Case Report

Renal Cell Carcinoma presented with An Intramedullary Spinal Cord Metastasis : A Case Report

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Abstract:

Intramedullary spinal cord metastasis is one of the rare manifestations of systemic neoplasm causing diagnostic dilemma. Here we report a case of metastatic renal cell carcinoma presented initially as spinal cord lesion. A 62 years old Bangladeshi gentleman presented with spastic paraparesis, bone pain with unexplained weight loss. After thorough examination & detailed investigations we found intramedullary spinal cord metastasis with infiltrative lesion on bone scan. We confirmed renal cell carcinoma after doing CT Abdomen and histopathology as primary site of lesion. He was treated by radical nephrectomy & radiotherapy. It is imperative that clinicians should be cautious of occult carcinoma as the cause of suspicious intramedullary spinal cord lesion.

Keywords: Intramedullary spinal cord lesion, Renal cell carcinoma.

Introduction:

Although metastatic neoplasms of the spine are common, pure intramedullary spinal cord metastasis (ISCM) is a rare manifestation of systemic cancer, which indicates the occurrence of remote dissemination, and thus, the terminal phase of cancer1, 2, 3. The most common primary tumors of ISCM are from the lung, breast and melanoma, which account for about three-fourth, but lymphoma, kidney, colon and thyroid are uncommon4, 5, 6. Despite its rarity, intramedullary metastasis should be considered in patients with systemic malignancy presenting with myelopathic symptoms7. The recognition of intramedullary spinal cord metastasis is an ominous finding as it generally occurs in the setting of widespread systemic and intracranial disease and is the prelude to cancer death by a few months8. The presenting symptoms of ISCM vary from pain, sensory loss, weakness, urinary incontinence to pseudo Brown-Sequard or Brown-Sequard syndrome9. The duration of symptoms before diagnosis of ISCM ranges from days to a few months9,10. The very rare occurrence of ISCM and the absence of pathognomic symptoms often lead to an undue delay until the underlying malignancy is discovered10. Surgery and radiotherapy have been recommended controversially in the treatment of ISCM. Although long-term survival is poor, treatment may preserve ambulation in the case of early diagnosis; it may also stabilize neurological function and this may change the patient’s health related quality of life unbelievably9,11,12. Careful history & thorough examination may lead to early diagnosis which can halt neurological deficits & effective palliation can be offered to the patient.

Case report:

A 62-years-old man presented with 1month history of progressive unremitting back pain, exacerbated at night & walking, associated with perianal paresthesia & urinary incontinence. He also had rapidly progressing paraparesis without any radicular pain. On examination, his motor power of both hip flexion was 2/5 and other distal lower extremity was 3/5 along with hypertonia, hyperreflexia, bilateral extensor plantar with sensory level at the level of T12. He had undergone series of investigations including neuroimaging, CSF analysis, routine blood tests. MRI of dorsal-lumbar spine revealed hypointense thickening of spinal cord extending from the T4 to L1 spine on T1WI, patchy enhancement on contrast T1WI & hyperintense on T2WI suggesting spinal cord edema. An intramedullary ovoid lesion with thick rim enhancement & intalesional hyperintense area on T2WI was seen at the level of T12 spine (Figure 1). No infectious etiology had been found despite performing investigation. The tissue biopsy of the involved area was not obtained as he had not consented to it. On further query, he complained unexplained weight loss of 5 kg in the last 2 months. We went for evaluation of unexplained weight loss & found raised ESR, normal chest X-ray. Enlarged, distorted right renal mass

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was revealed on ultrasonography. CT scan of abdomen revealed heterogeneously enhancing soft tissue mass involving right kidney suggestive of right renal mass with focal calycectasis (Figure 2). CT guided FNAC confirmed the diagnosis renal cell carcinoma. Radical nephrectomy was done afterwards and the histology confirmed renal cell carcinoma, conventional type. Early Postoperative period was uneventful. He complained of severe pain around left scapula afterwards which was constant, exacerbated by movement of shoulder, unremitting with pain killers along with intractable back pain. X ray of the left shoulder along with spine was unremarkable. Whole bone scintigraphy revealed increased uptake of radiotracer L5 vertebra & inferior aspect of left scapula suggestive of infiltrative bone lesion (Figure 3). He also had hypercalcemia probably due to paraneoplastic effect of the tumour producing PTHrP, and bony metastases as the PTH assay was undetectable. He received appropriate treatment for his hypercalcemia. Finally we diagnosed the case as renal cell carcinoma with intramedullary spinal cord metastasis with bone metastasis. Neurosurgical consultation was taken & patient refused surgery. Focal radiotherapy on the spine was initiated due to his refusal for surgery. Over the following 10 days during radiotherapy, his neurological deficit progressed to 4/5 on the right lower extremity and 4/5 on the left lower extremity. There was no significant improvement of modalities of sensation with mild painful paresthesia on both lower extremities. He was able to walk with the support of walking aid after 1 month.

**Fig 1:** MRI of dorsal-lumbar spine showing an intramedullary ovoid lesion with thick rim enhancement & intalesional hyperintense area on T2W1 at the level of T12 vertebrae.

**Fig 2:** CT scan of abdomen showing heterogeneously enhancing soft tissue mass involving right kidney suggestive of right renal mass with focal calycectasis.

**Fig 3:** Whole bone scintigraphy revealed increased uptake of radiotracer L5 vertebra & inferior aspect of left scapula suggestive of infiltrative bone lesion.

**Discussion:**

Spinal metastases may be seen in as much as 70% of patients with systemic neoplasia. Among these lesions, intramedullary spinal metastases are rare, comprising only 0.5% of spinal axis metastases. Majority of them arise from the lung neoplasia, with small cell carcinoma being the Predominant
histological subtype. Breast, colorectal, renal, melanoma, thyroid and lymphoma have all infrequently been reported to be the origin. However, metastasis to the intramedullary spinal cord is extremely rare as the initial sign of a disseminated carcinoma. There are several hypotheses to explain how to metastasize into the spinal cord parenchyma. Kalayci et al. reviewed articles and summarized them into 3 pathologic mechanisms.

First, haematogenous spread via the artery or vertebral venous plexus (Batson’s venous plexus) is believed to account for most cases. Second, tumor cells originated from carcinomatous meningitis infiltrate into the Virchow-Robin spaces of the vessels, penetrate the pial membrane and invade the spinal cord parenchyma. The third mechanism is direct invasion from contiguous structures. In our case, considering the high incidence of anastomosis between the left renal vein and the vertebral venous plexus, hematogenous spread can be a reasonable mechanism. In patients without history of systemic neoplasia such as the patient in our report, a detailed clinical history and physical examination coupled with selected laboratory and diagnostic imaging investigations may delineate primary or secondary spinal malignant lesions from other more benign differential diagnostic entities. MRI is useful in determining the extent of CNS involvement which may affect the therapeutic decision making in many patients. Treatment modalities for ISCM include radiotherapy, surgery, and chemotherapy. The modality of treatment should be decided by clinical and neurological status. Focal radiotherapy has been accepted for an effective treatment modality for ISCM with arresting tumor growth and preventing further neurological deficit. However, the efficacy of radiotherapy may be limited to radio-sensitive tumors, such as small cell lung cancer, breast cancer, and lymphoma. These radio-sensitive tumors are most frequently found as the origin of ISCM. Previous reports favoring radiation therapy are somewhat biased as their clinical elements mainly consist of these radio-sensitive tumors. Despite the radio-resistance of RCC itself, focal radiotherapy has been preferred for the first-line treatment modality of ISCM from RCC due to the absence of effective systemic therapy for metastatic RCC and short life expectancy which estimated at 3 to 9 months. The prognosis of a patient who has an intramedullary cord lesion is grave. Spine surgeons are playing greater role in the management of patients with metastatic disease. With the advent of new surgical strategies many patients may benefit from effective treatment modalities ranging from radical, open excision through minimally invasive surgery such as endoscopy to ultraminimal/ noninvasive spinal radiosurgery. Making an early diagnosis of ISCM is useful in planning either early or no major intervention. Providing patients with successful palliation and improving their quality of life demand multidisciplinary strategic treatment planning.

Conclusion:
This report presents an extremely unusual presentation of metastatic renal cell carcinoma as intramedullary spinal cord lesion. Careful history & thorough examination can lead to early diagnosis and management of such cases which can reduce the neurological deficits & ensure better functional outcome.

References: