Evaluation of parenchymal thyroid diseases with multiparametric ultrasonography

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Abstract

Aim: Differential diagnosis of parenchymal thyroid diseases by gray-scale ultrasonography is quite difficult for a radiologist as the findings are very similar to each other. In this study we aimed to assess some quantitative spectral Doppler parameters, resistivity index (RI), acceleration time (AT), and quantitative elastography [shear wave velocity (SWV)] together to show their reliability for differential diagnosis of parenchymal thyroid diseases. Materials and Methods: We retrospectively reviewed findings of 227 patients (179 females, 48 males) that underwent spectral Doppler ultrasound and acoustic radiation force impulse between October 2013 and March 2016. Ages of the patients were between 18 and 74 years (39.52 ± 12.67). Based on clinical and laboratory findings, patients were divided into five groups (N: Normal, EH: Early Hashimoto, H: Late Hashimoto, M: Nodular Thyroid Disease, HM: Hashimoto + Nodular Thyroid Disease). Detailed statistical analyses were done on parameters such as age, gender, volume information, and RI, AT (ms), SWV (m/s). Results: No significant effect of gender or volume on the differentiation of disease pattern (Chi-square test: \( P = 0.306 \), Kruskal-Wallis test: \( P = 0.290 \)) was found in this study. RI (0.41 ± 0.06) and SWV values (1.19 ± 0.18 m/s) were the lowest. AT values (>55 ms) were the highest in EH group (area under the curve: 0.913). Existence of H decreased RI and SWV values, while it extended AT in a different thyroid disease. Conclusion: Thyroid parenchymal diseases could be classified and differentiated from each other by measuring RI, AT, and SWV values quantitatively. So, in suspicious cases, these parameters could be a reliable asset for differential diagnosis. Key words: Acoustic radiation force impulse imaging; elastography; Hashimoto disease; thyroid gland; ultrasonography

Introduction

Differential diagnosis in advanced stages of diffuse and nodular thyroid parenchymal diseases is quite difficult with gray-scale ultrasonography because findings are usually very similar to each other. Also, nodular changes in multinodular (M) form and a chronic autoimmune disease Hashimoto (H) could be seen together in clinical practice.11

Actually, chronic autoimmune disease may show different radiologic characteristics depending on its stage: for early-stage disease (Early Hashimoto, EH) ultrasonography is done at the beginning, and for chronic-stage disease (Chronic Hashimoto, H) ultrasonography is done when the patient is under a medical treatment. Different pathologic stages during progression of the disease are hard to differentiate from each other with the

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Although there are many studies regarding radiological differential diagnosis of nodules (nodule–pseudo-nodule or benign–malignant nodule) in the literature, there are not enough studies on differential diagnosis of parenchymal changes in heterogeneous parenchyma of H, due to diffuse or other nodular parenchymal diseases with multinodular dysplasia.

In this study, in addition to conventional US examination, quantitative spectral Doppler parameters (resistivity index, RI and acceleration time, AT) and quantitative elastography values (shear wave speeds) were evaluated together to show their diagnostic capacity on differentiation of parenchymal thyroid diseases.

Materials and Methods

The study protocol was approved by Local Ethics Committee. We retrospectively reviewed the findings of 227 patients (179 females, 48 males) that underwent spectral Doppler US and acoustic radiation force impulse (ARFI) between October 2013 and March 2016. Ages of patients were between 18 and 74 years (39.52 ± 12.67). All the measurements of the patients were done by the same radiologist (D.Y. with 7 years of ARFI and Doppler US experience), who were blinded to all clinical and laboratory findings. All examinations were performed with the same system (Siemens ACUSON S2000, Siemens Healthcare, Erlangen, Germany). Total volume measurements of the thyroid lobes were done by Linear Transducer (9L4) after gray-scale examinations. We started Doppler US with the lowest pulse repetition frequency (PRF) value that causes artifacts and then increased PRF step by step (we performed all examinations within 700–1000 Hz) with medium wall filter. RI-AT values were obtained by measuring automatically from proximal segment of the first main parenchymal branch of the inferior thyroid artery. Then, with longitudinal examination, central poles of both lobes were evaluated and by using the ARFI-virtual touch quantification (VTQ) method, shear wave velocities (SWV, m/s) were measured. Mean values of five consecutive measurements were obtained for each localization. During elastographic study, areas with nodular or pseudo-nodular contents were excluded and the focus of the study was maintained on only more uniform areas. Gray-scale US combined with RI-AT-SWV were classified as multiparametric imaging.

Patients with normal thyroid gray-scale US findings (homogeneous parenchyma with regular normal echogenicity), normal thyroid function test, and normal auto-antibody levels were included in normal group, while patient with abnormal thyroid gray-scale US findings (heterogeneity, pseudo-nodular appearance with surrounding echogenic septations) and abnormal Doppler US findings were accepted as abnormal. According to these findings, after 2 years of follow-up, based on clinical, biochemical parameters [thyroid stimulating hormone (TSH), triiodothyronine (T3), thyroxine (T4), anti-thyroid peroxidase antibody (anti-TPO), thyroglobulin antibody (anti-Tg)], all the patients were divided into five different groups: control group (normal, N); first detected, early-untreated Hashimoto disease (EH); chronic Hashimoto patients that are under treatment and/or follow-up (H); multinodular parenchymal hyperplasia (M); and nodular hyperplasia with Hashimoto (HM). And all the data obtained from these groups were recorded.

Statistical method

All the data except demographic properties of the study group were analyzed statistically. Demographic data were excluded from this study because there was no significant relation between our study targets and demographic properties. Gray-scale findings, such as diffuse heterogeneity–diffuse nodularity or visual Doppler assessment such as vascularization degree or heterogeneity, were also excluded for their dependence on the operator and the subjectivity of the measurements.

First descriptive statistics of variables in the context of the data set is presented in assessment. Continuous variables are expressed by mean, standard deviation, while discrete variables are expressed by their frequency.

On comparisons, continuous normally distributed variables were evaluated by Student’s t-test or analysis of variance. Abnormal variables were evaluated by Mann–Whitney U-test or Kruskal-Wallis H-test. Binary post hoc analyses were performed for comparisons that were found statistically significant in multigroup comparison analyses. Chi-square test was used for comparison of discrete variables. In order to determine the direction and strength of the relationship between two continuous variables, Pearson or Spearman correlation coefficient tests were used based on whether the data were distributed normally or not. In continuous variables, receive operator characteristics (ROC) analyses, area under the curve (AUC), sensitivity–specificity results were used for determination of cut-off values. Statistical Package for the Social Sciences, version 15 was used for analyses, and the results were compared to P < 0.05 level, with maintaining the level of confidence interval by 95%.

Results

In statistical analysis, the evaluated parameters such as thyroid volume, RI, AT, and SWV values [Figure 1] that was determined by multiparametric US, age, and sex and descriptive statistics of the data set was compiled in Table 1. No statistically significant difference was detected between the diagnostic groups by gender; no selectivity for
There was no statistically significant difference between N-M, HM-H, HM-EH groups, in the evaluation of AT values (P > 0.005). AT values of EH group (70.00 ± 28.28) were significantly longer than N (27.35 ± 13.66), M (26.60 ± 11.74), and H (46.74 ± 12.09) groups. In addition to that, AT values were also significantly longer in H and HM groups (45.45 ± 15.65) than N and M groups.

When RI values were evaluated, HM (0.49 ± 0.05) and H (0.49 ± 0.05) groups had statistically lower values than N (0.56 ± 0.05) and M (0.58 ± 0.06) groups. SWV values of EH (1.19 ± 0.18) group were statistically lower than M (1.72 ± 0.33) and HM (1.64 ± 0.42) groups and values of M group (1.72 ± 0.33) were significantly higher than H group (1.45 ± 0.38). In thyroid diseases, presence of H decreased SWV, while presence of M increased SWV.

Results of ROC analysis were used for differentiation of groups due to measured values; AT values <35 ms with 0.76 sensitivity and −0.46 specificity and with 0.645 AUC is distinguishing for N group. In gray-scale nodular parenchyma, AT values <35 ms (0.83 sensitivity, −0.60 specificity, and with 0.737 AUC) show that this pattern is compatible with M group. In these cases, the characteristic features for M group would be that SWV value being faster than 1.45 m/s (with 0.82 sensitivity, −0.54 specificity and with 0.669 AUC) and RI being bigger than 0.53 (0.83 sensitivity, −0.65 specificity, and 0.669 AUC) [Figure 2] (M group).

In the presence of rough heterogeneous parenchymal changes and pseudo-nodularity, M and H groups might be confusable. In these cases if AT is longer than 55 ms (0.77 sensitivity, −0.89 specificity, and 0.87 AUC), EH could be a more plausible diagnosis. Also in these cases, age, especially older than 56 (0.54 sensitivity, −0.88 specificity, and 0.753 AUC) and RI values lower than 0.48 (1 sensitivity, −0.85 specificity, and 0.968 AUC), is important for differential diagnosis of EH [Figure 3] (EH group).

Parenchymal nodular changes and increase of SWV (1.72 ± 0.33) supported multinodular thyroid disease, but if AT was longer than 35 ms (0.73 sensitivity, −0.62 specificity,
and 0.715 AUC), it showed that these cases are compatible with HM group.

Differentiation between the characteristics of HM group and H group is also required, because pseudo-nodules that develop in H disease could be confusable with true nodules of HM group. The cases with AT values are between 35 and 55 ms and are compatible with H or HM group. The RI values also did not show statistically significant differences between H and HM groups, so only SWV and age variables could be used for differential diagnosis between these two groups. The cases with AT values were between 35 and 55 ms and SWV values were <1.75 (0.84 sensitivity, −0.45 specificity, and 0.641 AUC) in group H. For this reason, in cases that had rough nodular parenchymal structure, SWV was distinctive, while AT and RI values were not [Figure 4] (H group).

Discussion

Up-to-date, thyroid US was used for the measurement of parenchymal volume, assessing vascular characteristic of gland, screening, and differentiation of the nodules.[1,2] After the technologic developments about the transducers and high-resolution screens, gray scale and Doppler examinations became easier.[3,4] Additionally, SWV expended the scope of elastography and enabled the quantitative examination of the nodules and the thyroid parenchyma with the help of hardware and software.[5-10] Besides thyroid nodule evaluations, many works reported value of elastography to detect changes of thyroid parenchyma in diseases that affects thyroid parenchyma including HT.[11,12]

There are a lot of new methods and developments in the area of Doppler US and US elastography in thyroid diseases and
this information could be learned from proper literatures or guidelines.[13,14]

In this study, instead of finding new parameters for differentiation of benign–malignant nodules which have been investigated frequently, our aim was to produce a supportive method for differentiation of the thyroid parenchymal diseases which had similar visualization pattern that radiologists often encountered in daily practice. Thyroidal diseases are very common both in our country and worldwide. High prevalence of these diseases may generate certain risks in our population. Furthermore, in recent literatures, seronegative Hashimoto cases are seen in 13% of the population, and Hashimoto increases the risk of papillary thyroid malignancy. So these findings support that H should be differentiated from other chronic parenchymal diseases even in remission period.[15‑18]

We had some limitations in our study. The same radiologist did all measurements. However, as the evaluation was done by quantitative features and statistical analysis, we believe that this situation did not have any negative effects on our study. On the contrary, visual vascularity evaluation is a subjective parameter. Quantitative parameters, including peak systolic values and diastolic flow, have been evaluated in various studies. However, reproducibility of these parameters was found low as they are mostly angle- and direction-dependent.[19,20]

Hence, we used parameters which are not angle- and direction-dependent, including AT and RI. We found one study that evaluated RI in the diagnosis of Hashimoto’s disease, in which results were compatible with our study.[21]

The values we obtained from this study are supporting the current literature. As a conclusion, we believe that minor changes observed in our study are based on the differences of the devices and the measurement methods, which the radiologist used.[22,23]

Consequently, diseases that manifest similar gray-scale and color Doppler US findings and which affect thyroid parenchyma could be differentiated from each other quantitatively by measurement of RI, AT, and SWV values. In our study, the AT values should be <35 m/s in thyroid parenchyma, which has homogenous parenchyma and should show normal echogenicity pattern. If thyroid parenchyma had rough-nodular structure, and AT was <35 ms, RI was equal or more than 0.54 ms and
SWV was equal or higher than 1.45 m/s, then diagnosis should be multinodular goiter. In the same group, if AT was between 35 and 55 ms, and RI was <0.54 ms, then the diagnosis should be HM. The most important parameter for differentiation of HM and pseudo-nodular structure of H with rough and heterogeneous parenchyma was SWV values being <1.75 m/s in H group. Differentiation of EH group was easier than other groups, with the help of long AT (>55 ms), low RI (<0.48), and low SWV (<1.55 m/s) values. In order to facilitate the readers to follow the results more conveniently, all the data were summarized and enrolled in Table 2.

In the light of these parameters, early and chronic Hashimoto disease, nodular hyperplastic thyroid diseases, or overlaps of these diseases could be differentiated from normal group or isolated forms of diseases. Also, quantitative data could be obtained for supporting clinical laboratory findings.

In conclusion, apart from subjective gray-scale US findings, parenchymal thyroid disease might be further classified with the aid of spectral Doppler US and quantitative ARFI elastography measurements, especially in suspicious cases (differentiation of senile parenchymal changes and chronic Hashimoto disease, diagnosis of autoantibodies negative Hashimoto disease, differentiation of pseudo-nodular structure from multinodular structure); diagnosis and follow-up could be done with more reliable parameters. We believe that the algorithm we used or similar ones could be considered in daily practice.

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Conflicts of interest
There are no conflicts of interest.

References
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Table 2: Distinctive parameters for each group

<table>
<thead>
<tr>
<th>Group</th>
<th>AT (ms)</th>
<th>RI</th>
<th>SWV (m/s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1-N</td>
<td>&lt;35</td>
<td>0.56±0.05</td>
<td>1.55±0.43</td>
</tr>
<tr>
<td>2-M</td>
<td>&lt;35</td>
<td>≥0.54</td>
<td>≥1.45</td>
</tr>
<tr>
<td>3-HM</td>
<td>35-55</td>
<td>&lt;0.54</td>
<td>1.64±0.42</td>
</tr>
<tr>
<td>4-H</td>
<td>35-55</td>
<td>&lt;0.53</td>
<td>&lt;1.75</td>
</tr>
<tr>
<td>5-EH</td>
<td>&gt;55</td>
<td>&lt;0.48</td>
<td>&lt;1.55</td>
</tr>
</tbody>
</table>

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