Myogenic and Melanotic Differentiated Medulloblastoma: Case Report

INTRODUCTION

Medulloblastoma was first named after Bailey and Cushing in 1925 and described as an aggressive, highly malignant small cell tumor of midline cerebellum (9). Although they are rare tumors in adulthood, medulloblastoma is the most common malignant central nervous system tumor of the childhood and the most common primitive neuroectodermal tumor (PNET). Approximately 20% of these tumors are seen under the age of 19 (3,9,14,16). These neuronal differentiated embryonal tumors are usually diagnosed in the first decade of life with a male preponderance. Although the majority occurs sporadically, hereditary conditions have been associated with medulloblastoma, including Gorlin syndrome (nevoid basal cell carcinoma syndrome), Rubinstein-Taybi syndrome, ataxia telangiectasia, Turcot syndrome (e.g. glioma polyposis syndrome), Li-Fraumeni syndrome, neurofibromatosis and tuberous sclerosis (9).

Medulloblastomas arise from inferior medullary velum of the cerebellar vermis and may cause symptoms of raised intracranial pressure and truncal or appendicular ataxia. Occlusion of the fourth ventricle worsens the clinical course due to obstructive hydrocephalus. The child’s common complaints consist of the classical triad of morning headaches, vomiting and lethargy. Craniospinal spread via cerebrospinal fluid (CSF) can be seen.

We present a case diagnosed as myogenic medulloblastoma with focal areas of melanotic differentiation.

CASE REPORT

A 4-year-old boy was admitted to our clinic after headache, nausea and vomiting attacks. He described diplopia for a week. He had a gait ataxia and abducens nerve palsy on the left side in his neurological examination. Computerized tomography and gadolinium enhanced cranial magnetic...
resonance imaging (MRI) revealed a heterogeneous enhanced posterior fossa tumor rising from the fourth ventricle (Figure 1). The size of the ventricles was normal and there was no sign of hydrocephalus. There was no metastatic seeding to the spinal cord in his spinal MRI. He was operated and a gross total resection was achieved. He developed lethargy and upward gaze palsy on the fifth day of the operation. Computed tomography revealed tri-ventricular hydrocephalus. He was operated again and a ventriculo-peritoneal shunt was inserted. The post-shunting course was uneventful.

Microscopic examination revealed a poorly differentiated tumor composed of sheets of small-to-medium sized cells with focally molded, hyperchromatic nuclei and scanty cytoplasm (Figure 2). Numerous apoptotic cells and mitotic figures were observed. Some of the tumor cells contained finely granular brown pigment consistent with melanin (Figure 3). A different proportion of cells forming nests and exhibiting rhabdoid appearance with eccentric, bright eosinophilic cytoplasm were identified as well (Figure 4). These cells were immunohistochemically positive for desmin, muscle specific actin and myogenin (Figure 5A-C). GFAP, HMB-45, S-100, neurofilament, NSE, chromogranin, synaptophysin, vimentin, EMA and pancytokeratin were focally positive throughout the tumor. The final histopathologic diagnosis with these findings was medulloblastoma with myogenic and melanocytic differentiation.

He was transferred to pediatric oncology clinic for further adjuvant therapy.

DISCUSSION

Medulloblastoma is believed to arise from precursor cells of the external granular layer of the developing cerebellum and is predominantly composed of densely packed cells with round-to-oval hyperchromatic nuclei with generally abundant mitotic activity (11). They account for 40% of all posterior fossa tumors in children and they represent only 0.5–1% of all intracranial neoplasms in adults. The classic subtype represents 80% of all medulloblastoma cases. The other subtypes are, however, rather uncommon (16).

The classification proposed by the World Health Organization (WHO) in 2000 describes subtypes of medulloblastoma as classic, desmoplastic, large cell, medulomyoblastoma and melanotic medulloblastoma (14-16). The classification was revised in 2007 and there have been changes in medulloblastoma classification. The new system defines subtypes of medulloblastoma as desmoplastic/nodular medulloblastoma, medulloblastoma with extensive nodularity, anaplastic medulloblastoma and large cell medulloblastoma.

The term of medulomyoblastoma was first used in 1933 by Marinesco and Goldstein and, up to now approximately 50 cases have been reported in the literature (12,14). They originate from the vermis and demonstrate similar clinical behavior to classical medulloblastomas. The management protocol of this subtype is similar to the classical ones. Histologically these medulloblastomas include areas focally exhibiting rhabdomyoblastic differentiation which can be demonstrated with immunohistochemical stains for myogenin, desmin and actin. Primitive sarcomeres and Z-bands might also be observed in electron microscopy (14).

The 2000 WHO classification named medulloblastoma with myogenic differentiation as medulomyoblastoma. However, since its clinical and genetic features are similar to those of other medulloblastoma, this lesion is no longer considered as a distinct entity. The authorities suggest the term "medulloblastoma with myogenic differentiation" for any variant containing focal rhabdomyoblastic elements with immunoreactivity to desmin, myoglobin and fast myosin (11).
Melanotic medulloblastoma was first described in 1962 by Fowler and Simpson as 'pigmented melanin forming tumor of cerebellum'. Since then approximately 12 cases have been reported in literature (14,15).
The fate of the term melanotic medulloblastoma is the same. The new classification no longer recognizes this subtype as a distinct entity and instead suggests the term “medulloblastoma with melanotic differentiation” for tumors that appear undifferentiated or epithelial, with formation of tubules or papillae and expressing S-100 protein (11). Tumors possessing both myogenic and melanotic differentiation was reported only in six papers up to date (2,7,8,12,13,14). The mean age of those were 4. They usually arise from cerebellar vermis or the 4th ventricle (14).

Banerjee and Duinkere have reported first examples of these tumors in 1973 and 1981, respectively (2,7). Sakata et al have presented a 6 year old boy with a classical medulloblastoma demonstrating melanotic and myogenic differentiation as well as neuronal and astrocytic differentation. They conclude that their case supports the multipotent origin of neuroblastomas (15). Kalimo et al have reported a 5 year old girl with a PNET demonstrating neuronal, astrocytic, melanotic and myogenic differentiation. The child has suffered from rapid recurrence and died on the 6th month of primary surgery. In their conclusion they site the most widely accepted theory in oncogenesis of medulloblastomas which is the multipotient origin from the neural crest (8). Polydorides et al have reported a large cell medulloblastoma with myogenic and melanotic differentiation in a 2-year-old boy. The have supported their diagnosis with genetic investigation. They concluded their discussion in agreement with the updated WHO classification that these tumors are histologic patterns of medulloblastoma with similar prognostic features (14). Nozza et al. have described a cerebellar medullomyoblastoma with melanotic tubular structures in a 2 year old boy. They discussed the possibility of a teratomatous origin of the tumor but concluded in favour of the theory that undifferentiated neuroepithelial cells differentiate along myogenic lines (13).

Mehta et al have reported a 3-year-old girl presenting with insidious dysarthria, urinary incontinence and walking difficulty. The symptoms were due to a pigmented myomedulloblastoma. The authors emphasize the importance of differential diagnosis including pediatric tumors demonstrating rhabdoid differentiation and teratomas. They conclude that besides the best available treatment modalities the prognosis is poor (12).

The histogenesis of medulloblastomas with myogenic differentiation is controversial. According to a view, these tumors originate form pluripotent cells of the fetal cerebellar meninges (12). Another view is that myoblastic cells originate from neoplastic transformation of endothelial cells based on the fact that these muscle fibers were located around blood vessels and showed mitotic figures (1,10).

One important entity is medullamyoblastomas demonstrating teratomatous elements. Although some believe that these tumors are actual teratomas, absence of endodermal elements and mesenchymal elements other than muscle fibers excludes this option (5,15).

For melanotic medulloblastomas, the histogenesis is even more controversial. Some authors consider these tumors as variants of melanotic neuroectodermal tumors of infancy. However, these tumors are not as aggressive as medulloblastomas (6). Another suggestion is that melanotic medulloblastomas are hybrid tumors: pigmented cells are from the neural crest while the non-pigmented cells are from the neuroectoderm, and tumors combining muscle cells and pigmented epithelial elements have links to teratomas (4,14).

The 5-year survival rate in medulloblastoma is between 80% and 85%. Although there are reports stating the prognosis of nodular medulloblastomas is more favorable, the prognostic reliability for histological subtypes is controversial (17).

CONCLUSION

Understanding the underlying mechanism of oncogenesis will serve as leverages for future treatment modalities. However, with the options of today’s medicine we can not make a difference in treating various subtypes of medulloblastomas. In our opinion, this case together with the previously reported 6 cases of medulloblastoma with melanotic and myogenic differentiation supports the theory that medulloblastomas originate form pluripotent stem cells.

REFERENCES

Borcek AO, et al: Myogenic and Melanotic Differentiated Medulloblastoma


