Surgical Management of Spinal Meningioma

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Abstract

Background: Meningioma is a common tumor that accounts for 25-46% of spinal neoplasms. It is generally benign, and slow growing. Spinal meningiomas occur after the fourth decade of life. Mostly, they are located in the intradural compartment, generally respecting the pial layer of the spinal cord. The most frequent site of spinal meningiomas in the thoracic region. Clinical findings vary from mild to significant neurologic dysfunction; the most frequent clinical findings are back pain, sensori-motor deficit and sphincter dysfunction. The goal of surgical treatment must be total resection, if possible and usually not difficult.

Objective: This study was made to present the incidence, clinical presentation, localization, techniques, used in the management of spinal meningioma. And to evaluate the functional outcome of surgically treated cases.

Patients and Methods: This is a retrospective study of Sixteen patients with spinal meningiomas who had undergone microsurgical resection were treated between 2012 and 2015 in our department. Clinical presentation, diagnosis, histological examination, microsurgical resection, functional outcome were evaluated, defining potential prognosis factors associated with these lesions.

Results: Tumors site was intradural, extramedullary the most common site was the thoracic region, posterolateral and antero-lateral. Surgical resection was complete in 14 patients (87.5%) and incomplete in 2 patients (12.5%). At the last follow-up the neurological state was improved or unchanged in 15 patients (93.75%) and worse in one patients (6.25%).

Conclusion: Magnetic resonance imaging is the best imaging technique for diagnosis. Total tumor resection improved the surgical results of spinal meningiomas.

Key Words: Extramedullary – Intradural – Meningioma – Spinal surgery – Spinal tumors.

Introduction

MENINGIOMAS involving the spinal cord are relatively rare in comparison to the intracranial compartment accounting approximately 1.2% of all meningiomas of the central nervous system [1,2]. Mostly, they are located in the intradural compartment, generally respecting the pial layer of the spinal cord. Isolated extradural spinal meningiomas are rare [3]. By the introduction of modern neuroimaging techniques and standard microneurosurgical procedures, these tumours nowadays can be removed with a low morbidity and mortality [4]. Spinal meningiomas occur after the fourth decade of life, over 70% of the patients are between the ages of 40 and 70 years with a mean age of 50 years [1,5], thus, the occurrence in younger patients should raise the suspicion of a genetically determined disorder as neurofibromatosis 2 or an association with aggressive histological subtype [6,7]. They are slow growing tumours and therefore, they lead to symptoms due to significant spinal cord compression. Local pain is one of the leading symptoms, however, in a considerable number of patients diagnosis is confirmed not until neurologic deficits or gait disturbances are present [3,8,9].

Patients and Methods

Between January 2012 and December 2015 a total of 16 patients with a spinal meningioma underwent micro neurosurgical resection. Exclusion criteria included. (1) Meningioma en plaque or totally calcified tumors, (2) Foramen magnum meningioma as they have intra cranial compartment. (3) Patient with complete hemiplegia, (4) Recurrent cases The female gender was clearly predominant with 14 female patients and 2 male patients (female/male ratio 7:1) All these patients were examined preoperatively including: Age-the mean age was 69 years (range 17-88 years), the mean duration of symptoms was 13.7 months, scoring of motor weakness and sensory deficits, was done. All patients were followed-up clinically and by spinal MRI. The mean follow-up period was 12 months Histological examination of the specimen confirmed a grade I meningioma accord-
Surgical Management of Spinal Meningioma

Contrast enhanced spinal MRI was the diagnostic tool of choice and performed routinely for preoperative evaluation in all patients. To determine the spinal level, size and the dural attachment of the meningioma and its relation to the spinal cord. All meningiomas were operated in the prone position using a microsurgical technique via a midline skin incision extending two levels above and below the extent of the lesion. A monosegmental or multisegmental laminectomy was performed above and below the extent of the tumor removal of the meningioma was performed under standard microsurgical conditions, when necessary, a surgical aspirator (Cooper Ultrasonic Surgical Aspirator) and laser were also applied. In 5 cases intraoperative monitoring of somatosensory evoked potentials (SSEP). Was used the dural attachment was completely resected if the spinal meningioma was located dorsally or dorsolaterally. In these cases, duraplasty was performed with autologous fascia obtained during the operative approach. In ventrally located tumours the dural attachment was not excised but extensively bipolar cauterized After tumor completely removal and careful hemostasis, the dura was primarily closed in a watertight manner. The grading system according to Simpson (1957) for meningiomas in this area. Therefore, meningiomas of this study resection was defined as “complete” according to intraoperative observations and postoperative MRI corresponding to Simpson’s grade I or II.

Results

There were 14 female and 2 male patients. Age ranged from 17 to 88 years (mean 69 years). The mean follow-up period was 12 months including a complete neurological examination and postoperative MRI studies Tumors site was thoracic, intradural, extramedullary, most common, posterolateral 11 cases, antero-lateral 2 cases, anterior to the spinal cord 1 cases; cervical intradural, extramedullary, postero-lateral 1 cases, anterior to the spinal cord 1 case and no cases were reported in the lumbar region in this study motor deficits were in 81.25% of the patients. Gait disturbances were observed in 75%. In 37.5% the patients were unable to walk due to significant neurological deficits and half of the patients (50%) complained about local pain. Sensory loss in (50%) Sphincter disturbances were found in 13 patients (3 1.25%). Histopathology revealed the presence of meningiomas WHO grade I lesions: Meningotheliomatous type 11 cases, psammomatous type 1 cases, transitional type 1 cases, fibrous type 1 case, microcystic one case and only 2 cases a typical meningioma WHO grade II in 14 case (87.5%) the spinal meningioma was removed completely (Simpson’s grade I or II) and in 2 cases (12.5%) incompletely (Simpson’s grade III). Partial tumour calcification was observed in two patients (12.5%) and complete calcification in one cases (6.25%). Transient motor deficits were observed in 2 cases anteriorly placed tumors one thoracic and one cervical A postoperative DVT occurred in two cases, a cerebrospinal fluid fistula in 2 cases and a prolonged wound healing in one case. None of these patients required surgical repair and all recovered completely with conservative treatment we had recurrence in one patient which had a complete tumour resection initially his histological examination was WHO grade I the patient with recurrent tumour underwent a second operation. The tumour was resected completely and the new histopathology was semimalignant meningioma. Considering the functional outcome 63.5% who were able to walk indecently became 81.25%.

Case presentation:

Thirty five year old, housewife, female patient.

Complaints:
- Back pain for 9 months.
- Bilateral lower limb heaviness for 1 months.

Examination:
- Bilateral lower limb weakness Grade IV.
- Mild hypertonia in both lower limb.
- Mild hyperreflexia in both lower limb.
- Sensory level at the umbilicus.
- Continent.
- No pathological reflexes.

Imaging:
- MRI dorsal spine was done showed: D11,12 well-circumscribed, broad-based dural attachment mass, homogeneous enhancement compressing the spinal cord.
- In T1 isointense to slightly hypointense.
- In T2 isointense to slightly hyperintense.

Operative findings:
- Excision was performed through posterior approach.
- Midline back skin incision.
- Lamineotmy of D11, 12.
- Opening of the dura matter.
- Total surgical excision of the Intradural extramedullary mass. Extracted tumor showing a well encapsulated mass measuring 1.6x1 x0.8cm.
- Closure of the dura and wound in layers.

Postoperative: The patient has excellent result pain and weakness improved.

Post operative MRI: Was performed showing mass removal at D11,12 level.

Pathology: Transitional meningioma WHO G 1.

Fig. (1): Preoperative sagittal T1-weighted MRI with contrast showing homogeneous enhancement D11, 12 lesion.

Fig. (2): Preoperative sagittal T1-weighted MRI showing slightly hypointense D11,12 lesion.
Fig. (3): Preoperative sagittal T2-weighted MRI slightly hyperintense D11, 12 lesion.

Fig. (4): Preoperative axial T2 weighted MRI slightly hyperintense D11, 12 lesion compressing the cord.

Fig. (5): Preoperative axial T1 weighted MRI with contrast showing homogeneous enhancement D11, 12 lesion.
Fig. (6): Intra-operative findings of intradural extramedullary mass excision.

Fig. (8): Post-operative sagittal T1W MRI showing mass removal at D11-D12 level.

Fig. (7): Post-operative T2W MRI showing mass removal at D11-D12 level.

Fig. (9): Post-operative sagittal T1W MRI with contrast showing total mass removal at D11-D12 level.

Fig. (10): Post-operative axial T1W with and without contrast MRI with contrast showing total mass removal at D11-D12 level.
Discussion

Meningiomas are benign tumours mostly located in the intracranial compartment. Spinal meningiomas are rare and account about 1.2% of all meningiomas and 25% of all spinal cord tumours [8]. The first case of a successful removal was accomplished by Victor Horsley more than a century ago [10,11]. The primary goal of surgery is to achieve complete tumour removal and to avoid additional neurological damage. In the era before the introduction of the microscope in neurosurgery, good results were reported although some postoperative transient neurological deterioration was observed by some authors [7,12,13]. In the last decades, safety of neurosurgical procedures increased due to technical developments for tumour resection and the routine use of neuropsychological intraoperative monitoring [14-16]. There is a female predominance in the adult population: female to male ratio is 2:1 in intracranial meningiomas and 9:1 for spinal meningiomas, no sex predilection in children [17-20]. In my present series, the female/male ratio was 7:1. Female predominance could be explained by hormonal factors. Although the theory about the effect of hormones on meningioma has been controversial, hormonal studies have shown the existence of progesterone and estrogen receptors frequently found on histological examination, as well as the reports of an association between meningioma and breast cancer.

The presumed site of origin of spinal meningiomas from arachnoidal cap cells located in the leptomeninges at the spinal nerve root exit zones, adjacent to them denticulate ligaments or entry zones of arteries in the spinal canal; explaining why these tumors frequently arise in a lateral location to the spinal cord [21,22]. The most common topography is the thoracic spine 80%, (laterally 45-71%, postero-laterally 10-31% or antero-laterally to the spinal cord 15-27%) in our study the results were 87%, in the thoracic region and their distribution was (laterally 68.75%, postero-laterally 12.5% or antero-laterally to the spinal cord 6.25%) in the cervical it is present in 15% of cases and tends to be more anterior in our study it was 12.5% of cases spinal meningioma is very rare in lumbar spine 1-2% our study did not include any lumbar cases. Unspecific symptoms as local pain are often misinterpreted, until diagnosis is confirmed by neuroimaging procedures. At the time of diagnosis, however, a considerable number of patients were not able to walk independently. Delay of diagnosis as the result of failure to consider a slow growing spinal tumour responsible for longstanding back pain or neurological deficits was already observed by Pena et al., [23]. In our series the rate of patients unable to walk was 37.5%. This is in accordance with the literature. Klekamp et Samii reported that 59% demonstrated gait ataxia as predominant neurological symptom on admission, and a total of 31 out of 117 patients (26.5%) were unable to walk [18]. In other series, the rate of patients unable to walk independently ranges between 21 and 53% [4-6,8,10,15,18]. In our study, the rate of complete resection of spinal meningiomas was 87.5%. This rate is in accordance with the literature, where the rates of complete tumour removal are reported to be between 82 and 98% [4-8,13,15,18]. Complete resection according to Simpson’s grade II was performed in our study group. The rate of resection of the dural attachment is reported to be between 14 and 58% [8,13,15,18]. This is the result of the posterior operative approach in almost all cases and the difficulty of dural reconstruction ventral to the spinal cord. Tumour resection and following bipolar coagulation is generally considered to be adequate and effective. Concerning the functional results of surgery. In our study the outcome was improved or unchanged in 93.7% at the time of last follow-up. Before surgery only 63.5% of our patients were able to walk independently, after surgery significant improvement was observed and this rate increases up to 81.25%. Similar results were reported by others. In Klekamp. Samii’s series, 31 out of 117 patients were unable to walk preoperatively. Of these, 29% could walk again before discharge and 57% within 3 months of rehabilitation [18]. In earlier series, others observed that the immediate postoperative course was characterized by a more or less pronounced transient neurological deterioration [18]. In our study we faced it in 2 cases After the introduction of the operative microscope and microneurosurgical procedures, the percentage markedly declined. Excellent or good results range between 79 and 98% [4,7,8,13,15,18]. Potential risk for permanent neurological deterioration. In our study, the existence of tumour calcification bears an increased risk for poor neurological outcome [8]. The evidence of calcification makes surgical removal difficult and increases the risk for neurological damage. Moore et al., evaluated 30 patients over 70 years and reported improved functional outcome in all cases and thus denying the relation between age and outcome [5-22]. In our study we had recurrence in one case 6.25%. The recurrence rate for spinal meningiomas is generally low ranging between 0 and 13% although the grade of tumour resection was Simpson’s grade II in the majority of in the presented series [4,8-10,18]. Mirmanoff et al., evaluated the recurrence rate of
meningiomas and their progression after neurological resection. They observed no tumour recurrence within the first 5 years, but a rate of 13% at the 10 years follow-up [9]. In the literature, there seems to be no doubt that there is no correlation between the extension of dural resection and the risk for tumour recurrence. Some authors rather observed a higher recurrence rate in patients with radical tumour resection compared to patients, where the dural attachment was coagulated only [8, 18]. Reoperation with total tumour resection is generally possible, but remains a surgical challenge.

Some centers advocate adjuvant radiotherapy for recurrent tumours, but there is still a lack of evidence that radiotherapy reduces the risk of tumour regrowth [24,9,13]. Thus, adjuvant radiation treatment should be preserved for malignant spinal meningiomas only.

**Conclusion:**

Despite the retrospective character of this study and its limited cases we can conclude that resection of spinal meningiomas is a safe and most effective procedure. There might be a higher surgical risk for calcified tumours, incomplete resection. According to Simpsons grade II is the result of the posterior operative approach in almost all cases and the difficulty of dural reconstruction ventral to the spinal cord. Tumour resection and following bipolar coagulation is generally considered to be adequate and effective. And does not increase the recurrence rate.

**References**

Surgical Management of Spinal Meningioma

التدخل الجراحي للورم السحائي في العمود الفقري

الخلاصة: الورم السحائي هو الورم الشائع الذي يمثل 4-26% من الورم الشعوبي. هو عموداً صغيراً، و بطينه النمو. تحدث الأورام السحائية في العمود الفقري بعد العقد الرابع من الحياة، معظمها تقع في الحورة الدماغية. تحتوي عموماً على نمط النخاع الشوكي للحبل الشوكي الموضع الأكثر شيوعاً للأورام السحائية الشوكي في منطقة الصدرية. تتوزع الأعراض من الخلل الوظيفي العصبي الخفيف إلى الكبيرة. الأعراض الأكثر شيوعاً هي الألم الظهر وعجز الأحماس بالحركة. يجب أن يكون الهدف من العلاج الجراحي هو الاستئصال الكامل. إن أمركاً، وعادة ليس صعباً.

الهدف: تم إجراء هذه الدراسة لعرض الإصابة والأماكن، النقاط، المستخدمة في علاج ورم سحائي نخاعي الشوكي وتقييم النتائج الوظيفية للحالات المعالجة جراحياً.

المرضى والأساليب: هذه الدراسة استرجاعية لستة عشر مريضاً الذين يعانون من الأورام السحائية الشوكي التي يمضوا لاستئصال المجرىية تعاملوا بين عامي 2012 و2015 في فضننا. تم تقييم الحالات، التشخيص، والفحص النسيجي، والاستئصال الجراحي المجهري، والنتائج الوظيفية، وتحديد عوامل التشخيص المحتملة المرتبطة بهذه الأورام.

النتائج: يعتبر التدخل الجراحي لإزالة الأورام السحائية طريقة فعالة وأمنة مع وجود مصوبات جراحية في حالات الأورام المتكلسة. وعدم أزة أورام عاماما في بعض الحالات وذلك لاحتمالها بالليمفاوية أو وجود أورام أمام النخاع الشوكي ومع ذلك فإن الكلي كافي لازالة المتكتة من الورم مع عدم ارتفاع نسبة الأورام المرجعة. استئصال كاملاً في 14 رضي (81.2%) وغير كاملة في 3 رضي (18.8%). (χ2 = 12.5) في آخر مناية لهم تم تحسين الحالة العصبية أو دون تغيير في 15 رضي (82.7%) وأسوا في رضي واحد (17.3%).

الخلاصة: التصوير بالرنين المغناطيسي هو أفضل تقنية لتصوير التشخيص. تسجيل استئصال الورم الكلي يعني تحسين ملحوظ

في النتائج الجراحية لأورام سحائية العمود الفقري.