Chondrosarcoma of the Head and Neck

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Chondrosarcoma of the head and neck region is a rare disease, representing approximately 0.1% of all head and neck neoplasms.¹ This malignant tumor, in which the tumor cells form chondroid matrix, represents 10-20% of primary bone tumors and is the second most common sarcoma of bone origin.²,³ Due to the low incidence of this disease, there have been few studies with large numbers of subjects, and therefore, various results have been reported.⁴⁻⁶ In this study, we reviewed cases of chondrosarcoma experienced at Yonsei University Medical College during the last 15 years to determine the clinical manifestations, treatment methods and treatment results of this uncommon disease.

MATERIALS AND METHODS

Using the cancer registration for reference, patients who were admitted to our institute between 1990 and 2003, and whose final diagnosis at discharge was chondrosarcoma, were reviewed. There were 100 cases of chondrosarcoma, and of those, the head and neck region was the primary site in only 8 cases. The charts were reviewed retrospectively in terms of chief complaint, primary site, treatment modality, pathology finding, recurrence, follow-up and treatment results.

RESULTS

2010 patients were diagnosed with head and neck neoplasms at our institute from 1990 to 2003 and of these, there were only 8 cases of chon-
Surgical Treatment of Head and Neck Chondrosarcoma

Table 1. Clinical Characteristics of Head and Neck Chondrosarcoma

| Case | Age | Sex | CC       | Site                | Treatment                        | Pathology            | Grade | Margin | Recur | F/U (month) | Outcome |
|------|-----|-----|----------|--------------------|----------------------------------|----------------------|-------|--------|-------|-------------|---------|
| 1    | 34  | M   | Epiphora | Ethmoid sinus      | Craniofacial resection & postop. RTx | Chondrosarcoma       | II    | +      | No    | 111          | NED     |
| 2    | 56  | F   | Visual loss | Ethmoid sinus | Medial maxillectomy & ethmoidectomy & postop. RTx | Chondrosarcoma       | III   | +      | No    | 30           | NED     |
| 3    | 29  | F   | Facial swelling | Maxillary sinus | Subtotal maxillectomy | Mesenchymal chondrosarcoma | -    | No     | 25    | NED          |
| 4    | 26  | M   | Facial swelling | Maxillary sinus | Partial maxillectomy | Chondrosarcoma       | II    | -      | Yes   | 171          | NED     |
| 5    | 24  | F   | Facial palsy | Mastoid bone       | Mass excision via transmastoid approach | Chondrosarcoma       | II    | -      | Yes   | 40           | NED     |
| 6    | 49  | F   | Facial palsy | Mastoid bone       | ITFA type C | Chondrosarcoma       | I     | -      | No    | 51           | NED     |
| 7    | 42  | M   | Facial swelling | Jugular foramen | ITFA type A | Chondrosarcoma       | I     | -      | No    | 130          | NED     |
| 8    | 60  | M   | Neck mass | Thyroid cartilage | Mass excision | Chondrosarcoma       | II    | -      | Yes   | 54           | DOD     |

CC, chief complaint; F/U, follow-up; NED, no evidence of disease; DOD, died of disease; ITFA, infratemporal fossa approach.

Table 2. Chondrosarcoma of the Head and Neck by Site

| Site                  | Number |
|-----------------------|--------|
| Paranasal sinus       | 4      |
| Temporal bone         | 3      |
| Larynx                | 1      |
| Total                 | 8      |

drosarcoma, which represents 0.4% of all head and neck cancers. Of the 8 cases, 4 were male and 4 were female, with a mean age of 40 years ranging from 24 to 60. The follow-up period ranged from 25 months to 171 months with a mean of 76.5 months (Table 1). The primary sites were paranasal sinuses, temporal bone and larynx (Table 2). The chief complaints at the time of first visit were related to the primary sites, such that an ethmoid sinus origin tumor presented with ocular symptoms, maxillary and jugular foramen origin tumors with facial swelling, a mastoid origin tumor with facial paralysis, and a thyroid cartilage origin tumor with neck mass. Pathologically, of the 8 cases of head and neck chondrosarcoma, 7 were conventional chondrosarcoma and 1 was mesenchymal chondrosarcoma. Of the 7 cases of conventional chondrosarcoma, 2 were grade I, 4 were grade II and 1 was grade III (Fig. 1 and 2). Surgery was performed as the initial treatment modality in all cases. Radiotherapy was applied in 4 cases, which included 2 cases of recurrence and 2 cases with positive resection margin. The 2 cases with positive resection margin were successfully treated after radiotherapy. Recurrence occurred in 3 cases. The pathological grade of all 3 cases was grade II and salvage treatment was performed in every case. Of the recurring cases, the case of mastoid origin tumor was successfully treated surgically with a transmastoid approach, whereas in the case of maxillary sinus origin, successful salvage was achieved 7 years from the point of diagnosis after receiving chemoradiation therapy and bilateral maxillectomy. However, the case of thyroid cartilage origin recurred in the tonsil 13 months...
after the initial treatment. Despite treatment with radiotherapy and two salvage operations, death occurred 54 months after diagnosis because of failure to completely excise the tumor at the last operation due to anatomical reasons. Seven of eight cases have no current evidence of disease, and the survival rate is 87.5% (Table 1).

**DISCUSSION**

Chondrosarcoma is a slow growing but malignant tumor with a relatively high local recurrence rate. Chondrosarcoma represents 10 to 20% of all malignant bone tumors and of these, 1 to 12% originate in the head and neck region. This tumor can originate from cartilage and soft tissue as well as bone with common primary sites being the pelvic bone and long bones, such as the femur. The common primary sites in the head and neck region include the mandible, nasal cavity, sinus, and maxilla; however, it is generally accepted that the tumor mostly arises from bony tissue of the head and neck. In this study, the 8 cases were all of bone origin except for one case that originated from thyroid cartilage. The precipitating factors of this tumor are multiple hereditary exostosis, Ollier's disease, Maffucci's syndrome, previous intravenous thorotrast contrast use, Paget's disease of bone, chondromyxoid fibroma, and previous irradiation, but none of these factors were found in this study. Chondrosarcoma has a slight male predilection and occurs most commonly between the 4th and 7th decades. Chondrosarcomas of cartilaginous or soft tissue origin are more common in males and in patients over 50 years of age, whereas those of bone origin have higher incidence in females and in patients less than 50 years of age. According to one report, the rates of lymph node and distant metastasis were relatively low: 5.6% and 6.7%, respectively. In addition, metastasis was usually observed in patients who had undergone multiple operations over a long period of time. In this study, no case of metastasis was seen among the 8 cases, including the 3 patients who received multiple salvage operations.

Chondrosarcomas show various histological patterns ranging from benign chondroid tumor to undifferentiated neoplasm, which make them difficult to diagnose pathologically. Evans et al. classified chondrosarcomas into 3 grades, from grade I to grade III, according to cellular density, nuclear differentiation, and the size of nucleus. This classification is still currently used. Nevertheless, there have been efforts to classify chondrosarcomas simply into high grade and low grade for better correlation with prognosis. In this study, no recurrence occurred in grade I tumors, showing a relatively good correlation between tumor grade and prognosis. Chondrosarcoma has several histological types, of which the conventional type is the most common. Other types include clear cell, myxoid, mesenchymal and dedifferentiated variants. The clear cell variant is otherwise called malignant chondro-
blastoma, but there has been controversy over whether to classify it as a variant of chondrosarcoma or to separately categorize it as malignant chondroblastoma. Although there are differences exist among reports, the myxoid variant is known to originate from soft tissues rather than bone tissues. The mesenchymal variant is also called the aggressive variant, since 2/3 of cases present before the age of 30, and in advanced stages and grades. The only patient with mesenchymal chondrosarcoma in this study was a young female aged in the 3rd decade. Dedifferentiated chondrosarcoma characteristically contains an anaplastic component histologically. In addition, chondrosarcomas of the larynx have several characteristics that distinguish them from those of different origin: they are low grade tumors, presentation occurs after 50 years of age, and they have a better prognosis compared to bone origin chondrosarcomas of the head and neck region.

Surgical treatment is known as the most effective treatment modality for chondrosarcoma. The most important point in performing surgery is to ensure an adequate safety margin histologically since residual disease is known to be an important cause of recurrence. In general, the principle technique for extralaryngeal chondrosarcoma is a wide bloc resection, whereas that for laryngeal chondrosarcoma is conservative resection. The reason for this is that laryngeal chondrosarcoma is usually of low grade, showing a relatively favorable prognosis so that mass excision without total laryngectomy is sufficient to achieve an acceptable survival through salvage surgery. Neck dissection is not routinely performed because of a low incidence of lymph node metastasis. Concerning the radiosensitivity of chondrosarcoma, diverse opinions have been reported, ranging from radioresistant to radiosensitive and even curative. However, it is generally accepted that radiotherapy should be used for palliative purpose in unresectable cases, or as an adjuvant therapy in cases of residual disease rather than as an initial treatment. Chemo-therapy has a limited role in chondrosarcoma, but can be applied as an adjuvant therapy in high grade mesenchymal chondrosarcomas, in cases of rapid local recurrence with aggressive behavior, or in cases with potential for metastasis. In this study, all 8 cases were initially treated with surgery and of these, 4 cases were treated successfully with surgery alone. The 2 patients who had residual disease after initial treatment received radiotherapy and are disease free at the present day, suggesting that radiotherapy does have some role in controlling the disease.

The survival rate of chondrosarcoma is reported as 44-87%, and there are two possible explanations for this wide range. One is that it is hard to perform large-scale studies due to the rarity of this disease. The other is that the survival rate has improved a great deal recently thanks to the advancement in radiographic imaging and surgical technique. The survival rate in this study was 87.5%, which is relatively high compared with other reports. The prognostic factors of chondrosarcoma are resectability, stage, grade and primary site, while myxoid and mesenchymal variants are known for their poor prognosis. The most important prognostic factor is resectability, which makes complete excision of the tumor the single most significant factor in determining the prognosis. This is also supported by the fact that the most common cause of death in chondrosarcoma is local recurrence, not metastasis. The cause of death of the case in this study was residual disease that remained because of an inability to ensure an adequate safety margin at the last operation, despite multiple previous operations. In cases of recurrence, chondrosarcoma grows progressively for more than 2 years from recurrence to death. Aggressive surgical treatment should be considered, even for recurrent tumors, since the usual pattern of recurrence is local failure and not metastasis. In this study, all 3 cases of recurrence received aggressive treatment including surgery, and 2 of the 3 cases were successfully salvaged. One of the recurrent cases was salvaged 7 years after receiving multiple treatments due to multiple recurrences. There was no evidence of metastasis during the 7 years, proved the slow-growing nature of the disease with low incidence of metastasis. Moreover, this example demonstrates the importance of local control during treatment.

Chondrosarcoma of the head and neck region is such an uncommon disease that we only experienced 8 cases during a 15-year period. This dis-
ease has a slow-growing nature with low incidence of metastasis. Therefore, we conclude that complete excision of the primary site is of utmost importance for successful treatment. Although the number of cases was small, and despite considerable controversy, we speculate that postoperative radiotherapy may have some role in the treatment of chondrosarcoma, considering the 2 cases reported here with positive margin that were successfully treated with postoperative radiotherapy.

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