Intramedullary Spinal Cord Tumors: Part I—Epidemiology, Pathophysiology, and Diagnosis

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Abstract

Study Design Broad narrative review.

Objectives Intramedullary spinal cord tumors (IMSCT) are rare neoplasms that can potentially lead to severe neurologic deterioration, decreased function, poor quality of life, or death. As such, a better understanding of these lesions is needed. The following article, part one of a two-part series, addresses IMSCT with regards to their epidemiology, histology, pathophysiology, imaging characteristics, and clinical manifestations.

Methods The authors performed an extensive review of the peer-reviewed literature addressing the aforementioned objectives.

Results Numerous IMSCT exist with varying epidemiology. Each IMSCT has its own hallmark characteristics and may vary with regards to how aggressively they invade the spinal cord. These lesions are often difficult to detect and are often misdiagnosed. Furthermore, radiographically and clinically, these lesions may be difficult to distinguish from one another.

Conclusions Awareness and understanding of IMSCT is imperative to facilitate an early diagnosis and plan management.

Introduction

Primary spine tumors are rare neoplasms that can lead to significant patient morbidity and mortality.¹,² Intramedullary spinal cord tumors (IMSCTs) are the rarest of these neoplasms and can potentially lead to severe neurologic deterioration, decreased function, poor quality of life, or death.² Within the IMSCT category, the most common lesions are ependymomas, astrocytomas, and hemangioblastomas, followed by other rare lesions.²,³ Each specific lesion has its own hallmark characteristics, and the tumors vary with regards to the current understanding of their etiology or associations, their imaging and clinical characteristics, and/or how aggressively they invade the spinal cord. Due to the rarity of occurrence, these lesions are often difficult to detect and are often misdiagnosed, leading to delays in the appropriate patient care.⁴ Additionally, despite the hallmark characteristics and current diagnostic abilities, these lesions remain both radiographically and clinically difficult to distinguish from one another.⁵ Preoperative neurologic status and tumor histology are two of the most important variables affecting treatment outcome with these lesions, and thus awareness of these lesions is imperative to facilitate an early diagnosis.⁶–⁹ Early diagnosis allows early referral for treatment, which can be the critical difference in the treatment outcome. The following is a broad narrative review of the literature...
addressing IMSCTs with regards to their epidemiology, histology, pathophysiology, imaging characteristics, and clinical manifestations.

**Epidemiology**

**Incidence**

Primary spinal cord tumors are 10 times less common than their cranial counterparts, although they are histopathologically similar. Spine tumors are historically classified as (1) extradural, (2) intradural extramedullary, and (3) intradural intramedullary. Extrudural tumors are the most common spinal tumors and are usually of metastatic origin.

Intradural intramedullary lesions comprise 20 to 30% of all primary spinal cord tumors. The remaining 70 to 80% of primary intradural tumors are intradural extramedullary tumors. Intradural tumors are most likely to be found in the thoracic region, followed by the cervical and then lumbar spine.

Gliomas make up 80% of all intramedullary tumors. The gliomas are further subdivided into astrocytomas (60 to 70%) and ependymomas (30 to 40%). Astrocytomas have a higher occurrence in children than adults, whereas ependymomas are more common in adults than children. Hemangioblastomas are the third most frequent IMSC tumor, comprising 2 to 15% of all intramedullary tumors.

Metastatic intramedullary tumors are rare but present in 2% of all intramedullary tumors. Autopsy series have reported that 0.9 to 2.1% of all patients with cancer exhibit intramedullary spinal cord metastases. Thus, the incidence of these tumors may be higher than initially predicted. Of the intramedullary tumors with metastatic origins, 40 to 60% and 14% arise from primary neoplasms of the lung and breast, respectively. The remaining constituents of IMSCTs include lipomas (1%) and other rare tumors that have been documented sparingly in the literature (Table 1).

**Genetic Factors**

Several genetic factors are associated with IMSCTs. Understanding the various genetic mutations assists in illustrating the clinical manifestations, progression, and management of these tumors. Additionally, through a brief understanding of tumor genetics, the association of spinal tumors with lesions elsewhere in the body can be appreciated. The clinical syndromes currently associated with IMSCTs include neurofibromatosis 1, 2 (NF-1, NF-2) and Von Hippel-Lindau disease (VHL).

**Neurofibromatosis**

Neurofibromatosis is a common autosomal-dominant disorder with 100% penetrance in familial lines. Two subtypes have

| Tumor class                  | Associated tumors                                      |
|------------------------------|--------------------------------------------------------|
| Neuroepithelial tissue       | Astrocytic                                               |
|                              | • Astrocytoma                                           |
|                              | • Glioblastoma                                          |
|                              | Embryonal                                                |
|                              | • Primitive neuroectodermal                              |
|                              | Ependymal                                                |
|                              | • Ependymoma                                             |
|                              | • Subependymoma                                          |
|                              | Mixed glioma                                             |
|                              | Neuronal and mixed neuronal-glial                        |
|                              | • Gangliocytoma                                          |
|                              | • Ganglioglioma                                          |
|                              | • Ganglioneuroblastoma                                   |
|                              | Oligodendroglial                                         |
|                              | • Oligodendroblastoma                                    |
|                              | Uncertain origin                                         |
|                              | • Polar spongioblastoma                                  |
| Spinal nerves                | Neurofibroma                                             |
|                              | Schwannoma                                              |
| Nonmeningotheial, mesenchymal| Hemangioblastoma                                         |
|                              | Lipoma                                                   |
|                              | Melanoma                                                 |
|                              | Sarcoma                                                  |
| Germ cell tumors             | Germinoma                                                |
|                              | Teratoma                                                 |
| Cysts and tumorlike lesions  | Dermoid                                                  |
|                              | Epidermoid                                               |
| Hematopoietic neoplasms      | Primary central nervous system lymphoma (microglial)     |
| Metastatic tumors and other  |                                                         |
| rare neoplasms               |                                                         |
been established: NF-1 and NF-2. Fifty percent of patients with neurofibromatosis possess a family history of the disorder with new mutations developing in the remaining constituents.

The reported prevalence of NF-1 (also known as von Recklinghausen disease or peripheral neurofibromatosis) is 1 in 3,000 to 4,000 individuals. A Although the incidence of IMSCT in individuals with NF-1 is speculative, it has been suggested that almost 19% of subjects diagnosed with NF-1 develop IMSCT. Expressivity varies due to the heterogenetic characteristics of the disorder. The mutation is located on the long arm of chromosome 17 (17q), which encodes for neurofibromin, a negative regulator of the RAS cellular proliferation pathway. Neurofibromin is a tumor suppressor gene, whereby mutation disinhibits the RAS pathway leading to increased cellular division and proliferation and eventual tumor development. In the presence of NF-1, multiple neurofibromas may appear. Although many tumors are associated with NF-1, in relation to IMSCTs, astrocytomas are the most likely to develop (►Table 2). Less prevalent than NF-1, NF-2 (also known as central neurofibromatosis) occurs in 1 in 40,000 individuals and represents 2.5% of patients with IMSCT (►Table 2). The location of NF-2 has been identified as a mutation of the “Merlin” gene on chromosome 22q12. NF-2 is largely associated with ependymomas and occasionally meningiomas (extramedullary). The severe form of the condition exhibits multiple tumors with an earlier onset and a rapid clinical deterioration. The milder form is characterized by fewer tumors with a later onset and a gradual clinical progression.

### Von Hippel-Lindau Disease

VHL is a rare autosomal-dominant disease with 90% penetrance. Moller first suggested that VHL was a genetic disorder in 1929. Briefly summarized, mutation of a tumor suppressor gene located on chromosome 3p25–26 is responsible for this condition. The VHL tumor suppressor proteins form a protein complex that interacts with elongins B and C and other proteins, marking them for degradation by the cell, which inhibits hypoxia-related cell transcription factors (HIF1a, HIF2a). The VHL proteins also suppress the hypoxia-induced production of vascular endothelial growth factor, erythropoietin, and platelet-derived growth factor. Without the VHL protein present, an increased presence of these transcription factors occurs on the cellular level, eventually leading to tumor formation, and given the specifics of the transcription factors involved, these are often highly vascular lesions. VHL is characterized by widespread formation of both benign and malignant tumors throughout the body. According to Melmon and Rosen, VHL is diagnosed if the following manifest: the presence of more than one hemangioblastoma of the central nervous system (CNS), the presence of an isolated lesion associated with a visceral manifestation of the disease, or the presence of one characteristic of the disease and a family history of the disease.

### Table 2 Genetic basis of intramedullary spinal cord tumors

| Genetic disorder          | Associated tumor                                      |
|---------------------------|-------------------------------------------------------|
| Neurofibromatosis-1       | Acute nonlymphocytic leukemias                        |
|                           | Astrocytomas                                          |
|                           | Carcinoid tumors                                      |
|                           | Hematomas                                             |
|                           | Hypothalamic gliomas                                  |
|                           | Malignant nerve sheath tumors                         |
|                           | Meningiomas                                           |
|                           | Neurofibromas                                         |
|                           | Optic nerve gliomas                                   |
|                           | Pheochromocytomas                                     |
|                           | Primitive neuroectodermal tumors                      |
|                           | Rhabdomyosarcomas                                     |
|                           | Wilms tumor                                           |
| Neurofibromatosis-2       | Bilateral acoustic schwannomas                        |
|                           | Epignymomas                                           |
|                           | Gliomas                                               |
|                           | Meningiomas                                           |
|                           | Schwannomas                                           |
| Von Hippel-Lindau         | Adrenal pheochromocytomas                             |
|                           | Central nervous system hemangioblastoma (medulla oblongata or spinal cord) |
|                           | Epididymal cystadenomas                               |
|                           | Lindau tumor (cerebellum hemangioblastoma)            |
|                           | Nephritic cysts                                       |
|                           | Pancreatic cysts                                      |
|                           | Renal cell carcinomas                                 |
|                           | Renal cysts                                           |
|                           | Retinal hemangioblastomas                             |

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wide array of tumors is associated with this disease, and especially common are retinal hemangioblastomas († Table 2). Hemangioblastomas of the medulla oblongata and spinal cord as well as renal cell carcinoma may ensue and lead to significant mortality in 50% of individuals with the disease. Of CNS hemangioblastomas, 80% occur in the posterior fossa and 20% appear in the spinal cord. Furthermore, 10 to 15% of cranial hemangioblastomas are associated with VHL, whereas 25% of spinal cord hemangioblastomas are associated with VHL.1,3 In association with VHL, there is an earlier age of development of hemangioblastomas than in those of sporadic origin.

Syringomyelia

The classification and terminology related to fluid-filled cavities in the spinal cord remains controversial. In this text, we will refer to the tumor-associated fluid-filled cavity within the spinal cord as syringomyelia or syrinx.26 Syrinx occurs with 25 to 58% of patients with IMSCT, and in this association, syringes appear more commonly in the lower cervical and upper thoracic region. They occur most frequently with ependymomas, followed by hemangioblastomas and cavernomas.27-31 Irrespective of tumor type, there is a correlation between the degree of cephalad extension for a tumor and the presence of a syrinx. According to Samii and Klekamp, 49% of syringes occur above the tumor level, 11% occur below tumor level, and 40% are bipolar. A small percentage involves the medulla oblongata.29 The presence of a syrinx is considered a favorable prognostic sign. Noninfiltrative tumors with distinct cleavage planes are more likely to display syringes than more diffuse, infiltrative tumors. After tumor removal is performed, the syrinx will typically dissolve. Moreover, patients presenting with the concomitant finding of syringomyelia with their tumor tend to recover from surgery more rapidly.

As mentioned, the classification and etiology of syringomyelia is a current area of investigation and a continually evolving field, but it can be generally divided into three broad subgroups: (1) resulting from alteration of cerebrospinal fluid (CSF) flow dynamics related to hindbrain disorders, (2) resulting from intramedullary tissue damage secondary to hemorrhage or infarction, and (3) resulting from direct secretory ability of intramedullary tumor.29 The latter two mechanisms of syrinx formation pertain to IMSCT.27,32-37 The resultant syringes from hemorrhage and infarction can be seen occasionally in ependymomas, which often hemorrhage, and especially so in highly vascular lesions such as hemangioblastoma and cavernoma, which are the most common lesions to have associated syrinx. The secretory theory proposes that pathologic tumor vessels create transudation and the secretion of fluid from tumor cells, which thus account for syrinx development. Because the chemical composition of the tumor syrinx has an elevated protein concentration compared with CSF and syrinx fluid associated with non-IMSCT, the syringomyelia may be inherent to the tumor. These mechanisms of syrinx formation in relation to IMSCT are wholly different from those related to CSF obstruction and hindbrain disorders, which include such theories as Gardner’s “water hammer” theory, Williams’ “suck and slosh” theory, Ball and Dayan’s theory of infiltration through perivascular (Virchow-Robbin) spaces, and Oldfield’s theory, which combines features of all of the above.26,35,37 However, tumor obstruction of the CSF dynamics and subsequent relative blockage in CSF flow at the local level may also have a role in the pathogenesis of syrinx formation by increasing the focal transmedullary pressure in addition to the two main tumor-related etiologies of hemorrhage/infarction and secretory ability.29,38

Pathophysiology

Astrocytomas

Astrocytomas are red, gray, glossy tumors that are characterized by a poorly defined plane and are generally infiltrative in nature. Astrocytomas are the second most common IMSCT in adults at 30 to 35% of tumors and the most common in children at 90% of tumors.1,10,39 Of all astrocytomas arising in the CNS, 3% occur in the spinal cord.40 They primarily occur in the cervical spine, and they often involve multiple spinal segments due to the expansive nature of these tumors (→ Fig. 1). Nearly 20% of such lesions are associated with syrinx formation.29 Astrocytomas have an association with NF-1, occur predominantly in males, and rarely manifest in patients over the age of 60.41 Adults mainly exhibit high-grade lesions, whereas low-grade lesions are associated with the younger population.42 Malignant degeneration develops in 25% of adult astrocytomas.43

Ependymomas

Ependymomas are soft, encapsulated, reddish gray or yellow tumors with modest vascularity that present in the third or fourth decade of life. Half of all ependymomas are located in the region spine and a slight majority are located in the cord (55%), relative to the cauda equina (45%).1 They are not sex discriminant.44,45 The majority of them are classified as benign tumors, and ependymomas have a propensity to grow slowly. Spinal ependymomas are primarily present in the cervical or cervicothoracic region.46,47 The cysts associated with these tumors are predominantly found in the superior margin of the tumor and have an increased incidence in the cervical region (→ Fig. 2A).29,43,48 Approximately 65% of these tumors are associated with syrinx formation.29

Originating from ependymal cells, ependymomas are more centromedullary located as compared with astrocytomas, and they tend to blend with the cord (→ Fig. 2B). They appear as a focal enlargement within the spinal cord. Ependymomas have a mean extension of three to four vertebral levels.49 Abnormalities of chromosome 22 are associated with ependymomas, and this leads to their association with NF-2.19 Various histologic subtypes of ependymomas exist. The most common is the cellular variant or the “typical” form, which is a World Health Organization (WHO) grade II tumor that is well circumscribed. Pathologic analysis reveals both true ependymal rosettes and more commonly perivascular
Tanycytic ependymomas are also WHO grade II, are less common overall, and are found more commonly in the spinal cord than in the brain. Histologically, tanycytic ependymomas lack ependymal rosettes and their pseudorosettes are vaguely apparent, which can lead to misdiagnosis as pilocytic astrocytoma (WHO grade I). 

Fig. 1 (A) Sagittal T1-weighted post–gadolinium contrast cervical magnetic resonance image (MRI) noting a heterogeneously enhancing intramedullary astrocytoma (arrow). (B) Sagittal T2-weighted cervical MRI demonstrating an intramedullary astrocytoma (arrow). (C) Axial T1-weighted post–gadolinium contrast cervical MRI illustrating a cystic astrocytoma; the cystic portions are highlighted by arrows and are hypointense compared with the enhancing surrounding rim.
Myxopapillary ependymomas are WHO grade I tumors commonly found at the filum terminale, but these are considered extramedullary (Fig. 3). Rarely anaplastic ependymomas, WHO grade III, can be encountered, which develop more rapidly than lower grades, can occasionally develop from malignant degeneration, and have an overall poor prognosis.

Ependymomas may also be found outside the nervous system in the soft tissue, ovaries, and mediastinum. Subependymomas (WHO grade I), which may be histologically related to ependymomas, have also been reported as rare IMSCT.

Hemangioblastoma

Hemangioblastomas are small benign richly vascularized solitary neoplasms that rarely extend beyond one or two segments. Although more often located in the cerebellum, hemangioblastomas are also found in the spinal cord, predominantly located at the posterior or posterolateral region of the spinal canal (Fig. 4). Most hemangioblastomas develop sporadically but, as previously mentioned, they are associated with VHL, especially when presenting in the spinal cord. They constitute between 2 and 8% of all IMSCTs.\(^1\) Commonly, they are located in the cervical spine but may be found in any portion of the neuroaxis. Cyst formation is evident within 50 to 70% of tumors, and the tumors have an association with syrinx.\(^{52-54}\) Hemangioblastomas are more commonly found in men.\(^{52-55}\) Although hemangioblastomas are not age discriminant, most occur during the fourth decade of life. It is possible that individuals with a hemangioblastoma can remain asymptomatic throughout life, with tumor only apparent at autopsy.\(^55\)

Hemangioblastomas are composed of a dense network of vascular capillary channels containing endothelial cells, pericytes, and lipid-laden stromal cells.\(^{56}\) The cells of origin remain unknown, but based on the genetics of VHL and mutations found in sporadic lesions, they are likely vascular endothelial growth factor-secreting undifferentiated mesenchymal cells.\(^1,19\) Two histologic forms exist: reticular and cellular. The former is composed of irregular nuclei with conspicuous vessels. Conversely, the cellular form exhibits fewer conspicuous vessels with varying stromal cells; this form bears a similarity to astrocytomas and may be difficult to discriminate from these lesions.

Hemangioblastomas are well-demarcated tumors that are amenable to resection. With complete resection, the prognosis is excellent. Pain and sensory changes are the most frequent presenting complaints. Imaging is a crucial factor in surgical planning and should be thoroughly evaluated. An associated cord widening is commonly found adjacent to the tumor, possibly attributed to edema related to increased blood flow in surrounding draining veins or within the tumor creating vascular congestion. After tumor removal, widening regresses. In addition, spinal angiography is recommended to delineate the characteristics of the spinal draining veins.\(^{57}\)
The literature supports the belief that metastatic IMSCTs are rare. More recently, combining routine magnetic resonance imaging (MRI) and autopsy observation, it appears that metastatic IMSCTs may be more common than initially reported. These fast spreading tumors may escape diagnosis, as patients bearing these lesions may be asymptomatic. Although imaging is helpful in demarcating the location and extent of these tumors, the proper diagnosis is accomplished by surgical biopsy and the presence of a primary source of cancer.

The mechanism of cord infiltration of metastatic IMSCT is ambiguous. Because the majority of metastatic IMSCTs metastasize from the lung, the tumor is thought to spread from hematogenous dissemination leading to arterial embolization.\textsuperscript{15,58,59} Other proposed theories of tumor seeding involve retrograde infiltration via the spinal cord venous system (Batson’s plexus),\textsuperscript{60} spread through perforating veins in the bone, metastatic antidromic cellular migration via nerve root to the spinal cord, CSF dissemination through intraspinal perineural sheaths, and finally, penetration of the spinal cord parenchyma via penetrating vessels within the Virchow-Robin spaces.\textsuperscript{14}

Lipomas
Lipomas are rare congenital tumors that constitute 1% of intraspinal tumors. Commonly found in the cauda equina and conus medullaris, these tumors tend to violate the posterior cord and are usually extramedullary.\textsuperscript{61,62} Lipomas are commonly associated with spinal dysraphism and are thought to arise from premature disjunction of the cutaneous ectoderm from the neural ectoderm prior to neural tube closure, allowing mesenchymal cells to infiltrate into the neural groove.\textsuperscript{1} These tumors possess a higher water content than other intramedullary tumors and tend to attach firmly to the dura; their cellular content is indistinguishable from normal adipose tissue.\textsuperscript{62–64} These tumors are slow-growing, and they become symptomatic due to focal mass effect. They present no cleavage plane from the surrounding cord with blending of the neural and fatty tissue.\textsuperscript{1,65} As a result, complete tumor resection is extremely difficult, and some neurologic damage may be expected when attempting resection. Although little is known about these tumors, patients tend not to improve from their baseline neurologic state but the long-term prognosis is excellent with 90% progression-free survival at 16 years following total or near total excision and 35% progression-free survival at 10 years following a debulking procedure.\textsuperscript{1} Developmentally, these tumors prevent normal maturation of the surrounding neural tissue.

Other Rare Neoplasms
Numerous additional IMSCTs exist but are extremely rare. Cavernomas are rare vascular lesions that may hemorrhage repeatedly during the lifetime of an individual, and they can have associated syringes. Intramedullary lymphomas may occur in isolation or with other central nervous system lesions. They respond to chemotherapy. Gangliogliomas are benign tumors consisting of glial and neuronal cells that are estimated to comprise 3.8% of CNS tumors found in the upper cervical cord.\textsuperscript{66,67} Oligodendrogliomas possess ill-defined borders and less extensive cord growth compared with other IMSCTs; fewer than 50 cases have been literature reported. Other IMSCT neoplasms include melanocytomas, melanomas, fibrosarcomas, peripheral neuroectodermal tumors, and amyloid angiopathies (\textit{Table 1}).\textsuperscript{1,68–70}

Imaging
Radiographic evaluation is necessary to determine the location and extent of tumor involvement and may help to
features may help in this distinction. The presence of scoliosis can be indicative as advanced IMSCT can present with scoliosis even in adults. In adults, widening of the spinal canal is usually not noted except with large myxopapillary ependymoma in the conus medullaris or filum terminale. In children, widening of the spinal canal and kyphoscoliosis are more frequently encountered. Myelography has been utilized in the past to evaluate the cord contours; however, cord widening is sometimes difficult to appreciate and the procedure may exacerbate symptoms. Myelography has been supplanted by MRI except in rare circumstances.

MRI is the preferred modality in identifying and characterizing IMSCT. Each tumor has a defined T1-, T2-, and gadolinium-enhanced T1-weighted imaging pattern. MRI may be helpful in defining the presence or absence of a cord-tumor interface, illustrating associated cysts or syringomyelia, which may support the ability to achieve a favorable resection. With modern imaging capabilities, differentiation between ependymomas, astrocytomas, and even hemangioblastomas remains difficult; however, several radiographic features may help in this distinction. A recently published article by Arima et al scrutinized the ability of MRI to accurately differentiate between these three most common IMSCTs and were able to achieve an 89% accuracy. Although this appears to be a high rate of diagnostic accuracy, it does not yet achieve a value that allows treatment based on MRI features alone. Looking at gliomas specifically (ependymomas and astrocytomas), both lesions demonstrate fusiform configuration over numerous vertebral segments, and they both appear hypo- or isointense on T1-weighted images and hyperintense on T2-weighted images. Furthermore, these two tumors enhance with contrast. The rare subependymoma shares these MRI T1- and T2-weighted features, can have associated syrinx, and is usually eccentrically located and nonenhancing; contrast enhancement is faint if present.

Ependymomas tend to be centrally located within the cord and display symmetric expansion with diffuse heterogeneous enhancement (Fig. 2). They tend to occupy the whole width of the cord and generally produce enhanced margins (Fig. 2). Astrocytomas, in comparison, tend to be eccentrically positioned, may display an exophytic component, and can be nonenhancing or heterogeneously enhancing or even have an enhancing nodule (Fig. 1). Radiographically, they do not present with well-defined borders but large satellite cysts may be visualized. Polar cysts may appear and have a low signal intensity on T1-weighted images and high signal intensity on T2-weighted images. Intratumoral hemorrhage may be seen in both types of glioma but are more likely in ependymoma, with the imaging characteristics dependent on the age of the hemorrhage and possible secondary development of syrinx.

Hemangioblastomas are richly vascularized tumors that can have significant surrounding edema. They are associated with syringomyelia, can be found at any spinal level, and are most commonly sporadic. In patients with VHL, they are associated with additional visceral lesions or neuraxis hemangioblastomas. On MRI, they have mural nodules, which appear isointense on T1-weighted images and hyperintense on T2-weighted images with homogenous contrast enhancement, in contrast to the usual heterogenous contrast pattern of ependymomas and astrocytomas (Fig. 4). Vascular imaging or even spinal angiography may be useful to delineate feeding vessels and illustrate associated dilated pial veins due to vascular shunting and can be helpful for consideration of preoperative embolization in these highly vascular lesions.

Metastatic IMSCTs are well encapsulated and are unlikely to present with cystic change or intralesional hemorrhage. They most commonly present as single lesions in the thoracic cord but can be multiple or present in the cervical and lumbar spine. Metastatic lesions are most likely eccentrically located in the cord and expand the cord parenchyma, which was historically visible on myelography and illustrated clearly on MRI. MRI is the imaging tool of choice, and recently described “rim” and “flame” signs have been described to help differentiate non-CNS intramedullary metastases from primary spinal cord tumors, which occur significantly more frequently either alone or in combination in metastases than in primary cord tumors with high specificity but low sensitivity (specificity 97% for either alone and 100% when both present). The rim sign is defined by Rykken et al as a complete or partial rim of gadolinium enhancement, and the flame sign is defined as an ill-defined flame-shaped gadolinium-enhancing region at the superior or inferior margin of an otherwise well-defined lesion. Metastases are most likely to be T1 isointense and T2 hyperintense relative to the cord. The amount of edema may be out of proportion to the size of the tumor, and this is evidenced by extensive T2 hyperintensity, which can be on average 3.6-fold larger than that of the enhancing portion of the lesion. The estimated prevalence of spinal cord metastases from a post-mortem series ranged from 0.9 to 2.1% of patients with cancer. Any patient with a known history of cancer and an intramedullary spinal cord lesion should have metastatic IMSCT as part of the differential diagnosis.

Common IMSCTs may be difficult to differentiate from the rare tumors radiographically. Intramedullary lymphomas are rare tumors often linked to other supratentorial lesions. Oligodendrogliomas are characterized by ill-defined borders, no peritumoral edema, and slight T1-weighted image hyperintensity. Lipomas have imaging characteristics similar to fat (high T1-weighted image intensity).

Clinical Manifestations

IMSCtS present with a wide array of symptoms that vary in intensity and chronicity. The clinical features of each tumor are related to the growth rate, location, and longitudinal extent of the tumor. In general, spinal cord lesions should be suspected when patients present with bilateral motor and sensory symptoms not involving the head and face, often with other upper motor neuron symptoms consistent with a myelopathic syndrome. Hydrocephalus occurs in 1 to 8% of patients with IMSCT. The hydrocephalus may result from...
tumoral obstruction of the subarachnoid CSF flow or impaired CSF absorption induced by nondisseminated subarachnoid space tumors that increase CSF protein levels. Focal block of CSF flow by a tumor (or abscess formation) with increased CSF protein and presence of xanthochromia is known as Froin syndrome, which can lead to a "dry" lumbar puncture.87

The most common tumor presentation is back or neck pain. This is hypothesized to result from dural distension and irritation. The pain is of constant intensity and varies between individual patients; it is classically worsened in the recumbent position. Nerve root compression can produce weakness, spasticity, and clumsiness.88 Centrally located lesions can produce myelopathic symptoms. If the tumor extends cranially, cranial nerve involvement is possible. Paresthesias or dysesthesias can occur as a response to cord distension by the tumor mass. Decrease in temperature sensation and pain results from spinthalamic tract encroachment. Compression of the dorsal columns can manifest with defects in proprioception and gait abnormalities. Tumors affecting the autonomic pathways produce disturbances of the sympathetic and parasympathetic system. Severe cord defects can also complicate respiratory, bowel, bladder, or sexual function.19,88

Damage to cranial nerves is also a possibility with tumors located in the upper cervical spine. The hypoglossal nerve can be subject to compression by tumors located laterally at the foramen magnum leading to ipsilateral tongue paresis and atrophy. Arising from the C1–C5 anterior horn cells, the accessory nerve travels cephalad through the foramen magnum via the subarachnoid space between the ventral and dorsal roots and unites with its cranial counterpart. Thus, tumors around the upper cervical spinal cord can compress the accessory nerve, leading to weakness and atrophy of the sternocleidomastoid and trapezius muscles.89

Conclusion

IMSCCTs are rare manifestations that can severely impair neurologic function and the quality of life. The proper diagnosis of IMSCCT is essential to provide the best treatment and obtain the optimal outcome. The preoperative functional status, tumor histology, and degree of tumor invasion dictate postoperative results. Therefore, prompt diagnosis and intervention are essential. However, the manifestation of symptoms can be misleading and the diagnosis can be missed. Overall, various types of IMSCCT exist, and the management of these tumors demand individualized scrutiny and understanding.

Disclosures

Dino Samartzis, none
Christopher C. Gillis, none
Patrick Shih, none
John E. O’Toole, none
Richard G. Fessler, none

Acknowledgment

We would like to thank the Hong Kong Theme-Based Research Scheme (T12-708/12N) for their support of this work.

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