EVALUATION OF PAROTID GLAND TUMORS WITH MULTI-PHASE DYNAMIC HELICAL CT

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INTRODUCTION

The surgical procedure chosen for salivary gland tumors is strongly influenced by their location and histologic type. Benign tumors are treated by local excision or superficial parotidectomy, whereas a more aggressive resection is chosen for malignant tumors.1-3 Computed tomography (CT) and magnetic resonance imaging (MRI) are both commonly used in the preoperative evaluation of salivary gland masses. MRI findings of parotid tumors have previously been described and showed a good correlation between dynamic contrast enhancement pattern and histopathologic findings.5-6 On CT, although an irregular tumor margin and infiltration suggest malignancy, there is considerable overlap among malignant and benign masses. There are few reports indicating the use of a two-phase dynamic helical CT in their characterization.7 Thus the purpose of our study was to perform multi-phase dynamic helical CT in patients with parotid gland tumors and describe their enhancement characteristics.

MATERIALS AND METHODS

Nineteen patients (ten male, nine female patients; age range, 32-66 years; mean age, 48 years) with a parotid mass, referred for presurgical evaluation between March 2000 and November 2002, were included in this study. The histopathologic diagnosis was confirmed after surgical resection in all patients. Written informed consent was obtained from each patient for this study.

All examinations were performed with a helical CT scanner (HiSpeed Advantage; GE Medical Systems, Milwaukee, WI, USA) capable of a 1 s gantry rotation time. A total of 90 mL of contrast agent was administered into an antecubital vein at a rate of 3 mL/s using a power injector. Multi-phase axial CT images were obtained before and 30, 90, 180, and 360 s after the contrast material injection. A late enhanced axial image was also obtained after a delay of 30 s, the scanning began at the skull base and continued toward the thoracic inlet. For the non-enhanced and delayed-enhanced images the scanning range was limited to the parotid gland. Helical images were obtained with a 5 mm collimation 5 mm/s table speed, 120 kVp, and 80 mAs. From the volumetric data, contiguous transverse images were reconstructed at 5-mm intervals.

The images were transferred to a workstation and reviewed by two radiologists at one sitting. The radiologists evaluated image sets of a patient side by side at the same time and reached a consensus. The degree of lesion enhancement was graded as no enhancement, trace enhancement, moderate enhancement, or intense enhancement. The lesion enhancement pattern was visually assessed at each phase and classified as heterogeneous, hetero-
geneous enhancement with intratumoral low attenuation, or homogeneous. For quantitative assessment, density numbers in Hounsfield units (HU) of the lesions were recorded at each phase by means of the largest possible circular region of interest (ROI). When choosing the ROI, care was taken to exclude any cystic or necrotic areas. Density numbers measured in the early (30 s delay) and delayed (25 min delay) phase were compared. A plot representing the time-attenuation curve was constructed for each lesion. The tumoral enhancement ratio (ER) was calculated by using the function $ER = \frac{\text{Subscript}}{\text{CTlate}}$, where Subscript is the density number measured at 30 s delay, and CTlate is the density number measured at 25 min delay in Hounsfield units.

**RESULTS**

There were twelve pleomorphic adenomas, five malignant tumors and two Warthin tumors. The size of lesions ranged between 14 and 52 mm (mean, 28 mm). In the visual assessment five pleomorphic adenomas, one Warthin tumor, and two malignant tumors showed homogeneous enhancement in the early phase. Heterogeneous enhancement with intratumoral low attenuation was seen in two pleomorphic adenomas, one Warthin tumor, and three malignant tumors. Five pleomorphic adenomas showed heterogeneous enhancement.

In the quantitative assessment, tumoral mean density numbers in early phase scans were highest for the Warthin tumor (105±16 HU), followed by malignant tumors (83±19 HU), and pleomorphic adenomas (69±14 HU). In delayed phase scans, tumoral mean density numbers were highest for pleomorphic adenomas, followed by malignant tumors, and the Warthin tumor (Table 1). The resultant ER was highest for the Warthin tumor. The number of patients included for each category was too few to statistically evaluate the relationship between the density numbers and the pathological diagnosis. The results are summarized in table 1.

**DISCUSSION**

Preoperative differentiation between benign and malignant parotid tumors is important for adequate surgical planning and for the tumor prognosis. Aspiration cytology performed prior to surgery is often unreliable and cannot distinguish between benign and malignant tumors. CT imaging is widely used to assess the size and extent of the tumor, but it is less effective in distinguishing between benign and malignant tumors.

Figure 1a: Axial CT images acquired with a delay of 30 seconds (a) and 25 minutes (b) following contrast material injection of a 56 year-old man with Warthin tumor. The tumor shows a heterogeneous pattern of enhancement at early phase. Time-attenuation curve (c) for the same lesion depicts an early peak of enhancement with a high washout.

Figure 1b: Axial CT images acquired with a delay of 30 seconds (a) and 25 minutes (b) following contrast material injection of a 48 year-old woman with malignant tumor of the left parotid gland. The tumor shows a heterogeneous enhancement with intratumoral low attenuation at the early phase. Time-attenuation curve (c) for the same lesion depict an rapid enhancement with a slower washout.

Figure 1c: Axial CT images acquired with a delay of 30 seconds (a) and 25 minutes (b) following contrast material injection of a 45 year-old man with pleomorphic adenoma. The tumor shows a homogeneous pattern of enhancement at early phase. Time-attenuation curve (c) for the same lesion depict an moderate enhancement with a slow washout.
Table 1: Enhancement ratio, and early and delayed phase density numbers of parotid gland tumors on multi-phase CT scan.

<table>
<thead>
<tr>
<th>Tumor Type</th>
<th>Early Phase Density Number (HU)</th>
<th>Delayed Phase Density Number (HU)</th>
<th>ER</th>
</tr>
</thead>
<tbody>
<tr>
<td>Warthin Tumor</td>
<td>105±16</td>
<td>53±17</td>
<td>1.98</td>
</tr>
<tr>
<td>Malignant Tumor</td>
<td>83±19</td>
<td>58±15</td>
<td>1.43</td>
</tr>
<tr>
<td>Pleomorphic Adenoma</td>
<td>69±14</td>
<td>66±13</td>
<td>1.05</td>
</tr>
</tbody>
</table>

Note: Data are mean±standard deviation. HU: Hounsfield Units ER: Enhancement Ratio

In our study, the use of a multiple phase imaging protocol including late imaging 25 min after the contrast injection enabled us to determine time-attenuation curves and to accurately assess delayed tumoral enhancement. Warthin tumors showed an early and strong enhancement, whereas both malignant tumors and pleomorphic adenomas exhibited a slower enhancement pattern. Density numbers measured on delayed phase images were higher for pleomorphic adenomas. This finding is consistent with those obtained using dynamic MRI, which reported a peak signal intensity at 30 s following contrast injection for Warthin tumors and a gradual increase in signal intensity for up to 5 min for pleomorphic adenomas. The time-attenuation curves obtained in our study were consistent with the time intensity curves of previous dynamic contrast enhanced MRI studies.

The tumoral enhancement pattern is greatly influenced by tumor vascularity, histopathologic cell type, and the histologic stromal component. Warthin tumors consist of an epithelial, mostly oncocytic component with a lymphoid stroma, and have a great tendency to undergo cystic change. Pleomorphic adenomas on the other hand contain mixed epithelial and stromal elements. The stroma can show mucoid, chondroid, fascicular, and hyalin-fibrous differentiation. Malignant tumors can also have cystic components or show necrosis. This histological diversity of parotid gland tumors is well reflected in the visual assessment findings of the above presented study. It has also been hypothesized that early tumoral enhancement is related to tumoral microvasculature, whereas delayed enhancement reflects the cellularity and stromal component of the tumor. Warthin tumors are known to have rich microvasculature, which helps explain their early and intense enhancement pattern. Because Warthin tumors tend to undergo cystic changes, they have less cellularity and a smaller stromal component, and therefore fail to retain contrast material on delayed phase images. On the other hand, malignant tumors, in addition to their various amounts of microvasculature and possible cystic necrotic foci, contain a larger stroma, which in turn retains contrast media in the late phases. Although both pleomorphic adenomas and malignant tumors show cystic changes, the former have more abundant fibrous stroma, which accounts for their higher late phase enhancement.

A main limitation of our study is the small number of patients included. Measured ERs were higher for Warthin tumors, followed by malignant tumors and pleomorphic adenomas. These findings correlate well with previous reports, but the numbers of Warthin and malignant tumors included in this study were limited.
study are too small for statistical analysis or to allow us set a value for lesion characterization. Therefore, further studies with larger number of patients including different types of parotid tumors are needed.

In conclusion, multi-phase helical CT showed a pattern of strong enhancement in early phase scanning with a decrease in the delayed phase for Warthin tumors. Although all non-Warthin parotid tumors showed a slower enhancement pattern, an increase in enhancement in delayed phase images might be suggestive of a pleomorphic adenoma. These enhancement patterns can be helpful in the differential diagnosis of parotid gland tumors.

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REFERENCES