MICROSURGICAL MANAGEMENT OF LOW-GRADE SPINAL CORD ASTROCYTOMA IN ADULTS: A PERSONAL CASE SERIES REPORT AND BRIEF LITERATURE REVIEW

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SUMMARY – Astrocytoma is the second most common intramedullary tumor of predominantly low-grade malignancy in adult patients. Adult astrocytomas have better-quality prognosis compared with astrocytomas in children. Although a standardized surgical management protocol for spinal cord glioma is currently unavailable, surgery of low-grade astrocytoma should be aimed at gross total resection to preserve neurological function and to improve the outcome. Herein, we present a personal case series of four consecutive adult spinal cord astrocytoma patients who were operated on during the last few years. Tumor resection was performed in all patients utilizing microsurgical technique and intraoperative neurophysiologic monitoring. We also provide a literature review of the treatment of intramedullary astrocytoma in adults and discuss contemporary surgical management and prognosis.

Key words: Intramedullary tumor; Low-grade astrocytoma; Microsurgical management

Introduction

Spinal cord astrocytoma is an intradural intramedullary glial cell tumor accounting for 6%-8% of all primary spinal cord growths1-5. It is the most common intramedullary pediatric tumor and occurs secondary to ependymoma in adult patients. Adult astrocytomas have better-quality prognosis compared with astrocytomas in children. Although classified as histologically malignant according to the 2016 World Health Organization (WHO) tumor classification6, these tumors are predominantly of low-grade malignancy, including diffuse and pilocytic astrocytoma1-5. Since the average age at presentation in adults is 29±18 years7, it can be said that it predominantly affects younger patients, but steady age distribution from early childhood to the advanced age is not rare7. In general, low-grade astrocytoma has longer disease history with a range of various symptoms that are reliant on tumor volume and location, growth pattern, and histology8.

Notwithstanding their low-grade histologic nature, these tumors can be well enhanced on contrast magnetic resonance imaging (MRI) of the spine2, which improves distinction between more common solid and less common cystic types of the tumor, which are usually more malignant4 but can be better gross totally resected due to their less infiltrative growth pattern2. Nevertheless, a standardized surgical management
protocol for spinal cord glioma is currently unavailable\textsuperscript{4,10,11}, although the treatment of low-grade tumors should be aimed at radical, gross total microsurgical resection to improve and preserve neurological function\textsuperscript{2,8,12}.

Herein, we present a personal surgical case series of four spinal cord astrocytoma adult patients, and describe contemporary microsurgical techniques that are intended to assure complete tumor resection. We also provide a literature review and discuss the extent of resection, neurological outcome, and prognosis.

Case Series Report

During the last few years, four consecutive adult spinal cord astrocytoma patients were operated on by a senior neurosurgeon (K.I.A.). This personal series of surgical patients included three females and one male, mean age 55.5 years at the time of surgery. Immediate postoperative recovery was favorable in all patients, whose neurological and functional status much improved following surgery.

Case 1

A 61-year-old female presented with type 2 diabetes, diabetic polyneuropathy, and gait disturbances with minor motor weakness in her lower limbs (light paraparesis). She was previously diagnosed with an intramedullary tumor mass at the T1 level and was clinically and radiologically followed-up for several years. Her preoperative cervical and thoracic spine contrast-enhanced magnetic resonance imaging (MRI) at 8 years, 2 years, and immediately prior to surgery are depicted in Figures 1 and 2.

Due to worsening of neurological deficit related to tumor progression, she was eventually operated on.

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{fig1.png}
\caption{Sagittal T2-weighted magnetic resonance imaging (MRI) of the thoracic spine (A) showing the hyperintense intramedullary lesion at T1 (arrow); axial T2-weighted MRI of the thoracic spine showing lesion at the same level (arrow) (B); sagittal (C) and axial (D) T2-weighted MRI of the thoracic spine at 6-year follow-up revealing lesion growth (arrow).}
\end{figure}
The tumor was completely resected and histopathologic analysis of tissue samples confirmed the tumor as a low-grade astrocytoma (WHO grade II). Postoperative MRI of the cervicothoracic spine showed complete tumor resection (Fig. 2A-H).

**Case 2**

A 45-year-old female presented with lower back pain, decreased motor strength of lower limbs, and perineal numbness. Sagittal and axial MRI reformations of the entire spine showed an intramedullary tumor growth at the T12-L1 levels (Fig. 3A-D). The tumor was resected using a meticulous microsurgical technique and intraoperative neurophysiological monitoring to record motor evoked potentials (MEP). The immediate postoperative imaging of the lumbar and thoracic spine are depicted in Figure 4(A-C). Histopathologic diagnosis defined the tumor as anaplastic

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**Fig. 2.** Preoperative sagittal T2-weighted (A), sagittal T1-weighted post-contrast (B), axial T2-weighted (C), and post-contrast axial T1-weighted (D) magnetic resonance imaging (MRI) of the cervicothoracic spine showing an intramedullary tumor (arrows); postoperative sagittal T2-weighted (E), sagittal T1-weighted post-contrast (F), axial T2-weighted (G), and axial T1-weighted post-contrast (H) MRI of the cervicothoracic spine demonstrating complete tumor resection (arrows).

**Fig. 3.** Sagittal T1-weighted post-contrast magnetic resonance imaging (MRI) of the lumbar spine showing well-demarcated homogeneously enhanced intramedullary lesion (arrow) (A); sagittal T2–weighted (B), sagittal T2–weighted (C), and axial post-contrast T1–weighted MRI (D) of the lumbar and thoracic spine depicting intramedullary tumor at T12 level (arrows).
astrocytoma (WHO grade III). Postoperative focal irradiation of the tumor field was applied.

Case 3

A 58-year-old male presented with lower limb motor weakness and sensory loss at the T1 level. Whole spine MRI identified an intramedullary tumor growth at the T1-2 level (Fig. 5A-C), which was successfully resected by microsurgical technique with the help of intraoperative neurophysiological monitoring (Fig. 5D-F). Histopathologic diagnosis confirmed it as low-grade astrocytoma (WHO grade II).

Case 4

A 58-year-old female presented with lower limb motor weakness (paraparesis). Preoperative contrast-enhancing spinal MRI showed an intramedullary tumor at the T4 level (Fig. 6A-D). Gross total tumor microsurgical resection was achieved, which was confirmed by postoperative MRI of the thoracic spine (Fig. 7A-C). Histopathologic diagnosis identified the tumor as low-grade spinal cord astrocytoma (WHO grade II).

Discussion

After reviewing the literature, we found the mean patient age at presentation to range from 28.3 to 41±17 years. Seven,12,13 Five-year survival of patients with low-grade tumors (WHO grades I and II) ranged from 54% to 78% with median survival of 8-13 years. A detailed literature review of spinal cord astrocytomas has been published recently by our group.8

Preoperative neuroradiological evaluation includes MRI with and without contrast of the selected areas of the spine. Enhancement of spinal cord astrocytomas appears to vary and may include cystic formations within. Anteroposterior/lateral and flexion and extension x-rays are important to assess the potential spinal instability and the need of instrumentation.

Laminectomy with the patient in prone position is the most common surgical approach used to expose the posterior/dorsal part of the vertebral canal, spinal cord, and intramedullary lesions. Laminoplasty can also be utilized, especially in younger patients and in patients with multiple levels of involvement. Instrumented fusion can be considered especially in cervicothoracic and thoracolumbar junctions. Dura opening, arachnoid dissection, and myelotomy are usually performed dorsally through a midline incision via the posterior midline septum. The aim of surgery is to separate tumor margins meticulously from the surrounding cord tissue, which is not always easy to achieve when infiltrating tumors of more malignant grades are encountered.6,14 Yet, sharp dissection and creation of clear tumor planes is necessary to better expose the tumor, but may be challenging, which calls for high surgical skills.

We follow these rules for microsurgical technique and employed the same operative technique in the patients from this series.6,14 Recently, we have published a video clip of the surgical removal of spinal cord astrocytoma, which is available with free access.14
Recent analyses have shown that intraoperative neurophysiological monitoring and computed tomography neuro-navigation with advanced multi-modal options of microscope-based augmented reality, as well as other up-to-date, cutting-edge microsurgical armamentarium may be very helpful in providing accurate
tumor visualization and in ensuring high-precision dissection when intradural spinal growths are concerned\textsuperscript{15,16}. In spite of that, literature data on gross tumor resection rates for low-grade spinal cord glioma range between 5% and 72.7% of cases\textsuperscript{13,17-20}, although population-based studies confirmed improved outcomes and overall survival in these patients\textsuperscript{21-25}, especially those of younger age\textsuperscript{26,27}. However, preoperative and postoperative neurological condition, extent of resection, and low histological grade remain the strongest outcome predictors\textsuperscript{3,4,8,10,22,28-32}.

Gross total tumor resection was achieved in all four patients in our series. Their immediate postoperative recovery was favorable, which allowed for marked improvement in neurological functioning in all of them. Nonetheless, having in mind that a significant percentage of low-grade astrocytomas are more likely to be removed completely, it seems that the extent of resection alone may not be the only predictive outcome parameter\textsuperscript{4,8}.

Since the postoperative recurrence rate of low-grade astrocytoma is usually low, it appears that adjuvant therapy is not obligatory in all cases. Accordingly, some authors recommend radiotherapy for patients with high-grade tumors only\textsuperscript{7}. We have opted for the same strategy in this series of patients, bearing in mind that the main factor influencing prognosis and outcome of these, as well as other primary intramedullary spinal cord tumors is not the mode of treatment, but tumor grading, preoperative and postoperative neurological condition, and the extent of resection\textsuperscript{26,33,34}.

In conclusion, we believe that surgery of adult low-grade spinal cord astrocytoma should be aimed at gross total tumor resection in cautiously selected patients, which could result in a favorable outcome. Simultaneously, cutting-edge microsurgical equipment and technique should be engaged to achieve the goals of surgery. Neurophysiological monitoring during surgery is of utmost importance.

Nonetheless, continuous supplementary research in a greater number of patients is required in order to endorse these suggestions further.

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ASTROCITOM KRALJEŽNIČKE MOŽDINE NISKOG STUPNJA MALIGNOSTI U ODRASLIH: PRIKAZ OSOBNE SERIJE BOLESNIKA I PREGLED LITERATURE

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Astrocitom kralježničke moždine je tumor pretežito niskog stupnja malignosti koji je po učestalosti drugi intramedularni tumor u odraslih u kojih je njegova prognoza znatno povoljnija u usporedbi s prognozom u djece. Unatoč tomu što standardizirani protokol za kirurško liječenje gliomskih tumora kralježničke moždine zasad ne postoji, cilj ovakvog liječenja trebao bi biti usmjeren ka uklanjanju tumora u cijelosti kako bi se očuvala neurološka funkcionalnost i poboljšala uspješnost liječenja. U ovom radu dajemo prikaz osobne serije tijekom nekoliko posljednjih godina susljedno operiranih četvero odraslih bolesnika s astrocitomom kralježničke moždine u kojih je tumor mikrokirurški uklonjen uz pomoć intraoperacijskog neurofiziološkog praćenja. U radu također donosimo pregled literature o liječenju intramedularnih astrocitoma u odraslih, raspravljajući o suvremenom kirurškom liječenju ovakvih tumora i prognozi bolesti.

Ključne riječi: Intramedularni tumor; Astrocitom, niski stupanj malignosti; Mikrokirurško liječenje