method might prove useful for the differentiation of pancreatic tumors, as well as to target the EUS-FNA biopsy into harder regions suspicious of malignancy. However, a post-processing analysis through the use of special software is required to correctly characterize the patients, while future studies on an increased number of patients are awaited.

15th United European Gastroenterology Week, October 29th – 31st, 2007, Paris, France

USEFULNESS OF THE CHARACTERIZATION OF TISSUE HARDNESS OF PANCREATIC MASS

06-07-09
USING ELASTOGRAPHY ENDOSCOPIC ULTRASOUND
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INTRODUCTION: Cancer is starting the change of tissue hardness by its own fibrosis from early stage. The reconstruction of tissue elasticity provides the sonographer with important additional information which can be applied for the diagnosis of these diseases.

AIMS & METHODS: The aim of our study was to evaluate the ability of endoscopic ultrasound elastography to differentiate between benign and malignant pancreatic masses. The subjects were 32 patients performed an endoscopic ultrasound (EUS) for pancreas in our hospital till September 2006 to April 2007. The disease were 4 with tumor forming pancreatitis (TFP), 20 with pancreatic cancer (PC), I with GIST and 7 with normal control. A histological diagnosis by surgery or endoscopic ultrasonography fine needle aspiration (EUS-FNA) was performed for all subjects. The ultrasound was used the HITACHI HI VISION900, and EUS scope was PENTAX EG-3630UR and EG-3670URK. The calculation of tissue elasticity distribution is performed in realtime and the results are represented in color over the radial B-mode image. Malignant tissue appeared in blue color, fibrosis in blue to green, and normal tissue in green to red. In addition, we performed the quantification by using strain ratio (non mass area / mass area: SR) in order to evaluate the objective hardness as numerical value between mass to non mass area especially to distinguish TFP from PC.

RESULTS: Elastography for all PCs showed intense blue coloration, which indicated that the mass lesions had malignant aspects. While TFP presented the coloration pattern of mixed green, yellow and low intensity of blue. GIST presented green and yellow pattern. Normal control was an even application of green to red. The mean SR of TFP and PC were each 25.23±11.06, and 46.53±19.48, respectively, which was significant difference (p ~ 0.0026).

CONCLUSION: EUS elastography is potentially capable of further defining the tissue characteristics of benign and malignant lesions. This study suggested that it was useful for the quantification by using strain ratio to characterize the tissue hardness of pancreatic disease and distinguish TFP from PC.

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ENDOSONOGRAPHIC ELASTOGRAPHY OF THE ANAL SPHINCTER – A PROSPECTIVE STUDY
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(Translated from German)

Background: Inflammatory conditions and tumours can lead to an alteration of the tissue elasticity of organs, which can be confirmed and displayed with Elastography (so-called SELA) as a colour overlay of the B-mode. For certain tumours and inflammatory diseases it has been proven that SELA can provide the physician with potentially important information. We examined patients suffering from rectal incontinence to determine if the Endosonographic Elastography (so-called e-SELA) appearances of the anal sphincter correlate to the functional parameters as well as if it is in relation to traditional incontinence treatment.

Method: 30 Patients suffering from rectal incontinence (n=19 Rectum/Sigmoid resection, n=5 Crohn’s disease, n=6 other). After evaluation of the rectal incontinence by using incontinence scores (CACP) and measuring manometer functional parameters, an anal 2D-Endosonography (EUS) examination of the internal/external sphincter (IS/ES) was performed with SELA (SonoElastography) (Hitachi EUB-8500). A semi-quantitative evaluation was made with a colour scale from 0-3: red (soft) through to blue (hard). After 2 weeks bio feedback / pelvic floor training, a second evaluation was carried out.

Results: The elasticity of the internal sphincter was significantly higher than the elasticity of the external sphincter, as a result, the hardness of the external sphincter was significantly higher that the...
hardness of the internal sphincter. There was a negative correlation by stimulated pressure on the anal sphincter but not by non-stimulated pressure.

The hardness (blue colouring) of the external sphincter was clearly higher for men that for women (2.0±0.4 vs. 1.0±0.4, p<0.031) and decreases with age, with significantly high values in the age category from 40-49 years (2.0±0.44) in comparison with the age range from 70-79 years. (1.0±0.83, p<0.02).

**Conclusion**: This examination shows that the E-SELA provides important additional information in relation to clinical significant sphincter characteristics such as elasticity/hardness. Further clinical studies (to evaluate the effect of treatment and treatment decision) should be determined in prospective controlled studies.

ELASTOSONOGRAPHY IN MALIGNANT RECTAL DISEASE: PRELIMINARY DATA

G. Mezzi, P. G. Arcidiacono, S. Carrara, C. Boemo, P. A. Testoni:

Elastography is an ultrasound technique that allowed to obtain images from the mechanical properties of soft tissue. It is well known that some diseases, such as cancer, lead to changes in tissue hardness.

Elasticity imaging is a technique that reveals the physical properties of tissue and can determine changes in tissue hardness caused by disease [1±5]. We evaluated 20 consecutive patients (10 males, 10 females; mean age 61.15 years, range 29±84 years) with histologically confirmed distal rectal cancer for disease staging. Informed consent was given by all of subjects. All patients underwent rectal endoscopic ultrasound, which was performed using a flexible echo endoscope (an oblique-viewing instrument [fiber oblic or electronic video image] [Pentax, Hamburg, Germany]). The echoscope was inserted and advanced beyond the lesion, under direct vision, to the rectosigmoid junction. Tumors were targeted to determine the depth of infiltration into or through the rectal wall. Frequencies commonly used for T-staging range from 7.5 MHz to 9 MHz. The endoscopic ultrasound elastography score was subdivided into five types on the basis of the echopattern distortion of the examined area, from 1 (normal) to 5 (advanced malignant neoplasia) (l" Figure 1). Rectal staging was performed according to Tumor–Node–Metastasis staging system (TNM).

It was our experience that the elastography images showed a discreet correlation in the staging of advanced lesions (T3, 65%); and post-radiotherapy disease persistence has been correctly confirmed by elastography score (T3, 100%). These preliminary data have shown that elastography, performed during endoscopic ultrasound, is a method that can differentiate between benign and malignant rectal lesions, although further studies, in particular regarding the follow-up of neoadjuvant chemoradiation therapy and in benign lesions, are necessary.

References

1 Frey H. Realtime elastography: a new ultrasound procedure for the reconstruction of tissue elasticity (in German, English abstract). Radiologie 2003; 43: 850±855
2 Lorenzen J, Sinkus R, Adam G. Elastography: quantitative imaging modality of the elastic tissue properties [in German, English abstract]. Rofo 2003; 175: 623±630
Background: Initial clinical applications have shown that US elastography might be able to distinguish tissues because of their specific consistency.

Objective: (1) To investigate the feasibility of EUS elastography of the pancreas and (2) to describe elastographic patterns of the normal pancreas and the pancreas affected by inflammatory or focal disease.

Design: Prospective single-center study.

Setting: Academic center of the University of Witten/Herdecke.

Patients: Twenty patients with normal pancreas, 20 patients with chronic pancreatitis, and 33 patients with focal pancreatic lesion, histologically later proven in 32 of these 33 cases.

Interventions: Commercially available US equipment was used. The elasticity of tissue was reconstructed in real time within a sample area and was translated into a color scale imaging relative tissue elasticity within this area. Representative loops of at least 20 seconds were recorded regarding each region of interest.

Results: Adequate elastographic recordings could be obtained in all 73 patients. Patients with hypoechoic and intermediately echogenic normal pancreas revealed a relatively homogeneous elastographic pattern. Thirty-one focal lesions, including 30 neoplasms and most of the chronically inflamed pancreata had a honeycomb pattern dominated by hard strands. This pattern showed analogies to the histologic structure of 10 resected tumors. Other patients with chronic pancreatitis and those with hyperechoic healthy pancreas had miscellaneous elastographic appearances.

Figure 1: Elastosono-graphic images in a patient with a T3 rectal malignancy.
Conclusions: EUS elastography of the pancreas is feasible and produces plausible results. The examination of homogeneous tissue is impaired by the relative scale used. Chronic pancreatitis and hard tumors cannot be distinguished by elastography, probably because of their similar fibrous structure.


Dynamic Analysis of EUS Used for the Differentiation of Benign and Malignant Lymph Nodes
Adrian Saftoiu, Peter Vilmann, Tudorel Ciurea, Gabriel Lucian Popescu, Alexandru Iordache, Hazem Hassan, Florin Gorunescu, Sevastita Iordache, (Hellerup, Denmark, Craiova, Romania)

Background: EUS elastography was reported to offer supplemental information that allows a better characterization of tissue characteristics and that might enhance conventional EUS imaging.

Objective: Our purpose was to apply real-time elastography during EUS examinations and to assess the accuracy of the differentiation of benign versus malignant lymph nodes.

Design: Prospective cross-sectional feasibility study.

Setting: Department of Surgical Gastroenterology, Gentofte University Hospital, Hellerup, Denmark.

Patients: Patients diagnosed by EUS with cervical, mediastinal, or abdominal lymph nodes were included, with a total number of 78 lymph nodes examined. The final diagnosis of the type of lymph node was obtained by EUS FNA cytologic analysis or by surgical pathologic examination and by a minimum 6 months of follow-up.

Interventions: Hue histogram analysis of the average images computed from EUS elastography movies was used to assess the color information inside the region of interest and to consequently differentiate benign and malignant lymph nodes.

Main Outcome Measurements: Differentiate between malignant and benign lymph nodes.

Results: By using mean hue histogram values, the sensitivity, specificity and accuracy for the differential diagnosis were 85.4%, 91.9%, and 88.5%, respectively, on the basis of a cutoff level of 166 (middle of green-blue rainbow scale). The proposed method might be useful to avoid color perception errors, moving artifacts, or possible selection bias induced by analysis of still images.

Limitations: Lack of the surgical standard in all cases.

Conclusions: Computer-enhanced dynamic analysis based on hue histograms of the EUS elastography movies represents a promising method that allows the differential diagnosis of benign and malignant lymph nodes, offering complementary information added to conventional EUS imaging.


Pressed for an Answer: Has Elastography Finally Come to EUS?
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In this issue of Gastrointestinal Endoscopy, Saftoiu et al present their findings from a prospective study of elastography used during EUS evaluation of benign and malignant lymph nodes. The study represents a step forward in the transition of this fascinating technology from an investigational tool to a clinically meaningful test. In 1991, Ophir et al coined the term “elastography” to describe an imaging technique that conveyed information about a tissue’s
relative firmness in response to compression, meaning it is more akin to palpation than inspection. Various forms of elastography exist, being used within the fields of optics, magnetic resonance imaging, and US. In the area of US, elastography can be based upon a tissue’s response to compression to measure tissue deformation (compression elastography), a tissue’s displacement in response to short bursts of vibration (transient elastography), or a tissue’s effects on vibration amplitudes in response to continuous vibration (vibration elastography). Gastroenterologists may already be familiar with transient elastography in the form of the FibroScan (Echosens, Paris, France), a device that uses US to measure the effects of transient vibrations through the liver to measure the organ’s stiffness or degree of fibrosis. But to better understand compression elastography, it is prudent to first review how US images themselves are created.

Recall that a US transducer contains specialized (piezoelectrical) crystals that vibrate at specific frequencies when exposed to an alternating electrical field. These vibrations emit a pulse of sound waves that travel into target tissues, some of which ultimately “bounce” back off the tissues. Immediately after launching this burst of waves, the crystals become “ears,” awaiting the return of reflected sound waves (echoes) bouncing back from the tissues. Upon arrival, returning waves set the crystals vibrating once again, and these vibrations are converted to electrical impulses used to construct an image. The returning echoes, which permit creation of an image, carry key information about the tissues. First, the time it takes for the echo to return conveys how far away, or deep, that structure is from the transducer. Second, the echo’s intensity reflects the difference in densities (impedance) at the interface between adjacent structures off of which the sound wave “bounced.” The traditional B (“brightness”) mode images we are used to seeing in EUS convey each echo’s intensity on a black and white (mostly gray) scale. Many single beams of echoes are then aligned to construct the familiar 2-dimensional image. In “classic” compression elastography, stress is applied to the tissue by pressing the transducer slightly harder against the body. In the study by Saftoiu et al, however, the stress applied was simply the mild compression that tissues experience during respiration and vascular pulsations.

Stress applied to a tissue results in strain or deformation within the tissue. Harder tissues deform less than softer tissues, and, as discussed, US echoes carry information about tissue location and density, both of which change slightly with deformation. Thus, by comparing echoes made with and without compression, or simply over several seconds of normal breathing and blood circulation, one can obtain information about how hard or soft the tissues are relative to their surroundings. Calculations of the relative elasticity of all tissues within a viewing trapezoid-shaped window (similar to Doppler imaging) are made in real time by the EUS processor, and a false-color overlay is superimposed on the B-mode images. In the current study, the software expressed elasticity in numerical values that ranged from 0 to 255 and used a rainbow spectrum of color in the overlay to provide visual cues about the tissues’ hardness. Firm tissues appeared blue, soft tissues appeared red, and those in the intermediate range fell somewhere in the blue-green spectrum. How can we exploit this information? We know that cancer tends to be firmer than normal tissue. Therefore, Saftoiu et al, in addition to other endosonographers, attempted to use elastography to differentiate benign from malignant tissue based on EUS imaging. In the current study, the investigators focused on lymph nodes and compared EUS imaging criteria (ie, node size, shape, border, and echotexture characteristics), EUS-guided FNA results, and elastography findings. Their criterion standard was a composite of FNA results, surgical specimen findings, and clinical course. By using receiver operating characteristic curves, they determined that an elasticity cutoff of 166 yielded the greatest accuracy in distinguishing benign from malignant nodes. Not surprisingly, EUS criteria performed poorly in differentiating benign from malignant nodes, as others have demonstrated. FNA performed quite well, although this is in the context of being used as part of its own criterion standard. But, the exciting finding in the study was a reasonable sensitivity (85%) and good specificity (92%) for elastography when using the elasticity value of 166 to diagnose malignancy. The overall prevalence of malignant denopathy in the study was 53%, so the positive and negative predictive values of elastography were 92% and 85%, respectively. This means elastography in this study had a positive likelihood ratio (LR) of 10.5 and a negative LR ratio of 0.16. What does this mean clinically? A positive LR of 10.5 means that a positive test changes the disease probability by approximately 45%.

So, if you think a node has a 50% likelihood of being malignant based on EUS criteria or just the clinical context (eg, a 2-cm mediastinal node adjacent to a T3 esophageal cancer) and elastography suggests it is malignant, then the node is 95% likely to be malignant. Therefore, besides helping you choose which nodes to preferentially aspirate, elastography may prove useful in staging nodes that are deemed inaccessible, perhaps because of intervening tumor or vessels. A negative LR of 0.16 changes the post test probability of disease approximately –35%.
So, a node with a 50% pretest probability of harboring cancer is only about 15% likely to be positive if elastography suggests benignity. Is this cutoff value of 166 reliable? Not yet. The investigators are the first to acknowledge that this was based upon a single, relatively small data set and must be validated prospectively among new patient populations. Furthermore, there may be different optimal cutoffs based upon a node's location. For example, elastography must compare a node's hardness with the surrounding tissues to determine its response to compression. It is possible that the supporting structures in the mediastinum are different than those in the abdomen, and, therefore, future studies must determine if computer algorithms should consider a node's location when choosing the cutoff value for diagnosing malignancy.

A potential major limitation to elastography is the degree of fibrosis that may be present in benign tissue, making the tissue firm and decreasing the accuracy of the test for diagnosing malignancy. For example, do benign reactive mediastinal nodes in a chronic smoker contain sufficient fibrosis to cause false-positive results for cancer? Furthermore, just how much cancer must be present to make a lymph node harder than usual? We are always concerned about false negative FNA results when only very small amounts of cancer are present within a lymph node. It is unclear whether elastography would prove to be any more sensitive than FNA. In fact, there were 2 false-negative FNAs in the current study, but the investigators did not discuss the elastography findings from those cases. It would be interesting to know whether elastography classified those cases as positive for malignancy.

The investigatiors should be commended for tackling another important shortcoming of elastography, namely the difficulty in using a single image for calculating degrees of elasticity. This is problematic, because an arbitrary choice of image could introduce bias (the endosonographer may preferentially choose the "bluer" frames when most other frames depict a red hue). This would also create problems with inter- and intraobserver variation. To overcome this problem, the investigators integrated several seconds' worth of images into a computerized summation value to calculate the elasticity within the region of interest. They also developed software to drop images with hue distributions outside a reliable range, thereby excluding images most likely to represent artifact. These new protocols or some variation upon them will probably be incorporated into newer generations of the software used for elastography. Are there other foreseeable EUS uses for this technology? One can imagine that the elastography images of a gastric GI stromal tumor might be different from that of a pancreatic rest but probably not too different from a leiomyoma, limiting its utility in subepithelial lesions. Tiny intraductal probe-based systems are unlikely to reliably distinguish benign from malignant biliary or pancreatic strictures, because these are always firm, regardless of malignancy or benignity. Perhaps serous cystadenomas of the pancreas can be distinguished from mucinous cystic neoplasms based upon the viscosity of the cyst contents, a bit of an EUS "Holy Grail" if you will. Avoiding any FNA that carries a risk of pancreatitis is a laudable goal. In the end, this technology is likely suited for exactly what these investigators are doing: trying to improve our ability to stage malignancy by either providing a "virtual biopsy" or, more likely, guiding our needles to improve FNA accuracy and decreasing the number of FNA passes. The relative simplicity of applying elastography during EUS suggests that, if the "pesky details," eg, reliability, can be worked out, we are likely to see this technology develop into a standard adjunct to EUS. Just how pressed we will be to adopt elastography will undoubtedly depend upon the firmness of future data.


QUALITY ASSESSMENT OF FREE-HAND SONOELASTOGRAPHY IN VITRO

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Introduction: Sonoelastography is used to visualise elastic properties in soft tissue, and the method seems to have a promising ability to differentiate between malignant and benign pathologic lesions. We have evaluated the influence of different scanning parameters and the investigator performance on the quality of free-hand sonoelastography in vitro.

Material and method: A tissue mimicking phantom containing eight spherical inclusions of known elastic moduli was examined with 7.5 Hz and 9 MHz linear transducers connected to an ultrasound...
scanner with sonoelastography programme (Hitachi EUB-8500). The probes were moved in a
sinusoidal, repetitive manner to induce strain. Two investigators made separate scan series applying
first a 0-4 categoric scale and a subsequently a Visual Analogue Scale (VAS) to score 7 parameters:
dynamic range of elasticity (E-dyn), region of interest (ROI), frequency of transducer movement,
rejection of elastogram noise, frame rate, persistence and smoothing.

**Summary:** The hardest and softest inclusions were imaged more clearly than those with elasticity
close to the background material. Intraobserver agreement in image quality assessment was good
(Kappa: 0.67-0.75) and interobserver agreement average when using the categoric scale (Kappa:
0.55-0.56). The subsequent VAS evaluation showed better intra- and interobserver agreement with
Intra-class correlation coefficients (Intra-CC) 0.983 and 0.931 (95%CI) and Interclass correlation
coefficient (Inter-CC): 0.932 (95%CI).

**Conclusion:** Sonoelastography adequately visualised isoechoic inclusion bodies in a TM-phantom
with good intra- and interobserver agreement. The quality of the elastograms were mostly influenced
by dynamic range of elasticity and this variable must be taken into account when scanning humans.

EuroEUS, 4th - 5th May, 2007, Seville, Spain

ENDOSCOPIC SONOELASTOGRAPHY – INITIAL EXPERIENCE.

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**Background:** Sonoelastography is an ultrasound method to visualise elastic properties of soft tissue
based on the distribution of strain in response to stress. Tissue patholgy may alter the elastic
properties and malignancy seems to generate harder tissue in most cases. Hence, sonoelastography
may be a valuable tool in differentiating malignancy from other pathologic changes.

**Material & Method:** We performed sonoelastography in 23 patients using a linear echoendoscope
(Pentax GF3830-UT) and a Hitachi EUB-8500 ultrasound scanner. On admission, 18 of the patients
had suspected malignant tumour/metastasis in the mediastinum (9), pancreas (4), oesophagus (2) or
other locations(3). Five patients without suspected malignancy had chronic pancreatitis (2) or a
myogenic tumour in upper GI-tract (3). EUS-FNA was performed in 8 patients and EUS-guided tru-cut
biopsy in 4 patients.

**Summary:** Seven tumours with suspected malignancy appeared harder than the surrounding tissue,
all of which proved to be carcinomas. Two large GISTs with malignant changes presented with mixed
elastograms. Six suspected tumours failed to present areas with harder tissue, 3 of these were
sampled with no malignancy and none showed indications of malignancy during follow-up. One
malignant mesothelioma of the pleura appeared softer than the reference tissue. One pancreatic
cancer and one case of pancreatitis generated inconclusive sonoelastograms. The five patients
without initial suspicion of malignancy all had lesions with negative elastograms.

**Conclusion:** Endoscopic sonoelastography is a promising method to record tumour elasticity and
may be helpful in detecting malignancy. Increased hardness as compared with surrounding tissue
seems to indicate malignancy in most cases.

EuroEUS, 4th - 5th May, 2007, Seville, Spain

DYNAMIC ANALYSIS OF ENDOSCOPIC ULTRASOUND (EUS) ELASTOGRAPHY
USED FOR THE DIFFERENTIATION OF BENIGN AND MALIGNANT LYMPH NODES
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06-07-09
INTRODUCTION: Endoscopic ultrasound (EUS) elastography is a recently developed imaging technique that enhances conventional EUS imaging by visualizing the tissue elasticity distribution, because it shows differences in hardness between various structures.

AIM: The aim of our study was to apply real-time elastography during EUS examinations and to assess the accuracy for the differentiation of benign versus malignant lymph nodes. A postprocessing computer-enhanced dynamic analysis based on hue histograms of the EUS elastography movies was tested. The final diagnosis of the type of lymph node was obtained by EUS-FNA cytology analysis and/or by surgical pathology, as well as by a minimum six months follow-up of the patients.

RESULTS: Patients diagnosed by EUS with cervical, mediastinal or abdominal lymph nodes were prospectively included, with a total number of 78 lymph nodes examined by EUS elastography. Hue histogram analysis of the average images computed from EUS elastography movies was used to assess the color information inside the region of interest and to consequently differentiate benign and malignant lymph nodes. Based on mean hue histogram values the sensitivity, specificity and accuracy for the differential diagnosis were 85.4%, 91.9% and 88.5%, respectively, based on a cut-off level of 166 (middle of green-blue rainbow scale).

CONCLUSIONS: Computer-enhanced analysis based on hue histograms of the EUS elastography movies represents a promising method that allows the differential diagnosis of benign and malignant lymph nodes with a high sensitivity, specificity and accuracy, offering complementary information added to conventional EUS imaging.

EuroEUS, 4th-5th May, 2007, Seville, Spain

ANALYSIS OF ENDOSCOPIC ULTRASOUND ELASTOGRAPHY USED FOR CHARACTERISATION AND DIFFERENTIATION OF BENIGN AND MALIGNANT LYMPH NODES
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Abstract:
Purpose: Ultrasound elastography is a new imaging procedure which allows the reconstruction of elasticity distribution by characterising the difference of hardness between pathological and normal tissue.
Materials and Methods: The aim of our study was to apply real-time elastography during endoscopic ultrasound (EUS) examinations and to consequently characterise benign versus malignant lymph nodes. The pattern of real-time EUS elastography images was compared with the conventional EUS aspects of lymph nodes and with the final diagnosis obtained by EUS-FNA cytology analysis and/or by surgical pathology.
Results: Patients diagnosed by EUS with cervical, mediastinal or abdominal lymph nodes were prospectively included, with a total number of 42 lymph nodes examined by EUS elastography. By using a qualitative pattern analysis, we were able to differentiate between benign and malignant lymph nodes with a high sensitivity, specificity and accuracy (91.7%, 94.4% and 92.86%, respectively), based on five pre-defined patterns obtained on EUS elastography. A quantitative analysis based on histograms of the EUS elastography images also allowed an excellent discrimination between benign and malignant lymph nodes. Based on separate RGB channel histogram values, an "elasticity ratio" was further defined and yielded a sensitivity, specificity and accuracy for the differential diagnosis of 95.8%, 94.4% and 95.2% respectively, based on a cut-off level of 0.84.
Conclusion: EUS elastography is a promising method which allows characterisation and differentiation of benign and malignant lymph nodes with a high sensitivity, specificity and accuracy, offering complementary information added to conventional EUS imaging.
ELASTOGRAPHY AND CONTRAST-ENHANCED (SONOVUE) COLOR DOPPLER PANCREATIC ENDOSCOPIC ULTRASOUND TO CHARACTERIZE PANCREATIC MASS. RESULTS IN 25 PATIENTS

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INTRODUCTION: US contrast agent is widely used for abdominal US examination to differentiate benign than malignant liver tumors. It is well known that some diseases, such as cancer, lead to a change of tissue hardness. The reconstruction of tissue elasticity provides the sonographer with important additional information which can be applied for the diagnosis of these diseases. Elasticity imaging has recently attracted attention as a technique which directly reveals the physical property of tissue and enables us to determine the change of tissue hardness caused by diseases. The aim of this study was to applied US contrast agent injection (SONOVUE) and elastography guided by endoscopic ultrasound to characterize pancreatic mass.

AIMS & METHODS: Between May 2004 and April 2005, 25 patients (17 M and 8 F), mean age 62.5 years have had an endoscopic ultrasound (EUS) for a pancreatic mass. The lesion was located in the head of the pancreas (12 cases), in the body (5 cases) and in the tail (8 cases). The mean size of the lesion was 25 mm (range: 12-40 mm). An EUS-FNA was performed in all cases using a 22 gauges needle (Wilson-Cook). SONOVUE (8 ml) was injected intravenously and the lesion was studied in color-doppler before and after the injection. The US machine used was the platform HITACHI 6500 with harmonic system and the EUS scope was the PENTAX EG 38-UT. Pancreatic adenocarcinomas were described as without enhancement after SONOVUE injection. In another hand color Doppler enhancement after SONOVUE injection was showed in benign pancreatic nodule and endocrine tumor. Like colour Doppler examinations, tissue elasticity imaging was performed with EUS-scope EG 38-UT (PENTAX) and does not require additional instruments. The calculation of tissue elasticity distribution is performed in realtime and the examination results are represented in colour over the conventional B-mode image. Malignant tissue appeared in blue colour, fibrosis in green, normal tissue in yellow and fat in red.

RESULTS: No complication occurred during the study. The final diagnosis of the pancreatic lesion was obtained by EUS-FNA (22 cases) and by Surgery (3 cases). Final histology was pancreatic adenocarcinomas (15 cases), endocrine tumors (2 cases), nodule of chronic pancreatitis (4 cases), pancreatic sarcoma (1 case), and pancreatic metastasis (3 cases). Concerning pancreatic adenocarcinoma, 14/15 lesions were not enhanced after SONOVUE injection only 1 lesion was enhanced it was a malignant IPMT. Regarding the endocrine tumor, 2/2 lesions were enhanced after us contrast injection, Pancreatic sarcoma and pancreatic metastasis were also enhanced after SONOVUE as the nodules of chronic pancreatitis. Elastography showed malignant aspects (intense blue coloration) for all pancreatic adenocarcinomas, endocrine tumor, pancreatic metastasis and pancreatic sarcoma but all nodules of chronic pancreatitis presented benign aspects (mixed green and low intense of blue).

CONCLUSION: the association of a non enhanced pancreatic mass + intense blue lesion in elastography is highly in favour of a pancreatic adenocarcinoma (accuracy 93.3%). the association of a enhanced pancreatic mass + intense blue lesion in elastography is highly in favour of a pancreatic carcinoma but non adenocarcinoma (endocrine, metastasis) and the association of a enhanced pancreatic mass + mixed green and low intense blue lesion in elastography is highly in favour of a benign pancreatic nodule as nodule of chronic pancreatitis.

13th United European Gastroenterology Week, October 15th – 19th, 2005, Copenhagen, Denmark

IS ELASTOGRAPHY GUIDED BY ENDOSCOPIC ULTRASOUND CAN DIFFERENTIATE BENIGN THAN MALIGNANT LYMPH NODES?

06-07-09
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INTRODUCTION: It is well known that some diseases, such as cancer, lead to a change of tissue hardness. The reconstruction of tissue elasticity provides the sonographer with important additional information which can be applied for the diagnosis of these diseases. Elasticity imaging has recently attracted attention as a technique which directly reveals the physical property of tissue and enables us to determine the change of tissue hardness caused by diseases. The aim of this study was to applied elastosonography guided by endoscopic ultrasound to differentiate benign than malignant lymph nodes.

AIMS & METHODS: Between March 2004 and April 2005, 25 patients (16M and 9F), mean age 64.6 years have had an EUS-FNA of lymph nodes during the staging of lung cancer (6 cases), esophageal carcinoma (5 cases), gastric cancer (3 cases), pancreatic cancer (2 cases), for a suspicion of LN relapse of a kidney cancer (2 cases), of a breast cancer (2 cases). An EUS-FNA was also performed in 5 cases for isolated LN.35 LN were biopsied and studied in sonoelastography in the same session. LN were located in the mediastinum (17 cases), in the cervical area (3 cases), in the celiac area (5 cases) and in the aorto-caval area (6 cases). The mean size of the lesion was 19.3 mm (range: 8-35 mm). An EUS-FNA was performed in all cases using a 22 gauges needle (Wilson-Cook). The realtime elasticity imaging described in this study, was performed with the SonoElastography module that was integrated into the platform of the HITACHI EUB-8500 system. Like colour Doppler examinations, tissue elasticity imaging was performed with EUS-scope EG 38-UT (PENTAX) and does not require additional instruments. The calculation of tissue elasticity distribution is performed in realtime and the examination results are represented in colour over the conventional B-mode image. Malignant tissue appeared in blue colour, fibrosis in green, normal tissue in yellow and fat in red.

RESULTS: No complication occurred during the study. Final histology was malignant LN (24 cases) and inflammatory LN (11 cases). Sonoelastography of the LN concluded to a malignant LN in 24/24 cases. In the other hand, elastosonography concluded to a benign LN in 6/11 cases, 5 false positives were reported in this study. Sensibility and Specificity of Sonoelastography guided by EUS to differentiate benign than malignant LN were respectively of 100% and 61.5%, PPV and NPV were respectively 77% and 100%.

CONCLUSION: Sonoelastography will be in the future an adjunct to EUS and EUS-FNA to characterize the LN and to offer a better target in patient with multiple lymph nodes.

13th United European Gastroenterology Week, October 15th – 19th, 2005, Copenhagen, Denmark

ENDOSCOPIC ULTRASOUND ELASTOGRAPHY: THE FIRST STEP TOWARDS VIRTUAL BIOPSY? PRELIMINARY RESULTS IN 49 PATIENTS

Background and Study Aims: It is well known that some diseases, such as cancer, lead to changes in the hardness of tissue. Sonoelastography, a technique that allows the elasticity of tissue to be assessed during ultrasound examination, provides the ultrasonographer with important additional information that can be used for diagnosis. The aim of this study was to evaluate the ability of endoscopic ultrasound elastography to differentiate between benign and malignant pancreatic masses and lymph nodes.

Patients and Methods: During a 12-month period, 49 patients underwent endoscopic ultrasound (EUS) examinations with elastography, conducted by a single endoscopist. Twenty-four patients underwent evaluation of a pancreatic mass (mean diameter 24.7 ± 11.1mm) and 25 underwent evaluation of 31 lymph nodes. The mean diameter of the lymph nodes was 19.7 ± 8.6 mm, and they were found in the cervical area (n = 3), mediastinum (n = 17), celiac arterial trunk region (n = 5), and aortocaval region (n = 6).

Results: The sonoelastography images of pancreatic masses were interpreted as benign in four cases and malignant in 20. The sensitivity and specificity of sonoelastography in the diagnosis of malignant lesions were 100% and 67%, respectively. The sonoelastography images of the lymph nodes were interpreted as showing malignancy in 22 cases, benign conditions in seven,
and indeterminate status in two. The sensitivity and specificity of sonoelastography for evaluating malignant lymph-node invasion were 100% and 50%, respectively.

**Conclusions:** EUS elastography is potentially capable of further defining the tissue characteristics of benign and malignant lesions and can be used to guide biopsy sampling for diagnosis.

*Endoscopy 2006; 38: 1-5*

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**EUS SONO-ELASTOGRAPHY FOR EVALUATION OF PANCREATIC MASSES. RESULTS OF A PROSPECTIVE MULTICENTRIC STUDY.**

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US contrast agent is widely use for abdominal US examination to differentiate benign than malignant liver tumors. It is well known that some diseases, such as cancer, lead to a change of tissue hardness. The reconstruction of tissue elasticity provides the sonographer with important additional information which can be applied for the diagnosis of these diseases. Elasticity imaging has recently attracted attention as a technique which directly reveals the physical property of tissue and enables us to determine the change of tissue hardness caused by diseases. The aim of this study was to applied US contrast agent injection (SONOVUE) and elastography guided by endoscopic ultrasound to characterize pancreatic mass.

**Patients and methods:** Between October 2005 and February 2006, 121 patients (77 M and 44 F), mean age 63.6 years have had an endoscopic ultrasound (EUS) for a pancreatic mass. The lesion was located in the uninate process (14 cases) in the head of the pancreas (48 cases), in the isthmus (17 cases) in the body (29 cases) and in the tail (13 cases). The mean size of the lesion was 29.5 mm (range: 7-80 mm). An EUS-FNA was performed in all cases using a 19 or 22 gauge needle (Wilson-Cook). The US machine used was the platform HITACHI 6500 with harmonic system and the EUS scope was the PENTAX EG 38-UT. Like colour Doppler examinations, tissue elasticity imaging was performed with EUS-science EG 38-UT (PENTAX) and does not require additional instruments. The calculation of tissue elasticity distribution is performed in realtime and the examination results are represented in colour over the conventional B-mode image. Malignant tissue appeared in blue colour, fibrosis in green, normal tissue in yellow and fat in red.

**Results:** No complication occurred during the study. The final diagnosis of the pancreatic lesion was obtained by EUS-FNA (82 cases) and by Surgery (39 cases). Final histology was pancreatic adenocarcinomas (72 cases), endocrine tumors (16 cases), benign nodule of chronic pancreatitis (30 cases), and pancreatic metastasis (3 cases). Elastography showed malignant aspects (intense blue coloration) for all pancreatic adenocarcinomas, endocrine tumor, pancreatic metastasis and pancreatic sarcoma but all nodules of chronic pancreatitis presented benign aspects (mixed green and low intensity of blue). The LN sonoelastography classification was done in 5 scores 1 to 5. If we consider score 1,2 as benign and score 3, 4 and 5 as malignant. The sensibility, specificity, Predictive positive value and predictive negative value of EUS Sonoelastography to differentiate benign than malignant pancreatic mass were respectively 80.6%, 92.3%, 93.3% and 78.1%. The global accuracy of this new technology was 89.2%. The Predictive negative value for malignancy of score 1-2 was 77.4% and the predictive positive value for malignancy of score 3,4 and 5 was 92.8%

**To conclude:** EUS pancreatic sonoelastography is a very useful tool to differentiate benign than malignant pancreatic mass. EUS sonoelastography doesn’t concurrence EUS-FNA but gives more datas in case of non contributive EUS biopsy.

*EuroEUS, 28th – 29th April, 2006, Hamburg, Germany*

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**EUS SONO-ELASTOGRAPHY FOR LYMPH NODES STAGING. RESULTS OF A PROSPECTIVE MULTICENTRIC STUDY.**

06-07-09
It is well known that some diseases, such as cancer, lead to a change of tissue hardness. The reconstruction of tissue elasticity provides the sonographer with important additional information which can be applied for the diagnosis of these diseases. Elasticity imaging has recently attracted attention as a technique which directly reveals the physical property of tissue and enables us to determine the change of tissue hardness caused by diseases. The aim of this study was to applied elastosonography guided by endoscopic ultrasound to differentiate benign than malignant lymph nodes.

Patients and methods: Between October 2005 and February 2006, 101 patients (56M and 45F), mean age 61.1 years have had an EUS-FNA of lymph nodes during the staging of lung cancer (26 cases), esophageal carcinoma (25 cases), gastric cancer (13 cases), pancreatic cancer (12 cases), for a suspicion of LN relapse of a kidney cancer (2 cases), of a breast cancer (8 cases). An EUS-FNA was also performed in 15 cases for isolated LN. LN were located in the mediastinum (51 cases), in the cervical area (4 cases), in the celiac or mesenteric area (44 cases) and in perirectal space (2 cases). The mean size of the lesion was 20.1 mm (range: 7-50 mm). An EUS-FNA was performed in all cases using a 22 gauges needle (Wilson-Cook). The realtime elasticity imaging described in this study, was performed with the SonoElastography module that was integrated into the platform of the HITACHI EUB-8500 system. Like colour Doppler examinations, tissue elasticity imaging was performed with EUS-scope EG 38-UT (PENTAX) and does not require additional instruments. The calculation of tissue elasticity distribution is performed in realtime and the examination results are represented in colour over the conventional B-mode image. Malignant tissue appeared in blue colour, fibrosis in green, normal tissue in yellow and fat in red.

Results: No complication occurred during the study. Final histology was malignant LN (55 cases including 35 metastasis by an adenocarcinoma, 13 by a squamous cell carcinoma, 3 by an endocrine tumor, 1 melanoma and 5 lymphomas) and inflammatory LN (44 cases including 3 cases of sarcoidosis). The LN sonoelastography classification was done in 5 scores 1 to 5. If we consider score 1,2 and 3 as benign and score 4 and 5 as malignant.

The sensibility, specificity, Predictive positive value and predictive negative value of EUS Sonoelastography to differentiate benign than malignant LN were respectively 100%,83.3%,100% and 75%.

But, if we consider score 1,2 as benign and score 3, 4 and 5 as malignant.

The sensibility, specificity, Predictive positive value and predictive negative value of EUS Sonoelastography to differentiate benign than malignant LN were respectively 88.10%,88.13%,91.22% and 84.10%.

The accuracy of this new technique was between 88 and 89.10%.

Conclusion: Sonoelastography will be in the future an adjunct to EUS and EUS-FNA to characterize the LN and to offer a better target in patient with multiple lymph nodes.

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ENDOSCOPIC ULTRASOUND ELASTOGRAPHY - A NEW IMAGING TECHNIQUE FOR THE VISUALIZATION OF TISSUE ELASTICITY DISTRIBUTION

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Abstract
Endoscopic ultrasound (EUS) elastography is an imaging procedure used for the visualization of
tissue elasticity during usual EUS examinations. EUS elastography can be accomplished real-time with state-of-the-art ultrasound systems, with the images being represented in transparent color superimposed on the conventional gray-scale B-mode scans. The aim of this review was to introduce the potential range of applications of EUS elastography.

EUS elastography might be useful for the differentiation of benign and malignant lymph nodes, with a qualitative pattern analysis and a quantitative histogram analysis of the color images being used to adequately classify the lesions. Mapping of the tissue elasticity distribution might be useful for the differential diagnosis of focal pancreatic masses, especially in the setting of chronic pancreatitis where the accuracy of EUS-guided fine needle aspiration is also low. EUS elastography might also enhance the detection and differentiation of various solid tumors (adrenal tumors, submucosal tumors, etc.) situated nearby the gastrointestinal tract.

Routine use of EUS elastography thus offers supplemental information that enhances conventional EUS imaging, with a possible decrease in the number of unnecessary EUS-FNA procedures used for tissue confirmation. However, future enhancements of the EUS elastography technology, as well as prospective, randomized studies will probably establish the clinical impact of dynamic elasticity imaging.

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ACCURACY OF ENDOSCOPIC ULTRASOUND ELASTOGRAPHY USED FOR THE DIAGNOSIS OF MALIGNANT LYMPH NODES
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INTRODUCTION: Endoscopic ultrasound elastography is a recently developed imaging procedure that allows the reconstruction of elasticity distribution by characterizing the difference of hardness between diseased tissue and normal tissue.

AIMS & METHODS: The aim of our study was to use real-time EUS elastography for the differential diagnosis of benign and malignant lymph nodes. Due to the results of a previously published pilot study the pattern of real-time EUS elastography images combined with post processing histogram analysis were used to differentiate benign and malignant lymph nodes. Final diagnosis was based on a combination of EUS-FNA cytology analysis (used for confirmation of malignancy), surgical pathology or 6 months follow-up.

RESULTS: Patients diagnosed by EUS with cervical, mediastinal or abdominal lymph nodes were prospectively included, with a total number of 78 lymph nodes examined by EUS elastography. By using histogram analysis we were able to differentiate benign and malignant lymph nodes with a high sensitivity, specificity and accuracy (83.33%, 88.88% and 85.89%, respectively). Based on separate RGB channel histogram values, an “elasticity ratio” was further defined and yielded a higher sensitivity, specificity and accuracy, based on a cut-off level of 0.85.

CONCLUSION: EUS elastography is a promising method that allows the characterization and differentiation of benign and malignant lymph nodes with a high sensitivity, specificity and accuracy, offering complementary information added to conventional EUS imaging.

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EUS SONO-ELASTOGRAPHY FOR LYMPH NODES AND PANCREATIC MASSES STAGING - RESULTS OF A PROSPECTIVE MULTICENTRIC STUDY ON 222 PATIENTS
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06-07-09
INTRODUCTION: It is well known that some diseases, such as cancer, lead to a change of tissue hardness. Elasticity imaging has recently attracted attention as a technique which directly reveals the physical property of tissue and enables us to determine the change of tissue hardness. The aim of this study was to apply elastosonography guided by endoscopic ultrasound to differentiate benign than malignant lymph nodes and pancreatic masses.

AIMS & METHODS: Between October 2005 and February 2006, 101 patients (56 M and 45 F), underwent EUS-FNA of lymph nodes for staging of lung cancer (26 cases), esophageal carcinoma (25 cases), gastric cancer (13 cases), pancreatic cancer (12 cases), for a suspicion of LN relapse of a kidney cancer (2 cases), of a breast cancer (8 cases). The mean size of the lesion was 20.1 mm (range: 7-50 mm). During the same period, 121 patients (77 M and 44 F), underwent endoscopic ultrasound (EUS) for a pancreatic mass. The lesion was located in the uncinate process (14 cases), in the head of the pancreas (48 cases), in the isthmus (17 cases), in the body (29 cases) and in the tail (13 cases). The mean size of the lesion was 29.5 mm. An EUS-FNA was performed in all cases using 22 gauge needle. The realtime elasticity imaging described in this study was performed with the SonoElastography module integrated in the platform of the HITACHI EUB-8500 system.

RESULTS: No complication occurred during the study. 1. Concerning the LN, final histology was malignant LN (55 cases including 35 metastasis by an adenocarcinoma, 13 by a squamous cell carcinoma, 3 by an endocrine tumor, 1 melanoma and 5 lymphomas) and inflammatory LN (44 cases including 3 cases of sarcoidosis). The LN sonoelastography classification was done in 5 scores 1 to 5. Score 1, 2 was considered as benign and score 3, 4 and 5 as malignant, the sensibility, specificity, positive and negative predictive values of EUS sonoelastography to differentiate benign from malignant LN were respectively 88.10%, 88.13%, 91.22% and 84.10% with an accuracy of this new technique between 88 and 89.10%. 2. The final diagnosis of pancreatic lesions was pancreatic adenocarcinomas (72 cases), endocrine tumors (16 cases), benign nodule of chronic pancreatitis (30 cases), and pancreatic metastasis (3 cases). Elastography showed malignant aspects for all pancreatic adenocarcinomas, endocrine tumor, pancreatic metastasis and pancreatic sarcoma. All nodules of chronic pancreatitis presented benign aspects. The LN sonoelastography classification was done in 5 scores (1-5). If we considered the scores 1, 2 as benign and 3, 4 and 5 as malignant, the sensibility, specificity, positive and negative predictive values of EUS sonoelastography to differentiate benign from malignant pancreatic masses were respectively 80.6%, 92.3%, 93.3% and 78.1% with a global accuracy of this new technology of 89.2%.

CONCLUSION: Sonoelastography will be in the future an interesting adjunct to EUS and EUS-FNA to characterize the LN and pancreatic masses.

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