INTRODUCTION

Most thyroid diseases are evaluated by radionuclide scintigraphy, sonography or percutaneous needle biopsy. Sonography is perhaps the most definitive method for evaluation of nodules in a thyroid gland. However, the disadvantages of this technique include limited penetration in some anatomic locations, such as retrotracheal and substernal areas, and a limited field of view for assessing large masses. Further, current diagnostic methods may not enable the accurate detection of the malignant nature of a thyroid nodule, especially in a multinodular goiter.

Therefore, new techniques are being investigated to improve the evaluation of thyroid nodules.

The magnetic resonance (MR) imaging has a minor role in the evaluation of most patients with thyroid nodules or goiters, because of inadequate spatial and contrast resolution and only a few cases documented. However, technological advances have provided good spatial and contrast resolution in the neck including the thyroid gland, parathyroid gland and lymph node. Also it has made available the dynamic keyhole imaging of the thyroid nodule with spin echo sequences.

The aim of our study was to evaluate this new technique in the routine evaluation of possible thyroid cancer, and particularly for the determination of the characteristic MR dynamic enhancement pattern of malignant thyroid tumor.

Materials and Methods: Eight patients who were pathology proven to have a malignant thyroid tumor, preoperatively. There are 5 papillary carcinomas, 1 medullary carcinoma, 1 follicular carcinoma, and 1 fine needle aspiration biopsy proven atypical cell.

Results: Based on preoperative MR imaging, we compared the dynamic MR enhancement pattern relating to the pathologic type. On contrast agent-enhanced dynamic T1-weighted image (T1WI), 5 papillary carcinoma and one medullary carcinoma showed delayed enhancement compared to normal parenchyma. In addition, one follicular carcinoma shows stronger enhancement than normal parenchyma, with one papillary carcinoma showing a persistent decrease in enhancement compared to normal parenchyma.

Conclusion: Although this study is limited by a small patients population, the data suggests that delayed enhancement on enhanced dynamic T1WI is a possible characteristic MR finding of a malignant thyroid tumor. I think that the comparison of MR imaging between benign and malignant nodules is required for a correct characterization.
enhancement pattern of a malignant thyroid tumor.

**MATERIALS AND METHODS**

This study series included eight patients (1 male and 7 females; mean age: 56 years; age range: 22-81 years) who were pathologically proven to have a malignant thyroid tumor, preoperatively; 5 papillary carcinoma, 1 medullary carcinoma, 1 follicular carcinoma, and 1 fine needle aspiration biopsy (FNAB) proven atypical cell (with multiple metastatic lymph nodes in the neck). Preoperative MRI images had been obtained in all patients.

We got the approval from the Institutional review board and informed consent from each patient part of this study. A pathologic report and ultrasonography (USG) findings were collected retrospectively.

All images were obtained with a 3.0-T MR imaging unit, Magnetom Verio (Siemens Medical Systems, Erlangen, Germany).

The MRI protocol was constituted of pulse sequences including an axial fast spin-echo T2-weighted image (T2WI), axial T1-weighted image (T1WI), and contrast agent-enhanced dynamic T1WI.

1) The axial fast spin-echo T2WI was obtained in approximately 3 minutes, 41 seconds. Imaging was performed in the 30 axial planes with 3-mm slice thickness, and no interslice gap. The field of view was 140 cm and the voxel size was $0.4 \times 0.4 \times 3.0$ mm. The T2WI were acquired with a repetition time (TR) of 4,100 ms and an echo time (TE) of 137 ms. The flip angle was 90 degrees.

2) The axial T1WI was obtained in approximately 6 minutes, 57 seconds. Imaging was performed in the 30 axial planes with a 3-mm slice thickness, and no interslice gap. The field of view was 140 cm and the voxel size was $0.5 \times 0.5 \times 3.0$ mm. The T1WI were acquired with a TR of 2,740 msec and an TE of 48 msec. The flip angle was 70 degrees.

3) The dynamic spin-echo sequence was a routine axial T1 spin-echo sequence obtained during contrast infusion.

The image was obtained in approximately 9 minutes, 13 seconds. Imaging was performed in the 88 axial planes with a 0.7-mm slice thickness and no interslice gap. Slice oversampling was 36.4%. The field of view (read/phase) was 255 mm/50.0%, and the voxel size was $0.7 \times 0.5 \times 0.7$ mm. The images were acquired with a TR of 5.12 msec and an TE of 1.99 msec. The flip angle was 10 degrees and time to center was 26.4 sec.

Contrast enhanced dynamic T1WI were obtained immediately after injection (as a precontrast scan) and at 1 min 30 sec/3 min/4 min 30 sec/6 min after injection of Gadovist.

Paramagnetic contrast agent (gadobutrol, Gadovist; Schering, Berlin, Germany) was administered intravenously at a weight-adjusted dose of 0.15 mmol per kilogram of patient body weight.

**RESULTS**

Based on preoperative MR imaging, we compared the dynamic MR enhancement pattern relating to the pathologic type. All biopsy-proven malignant thyroid tumors showed hypoechogenicity on previous USG, except for one follicular carcinoma (isoechogenicity). In addition, only one patient with papillary carcinoma remained on the color Doppler USG. In this case, the malignant nodules showed decreased vascularity compared to normal parenchyma.

1) On a T1WI, one papillary carcinoma showed high signal intensity (SI) and one medullary carcinoma showed low SI compared to normal parenchyma. And, other cases were not differentiated compared to normal parenchyma.

2) On a T2WI, 3 papillary carcinomas and one follicular carcinoma showed high SI, and one papillary carcinoma showed low SI. The other case was not differentiated with normal parenchyma.

3) On contrast agent-enhanced dynamic T1WI, 4 papillary carcinomas and one medullary carcinoma, as well as 1 FNAB proven atypical cell (Fig. 1) showed delayed enhancement compared to normal parenchyma. And, one follicular carcinoma (Fig. 2) showed stronger enhancement than normal parenchyma, and one papillary carcinoma (Fig. 3) shows persistently decreased enhancement compared to normal parenchyma.

About 75% of all cases showed delayed enhancement pattern on contrast-enhanced dynamic T1WI (Table 1).

Our study found that the maximum gland-lesion contrast difference occurred 3 to 4 minutes and 30 seconds after contrast infusion.

In addition, two patients with benign nodules (nodular hyperplasia) underwent the same MR imaging. One patient had a ho-
Fig. 1. On ultrasonography, an approximately 8 mm, well defined hypoechoic nodule is seen in the mid level of the left lobe of the thyroid gland. This is dynamic images of the thyroid gland in the axial section. The nodule (arrow) was not clearly delineated immediately after contrast injection (as a precontrast scan). Early-phase dynamic images obtained 1 minute 30 seconds after injection best depicts the nodule (arrow) in left thyroid gland. The nodules show decreased enhancement in the early phase of a dynamic enhancement study, compared to normal parenchyma. In addition, the nodule (arrow) shows gradually centripetal delayed enhancement. A 6-minute image after contrast injection indicates that the nodule is masked into the background thyroid tissue.

Note. — FNAB = fine needle aspiration biopsy

Fig. 2. On ultrasonography, an approximately 3 centimeter, well defined hyperchoic nodule (white arrow) is seen in right lobe of the thyroid gland at the mid level. On a dynamic image obtained 3 minutes after injection, the nodule (white arrow) shows strong early enhancement compared to the parenchyma of the left lobe. In addition, the nodule (white arrow) shows persistent well enhancement, compared to normal parenchyma. In a 6-minute image after contrast injection, the nodule (white arrow) shows persistent enhancement, but background parenchyma is washed out.
characteristic dynamic enhancement pattern of magnetic resonance imaging for malignant thyroid tumor. The delayed enhancement pattern of thyroid cancer on dynamic T1WI is similar to that of pituitary adenoma. We hypothesize that the enhancement pattern of these tumors is related with tumor angiogenesis and its vascularity. Pituitary tumors are known to be less vascular than normal pituitary tissue, suggesting that angiogenesis may be inhibited in these tumors. This is consistent with the findings of Jugenburg and colleagues, but is in marked contrast to studies in other tissues including the breast, prostate, and lung, in which tumors are more vascular than the respective normal (3). In contrast to the majority of solid tumors, pituitary adenomas show a decreased expression of vascular endothelial growth factor and have signifi-

Table 1. Outcomes of Dynamic Enhancement Pattern, Thyroid Cancer

| Dynamic Enhancement Pattern                  | Patient Number | Percentage (%) |
|----------------------------------------------|----------------|----------------|
| Delayed enhancement                          | 4 papillary carcinoma | 75             |
|                                              | 1 medullary carcinoma |               |
|                                              | 1 FNAB proven atypical cell |      |
| Early strong enhancement                     | 1 follicular carcinoma | 12.5          |
| Persistently decreased enhancement           | 1 papillary carcinoma | 12.5          |
|                                              | 8               | 100            |

Note. — Compared to normal thyroid parenchyma.

FNAB = fine needle aspiration biopsy

Fig. 3. On ultrasonography, an approximately 5 millimeter, well hypoechoic nodule (arrow) is seen in the right lobe of the thyroid gland at mid level. This nodule has internal punctate tiny echogenicity, suggesting a tiny calcification. On a dynamic enhancement study, the nodule (arrow) shows a persistent decrease in enhancement compared to background parenchyma.

DISCUSSION

Although this study has the limitation of a small patient population, the data suggest that delayed enhancement on contrast enhanced dynamic T1WI is a possible characteristic MR findings of malignant thyroid tumor. The delayed enhancement pattern of thyroid cancer on dynamic T1WI is similar to that of pituitary adenoma. We hypothesize that the enhancement pattern of these tumors is related with tumor angiogenesis and its vascularity. Pituitary tumors are known to be less vascular than normal pituitary tissue, suggesting that angiogenesis may be inhibited in these tumors. This is consistent with the findings of Jugenburg and colleagues, but is in marked contrast to studies in other tissues including the breast, prostate, and lung, in which tumors are more vascular than the respective normal (3). In contrast to the majority of solid tumors, pituitary adenomas show a decreased expression of vascular endothelial growth factor and have signifi-
cantly lower microvessel densities compared to non-neoplastic pituitary glands (4). Similarly, the microvascular density was higher in normal tissue than in benign and malignant proliferative lesions of the thyroid gland (1). Microvessel density is a surrogate marker of tumoral angiogenesis. Microvessel density assessment is the most commonly used technique to quantify intratumoral angiogenesis in cancer. It was first developed by Weidner et al. (7) and used for panendothelial immunohistochemical staining of blood microvessels (8). In contrast to thyroid tumor and pituitary adenoma, hypervascular tumors such as breast and prostate cancer, shows early enhancement pattern on dynamic T1WI.

In addition, breast and prostate MR imaging is being widely used with increasing frequency to aid in the detection and evaluation of breast and prostate malignancies. We explained that, contrary to common cancers, but similar to other endocrine neoplasms such as pituitary and adrenal cortex tumours, angiogenesis is reduced in some thyroid proliferative lesions compared with normal tissue (1, 2). So, thyroid cancer shows a delayed enhancement pattern on contrast enhanced dynamic T1WI, probably due to its hypovascularity.

Two cases did not show a delayed enhancement pattern on dynamic T1WI. One case of follicular carcinoma showed strong early and persistent enhancement on contrast-enhanced dynamic T1WI, compared to normal parenchyma. This nodule is about a 3 cm sized lesion, the largest identified in our study. So, relatively well developed vascularity within the large cancer caused a different dynamic enhancement pattern. This patient had hashimoto thyroiditis, which could also explain the unknown underlying parenchymal change in the thyroid gland enhancement pattern. The other case of papillary carcinoma showed a persistently decreased enhancement pattern on dynamic T1WI, compared to normal parenchyma. The nodule was about a 5 mm sized lesion, relatively small compared to among the patients. Also this nodule has internal punctate tiny echogenicity, suggesting a microcalcification on USG. Hence, a relatively poorly developed vascularity within the small cancer caused a different dynamic enhancement pattern. Much to our regret, only one case of papillary carcinoma remained in the color Doppler USG. And, this malignant nodule shows hypovascularity compared to the parenchyma on USG and also showed a delayed enhancement pattern on the dynamic enhanced T1WI. But, because of the other 7 cases had not remained on the Doppler USG, it was difficult to perform a correlation analysis between Doppler USG and dynamic enhanced MR imaging in our study. We found that the maximum gland-lesion contrast difference peaked at 3 minutes to 4 minutes, 30 seconds after contrast infusion, suggesting that this is the most sensitive time for detecting lesions of the thyroid gland. Our dynamic technique was easy to perform and did not add a time penalty to the pituitary study. The practice of examining the gland twice, once during and after contrast infusion, was effective. In conclusion, delayed enhancement on contrast-enhanced dynamic T1WI is a possible characteristic MR finding of malignant thyroid tumors. Also, dynamic contrast-enhanced magnetic resonance imaging can be useful to detect or exclude thyroid carcinoma with high diagnostic accuracy in patients with a multinodular goiter when results of other diagnostic methods are inconclusive. The previous early results with MR of the thyroid indicate inadequate spatial and excellent contrast resolution for depicting disease in these glands (5, 6). The tool for the diagnosis of thyroid will be tempered by the traditional reliance on scintigraphy and sonography for this purpose. However, with the development of the MR technique, MR imaging has been shown to be a promising technique for imaging the thyroid. Moreover clear indications for MR in the evaluation of these glands are emerging. The limitations of our study was the relatively small patient population and the fact that we did not compare the benign and malignant thyroid lesions. Future studies should compare the MR imaging between benign and malignant nodules for the correct characterization of the tumor.

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악성 갑상선 종양의 특징적인 역동적 자기공명영상의 조영증강 소견: Preliminary Report

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목적: 이 논문의 목적은 자기공명영상 검사에서 악성 갑상선 종양이 어떤 역동 조영증강 소견을 보이는지 확인해 보는 것이다.

대상과 방법: 수술 전, 병리 조직 검사에서 악성 갑상선 종양으로 확인된 8명의 환자를 대상으로 하였으며, 각각 5명의 유두암종(papillary carcinoma), 1명의 수질암종(medullary carcinoma), 1명의 소포암종(follicular carcinoma), 1명의 세침흡입검사에서 확인된 비정형세포(atypical cell)의 환자로 구성되었다. 모든 자기공명영상 검사는 3.0-T magnetic resonance (MR) imaging unit, Magnetom Verio (Siemens Medical Systems, Erlangen, Germany)로 얻어졌으며, MRI protocol은 axial fast spin-echo T2-weighted MR image, axial T1-weighted image and contrast agent-enhanced dynamic T1-weighted images의 pulse sequence를 포함하였다. 조영제는 paramagnetic contrast agent (gadobutrol, Gadovist: Schering, Berlin, Germany)을 이용하였다. 그 외 병리 조직 검사결과와 초음파 검사결과는 후향적으로 확인하였다. 갑상선 자기공명영상 검사를 한 모든 환자에게 설명 후 동의를 받았으며 병원 윤리위원회에서 갑상선 자기공명영상 검사에 대한 허가를 받았다.

결과: 수술 전 자기공명영상 검사를 시행하였고, 각각 종양의 조직형에 따른 역동 조영증강 소견을 비교해 보았다.

1) T1 강조영상에서 1명의 유두암종 환자는 고신호 강도 종괴로 보였고, 1명의 수질암종 환자는 저신호 강도 종괴로 보였으며, 그 외 6명의 경우는 갑상선 실질과 구분되어 보이지 않았다.

2) T2 강조영상에서 3명의 유두암종 환자와 1명의 소포암종 환자만 고신호 강도 종괴로 확인이 되었으며 1명의 유두암종 환자와 척추신경의 고신호 강도 종괴로 확인이 되었으며 1명의 유두암종 환자는 저신호 강도 종괴로 확인이 되었으며 1명의 유두암종 환자는 저신호 강도 종괴로 확인이 되었다. 그 외 3명의 종괴는 갑상선 실질과 구분되어 보이지 않았다.

3) 역동적 조영증강 T1 강조영상에서는, 5명의 유두암종 환자와 1명의 수질암종 환자와의 조영증강 소견이 정상 실질과 비교하여 차이가 있는 소견을 보였다. 1명의 소포암종 환자의 경우 종괴는 정상 실질과 비교하였을 때 강한 초기 조영증강 양상을 보였으며, 1명의 유두암종 환자와의 조영증강 소견은 저속적으로 정상 실질보다 감소된 조영증강 양상을 보였다.

결론: 위 연구는 8명이라는 적은 환자군을 가지고 악성 갑상선 종양이 역동 조영증강 양상을 연구한다면 한계점이 있다. 하지만 위의 연구 결과로 미루어 볼 때 역동적 자기공명영상의 조영증강 검사에서 지연 조영증강을 나타낸 악성 갑상선 종양의 하나의 특징적인 소견일 수 있다는 것을 제시해 준다. 앞으로 양성 갑상선 종괴의 자기공명영상 양상에 대한 비교 연구가 좀 더 필요할 것으로 생각된다.

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