Primary pigmented meningeal melanocytoma originating in Meckel cave in a patient with carney complex: A case report

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Abstract

Rationale: Primary melanin-producing tumors are rare extra-axial neoplasms of the central nervous system. In the literature, few case reports have discussed neoplasms involving the cavernous sinus; of these, only 4 have reported on neoplasms originating in Meckel cave. The diagnostic approach, including clinical and radiological analysis, is challenging, and cytopathological assessment with a molecular basis is the best approach to discriminate between these lesions. Herein, we discuss the pathophysiology, diagnostic approach, intraoperative features, and postoperative management in a unique case of primary pigmented meningeal melanocytoma originating in Meckel cave in a patient who was diagnosed with Carney complex (CCx) and sickle cell disease (SCD).

Patient concerns: A 23-year-old man diagnosed with SCD had also been diagnosed previously with CCx, without any familial history or neurocutaneous melanosis. He had experienced headaches accompanied by left facial pain and paresthesia for 2 months.

Diagnosis: The initial computed tomography scan and magnetic resonance imaging (MRI) revealed a mass arising from the left Meckel cave. On MRI, it followed the signal intensity of melanin. He underwent subtotal resection of the mass. Considering the patient’s history of CCx, melanocytic schwannoma was the most relevant diagnosis. A postoperative histopathological examination was suggestive of benign pigmented meningeal melanocytoma.

Interventions: The patient underwent an uneventful subtotal resection of the mass through a left temporal linear incision.

Outcomes: The patient showed progressive improvement of neurologic deficits, and after 2 years of follow-up, he did not present with any new complaints.

Lessons: To the best of our knowledge, this is the first report of the unusual presentation of both SCD, as well as of primary pigmented meningeal melanocytoma in a patient with CCx. Complete surgical resection can be curative in most cases of melanocytoma. The presence of CCx with SCD suggests potential shared genetic contributions that will require further exploration.

Abbreviations: CCx = Carney complex, CD = cluster of differentiation, CT = computed tomography, HMB45 = human melanoma black 45, Melan-A = melanin antigen, MRI = magnetic resonance imaging, SCD = sickle cell disease.

Keywords: Carney complex, computed tomography, magnetic resonance imaging, melanocytoma, sickle cell disease.
1. Introduction

Primary melanocytic tumors of the central nervous system are extremely rare neuroectodermal neoplasms arising from the leptomeninges.[1] They commonly occur in the fourth and fifth decade of life and are very rare in children and adolescents.[2] The annual incidence of these lesions has been estimated to be approximately 1 per 10 million people, with a predilection for women.[3] They commonly involve the posterior fossa and upper cervical spine due to the presence of melanocytes in these sites.[4]

According to the World Health Organization classification of primary brain tumors, meningeal melanocytomas are classified under a subgroup of primary melanocytic neoplasms that have intermediate grade properties with low mitotic activity not fulfilling the criterion for malignant melanoma.[5] They either manifest focally, as is the case for melanocytoma or primary malignant melanoma, or in a diffuse manner, as is observed in cases of leptomeningeal melanocytosis and primary leptomeningeal melanomatosis.[6]

Only 8 patients have been reported with primary melanocytoma originating from the cavernous sinus, 4 of whom presented with melanocytoma originating specifically from Meckel cave.[2–5,11] Herein, we discuss the pathophysiology, diagnostic approach, intraoperative features, and postoperative management in a unique case of primary pigmented meningeal melanocytoma originating in Meckel cave in a patient who was diagnosed with Carney complex (CCx) and sickle cell disease (SCD).

1.1. Patient information

A 20-year-old male patient with a known case of SCD was diagnosed with CCx at our hospital when he was admitted with hypertension. During his first visit to the hospital, his work-up analysis revealed bilateral primary pigmented nodular adrenocortical disease, cardiac myxoma, and calcified testicular tumor (Fig. 1). He revisited our hospital 3 years later in June 2016, complaining of frequent headaches accompanied by left facial pain and paresthesia for the past 2 months.

1.2. Clinical findings

The patient’s symptoms gradually worsened, especially with chewing and exposure to the cold. Neurological examination revealed left fifth and sixth cranial nerve palsies. There was no history of neurocutaneous melanosis or CCx in his family.

1.3. Diagnostic assessment

An initial brain computed tomography (CT) scan revealed a well-defined mass that originated from the left Meckel cave. Its internal texture appeared heterogeneously hyperdense on an unenhanced CT scan (Fig. 2A). Cranial magnetic resonance imaging (MRI) further characterized the mass, presenting with a bright signal on T1-weighted images and very dark signal on T2-weighted images in accordance with the intensity of melanin. It showed a faint peripheral enhancement on the subtracted T1 post-contrast imaging (Fig. 3). The most relevant neuroimaging diagnosis was melanotic schwannoma as part of the CCx; other differential diagnoses included melanocytoma and melanoma. The maximum axial dimension of the mass increased from 0.4 cm in June 2016 to 1.9 cm in June 2017; this led to compression of the cisternal portion of the trigeminal nerve, thus explaining the patient’s symptoms.

1.4. Therapeutic intervention

Subsequently, subtotal resection of the mass was performed through a left temporal linear incision and by creating a burr hole above the zygoma, followed by craniotomy. The dura opened in a “C” shape, and the dura above the bulging mass was exposed; this revealed a coal-black, encapsulated mass, originating in Meckel cave and adhering tightly to the trigeminal nerve. The tumor was removed in pieces until the vessels were apparent and a subtotal tumor resection was achieved. Grossly, the tumor was a soft, fragile, pigmented mass covered with a thick white capsule (Fig. 4).

Microscopically, the tumor revealed nests of epithelioid-cell infiltration and spindle-shaped neoplastic cells with heavy melanin pigmentation. The cells contained typical macro-nuclei with few irregularities in the nuclear contours and prominent nucleoli. Nevertheless, the cytoplasm was abundant, and neither mitosis nor necrosis was observed.

The tumor was histopathologically examined for cell markers, including cluster of differentiation (CD)136, CD34, and KI67 protein. The tumor was also tested for immunoreactivity staining with epithelial membrane antigen, human melanoma black 45 (HMB45), microphthalmia-associated transcription factor, tyrosinase, vimentin, S100 protein, melanoma antigen (Melan-A), and reticulin. The results were difficult to interpret because of the heavy pigmentation; however, a positive expression of Melan-A, S-100, and HMB45 was noted. Significantly, there was total depletion of reticulin and an absence of Psammoma bodies. Other immunohistochemical findings were predominantly negative and cellular proliferation, measured by staining for Ki-67, was <1%.

However, based on these findings, a diagnosis of meningeal melanocytoma rather than melanocytic schwannoma or malignant melanoma was established (Fig. 5).

1.5. Follow-up and outcomes

The patient’s postoperative course was uneventful as observed through continuous follow-up imaging (Fig. 2B) and progressive improvement of the neurologic deficits. The patient was discharged after postoperative monitoring for 2 weeks, during which time no neurological symptoms were observed; he was followed-up regularly (every month for the first 6 months following discharge and every 3 months thereafter) at the neurosurgery clinic for 2 years and did not present with any new complaints.

2. Discussion

To the best of our knowledge, there are only 4 cases of primary meningeal melanocytoma originating in Meckel cave described in the literature (Table 1). Of interest, none of these were associated with SCD. Primary melanocytic tumors of the central nervous system arise from the leptomeningeal melanocytes, with a spectrum ranging from well-differentiated benign meningeal melanocytomas to malignant melanomas. However, atypical melanocytic melanocytomas, which have clinical and pathological characteristics between benign melanocytomas and malignant melanomas, are considered borderline tumors.[2]

These lesions typically appear on CT scans as extra-axial isodense to hypodense lesions, with dural and meningeal attachment. Further, they may show homogeneous enhancement
after contrast administration. On MRI scans, they usually present with homogeneous bright intensity on T1-weighted images and dark signal intensity on T2-weighted images because of the paramagnetic effects of free melanin radicals. Besides, they show homogeneous enhancement after gadolinium contrast administration. However, this signal pattern on MRI scans may vary based on the melanin and hemorrhagic concentration within the lesion; the lesion can present a heterogeneous signal intensity and a variable contrast-enhancing pattern.

Our case showed the typical CT and MRI appearance, which informed the provisional diagnosis of the lesion as melanocytic schwannoma, particularly in the context of all other findings and the presentation of CCx, which represents a spectrum of associated pathologies. Although melanocytomas reportedly...
Figure 2. Unenhanced axial computed tomography scans of the brain. (A) Heterogeneously hyperdense mass lesion occupying the left Meckel cave inducing mass effect along the ipsilateral lateral cavernous sinus and extending to a cisternal portion of the trigeminal nerve. (B) Follow-up post subtotal resection through left temporal craniotomy with the small residual lesion (asterisk).

Figure 3. Multisequential multiplanar magnetic resonance images of the brain with and without contrast. Well-defined extra-axial mass (arrowhead), arising from the left Meckel cave, showing bright T1 and dark T2 signal intensities (A–C). Interval increased maximum axial dimension of the mass at 1-year follow-up imaging (D–F). The subtracted image between post-contrast (E) and pre-contrast T1 (D) exhibits a faint peripheral rim of enhancement (asterisks) along the postero-lateral margin of the mass (F).
have imaging features similar to those of other dural tumors, after gadolinium contrast administration, the lesion in our case showed heterogeneous enhancement; in addition, after performing a subtraction of T1 post-contrast from conventional T1 pre-contrast sequence, a faint peripheral posterolateral enhancement was observed (Fig. 3). Interestingly, this was not reported in other cases in the literature (Table 1).

Histopathological analyses have played a significant role in the diagnosis of meningeal melanocytoma and have helped to rule out similar pathological conditions with similar characteristics. Microscopically, the majority of meningeal melanocytoma cases present as well-circumscribed, non-encapsulated lesions, consisting of several nests or fascicles of epithelioid and spindle cells with little nuclear atypia and abundant melanin pigmentation. Further, immunohistochemical diagnostic criteria require S-100 positivity, expression of Melan-A and/or HMB-45, as well as a nested positive staining pattern in the basement membrane with negative staining to the epithelial membrane antigen and Psammoma bodies.

Until recently, distinguishing between melanocytoma and melanotic schwannoma has been challenging, particularly with small specimens. However, melanotic schwannoma shows abundant deposition of reticulin material in the basement membrane around individual tumor cells; such deposition is usually absent in melanocytic tumors. In our case, we observed abundant extracellular melanin deposition with little nuclear atypia and no Psammoma bodies (Fig. 5). Therefore, the diagnosis of melanocytoma was favored owing to the depletion of reticulin and absence of Psammoma bodies. Other diagnostic approaches include the recent molecular study of GNAQ gene mutations, which permit highly accurate discrimination between leptomeningeal melanocytic lesions and melanotic schwannoma.

The survival rate among patients with an untreated meningeal melanocytoma in the central nervous system is 5 to 10 years. Among patients who undergo complete surgical resection or an incomplete resection with local radiation therapy, the survival rate might reach 100%; this percentage can drop to 46% in cases of incomplete resection. Thus, we believe that the optimal strategy for managing melanocytomas is complete resection; in case complete resection is impossible, as much of the tumor should be resected as possible, followed by postoperative local radiotherapy.

In our case, complete resection was inadvisable as the lesion was situated in a sensitive area with a high chance of vascular injury. The importance of post-resection radiotherapy depends on the degree of meningeal melanocytoma proliferation, which is controlled by Ki-67, a cellular marker that determines the growth fraction of the cell population. In our case, Ki-67 was very low (<1%). We therefore believed that the chances of recurrence or malignant transformation were low in our patient. Regardless, the possibility of local recurrence, regrowth, or malignant transformation should be considered, and follow-up neuroimaging is advised. This may help the referring clinicians to understand the progression of the disease and inform treatment decisions. In our case, follow-up imaging over 2 years showed no evidence of regrowth (Fig. 2B).

The ambiguity of this case presentation led us to consider the genetic bases for coexistent different mutations, that is, the autosomal recessive inheritance in SCD with an autosomal dominant inheritance of CCx. However, some individuals with CCx present with a sporadic type of inheritance or do not show...
an identifiable mutation of the PRKAR1A gene (a tumor suppressor gene). Researchers believe that additional and as yet unidentified genes may cause the disorder in these cases (genetic heterogeneity).\textsuperscript{[19]}

While studies have shown that the mutations of the Protein Kinase CAMP-Dependent Type I Regulatory Subunit Alpha (PRKAR1A) gene (which is located on chromosome 17), are mainly responsible for CCx, there have been several reports of mutations on the short arm of chromosome 2 being involved as well.\textsuperscript{[19]} Interestingly, there are several other diseases that have symptoms similar to those of CCx,\textsuperscript{[19]} further emphasizing the need to explore the genetic basis for diseases that coexist with CCx.

On the basis of these findings, investigators should identify additional genes that could link both conditions together, especially as SCD is composed of various genotypes. Nearly all genetic studies of SCD have concentrated on this β-globin genotype found on chromosome 11. However, other genotypes are the result of compound heterozygosity for Hbs and other hemoglobin variants, including HbC, HbE, HbD, and Hbs-β thalassemia.\textsuperscript{[20]}

### 3. Conclusions

This study is the first report of primary pigmented meningeal melanocytoma in CCx and of CCx with SCD. As meningeal melanocytomas are considered benign, complete surgical resection can be curative in most cases, and subtotal resection can also have a good outcome and prognosis. Altered gene mutations involved in variable morphogenesis and transcriptional regulation potentiate the further study of genes enriched in CCx, particularly those with coexisting SCD. The observation of coexistence of CCx and SCD in this case suggests the intriguing possibility of shared genetic contributions in both entities and provides opportunities for improved prognostic assessment and early therapeutic intervention in CCx patients.

### Acknowledgments

The authors would like to thank Dr. Abdulkader Marwh Mohammed, a pathology consultant, for her valuable assistance in obtaining the microscopic pathology images. This research did not receive any specific grant from funding agencies in the public, commercial, or not-for-profit sectors.

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### Table 1

Summary of reported cavernous sinus meningeal melanocytoma cases.

| No. | Authors, year | Age/sex | Symptoms | Location | CT attenuation | MRI signal | Management | Outcome |
|-----|---------------|---------|----------|----------|---------------|------------|------------|---------|
| 1   | Botticelli et al., 1983 | 43/F | Orbital pain, diplopia, enophthalmos, facial paresthesia, ptosis, nevus of Ota. | Left Meckel cave | Hyperdense | Not performed | Partial resection/ radiotherapy | Tumor recurrence 2 years after the operation. Second operation + radiotherapy for 10 years. |
| 2   | Faro et al., 1996 | 30/F | Severe frontal headache. | Right cavernous sinus | Hyperdense | High T1WI, low T2WI, no enhancement. | Total resection | One-month F/U and no recurrence |
| 3   | Chen et al., 1997 | 41/F | Facial numbness | Right Meckel cave | Hyperdense | High T1WI, low T2WI, no enhancement. | Total resection | One-year F/U and no recurrence |
| 4   | Kurita et al., 2000 | 29/M | Facial numbness | Right Meckel cave | Hyperdense | Low T1WI, very low T2WI, no enhancement. | Partial resection/ radiotherapy | 3-year F/U, regression/ no recurrence |
| 5   | Pan et al., 2011 | 36/M | Headache, ptosis, nevus of Ota. | Right cavernous sinus | Hyperdense | High T1WI, low T2WI, with enhancement. | Partial resection/ radiotherapy | One-year F/U and no recurrence |
| 6   | Doglietto et al., 2012 | 20/M | Diplopia, melanosis oculi | Left cavernous sinus | Hyperdense | High T1WI, low T2WI, with enhancement. | Partial resection/ chemotherapy/ radiotherapy | Seven-year F/U, recurrence, progression, lung metastasis, and death. |
| 7   | Lee et al., 2015 | 15/F | Diplopia, ptosis, facial hypotonia, and paresthesia. | Left Cavernous sinus | Hyperdense | High T1WI, low T2WI, no enhancement. | Partial resection/ gamma knife radiosurgery | 39-month F/U and no recurrence |
| 8   | Gamoh et al., 2018 | 43/M | Facial numbness and malocclusion | Left Meckel cave | Hyperdense | High T1WI, low T2WI. | Total resection/ gamma knife radiosurgery | Hyperdense Not performed Partial resection/ radiotherapy |
| 9   | Our case (2019) | 23/M | Diplopia, facial paresthesia, and headache. | Left Meckel cave | Hyperdense | High T1WI, low T2WI, peripheral rim enhancement. | Subtotal resection/ radiotherapy | 6.5 years F/U and no recurrence |

F/U = Follow-up, F = female, M = male, T1WI = T1-weighted image, T2WI = T2-weighted image.
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