Prevalence of primary CNS tumors in a diagnostic setup in Nairobi, Kenya

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Abstract
Introduction: Diagnosis of CNS tumors is often difficult for the general pathologist due to its rarity. However, high morbidity and mortality makes this the dreaded form of cancer and diagnosis requires precision to differentiate the various forms including secondaries.

Materials and Methods: Study was conducted in an independent laboratory over a period of 6 months in Nairobi, Kenya and all primary CNS neoplasms during the study period were included in the study.

Result: Amongst 42 lesions involving the CNS, 34 primary CNS neoplasms were identified. It was more common in females with a mean age of 49.41 years. Meningiomas were the most common type of CNS lesions followed by astrocytic tumors.

Conclusion: Incidence of meningiomas is high in the African population. Light microscopy is essential for initial diagnosis, however, immunohistochemistry, ultrastructural studies including molecular studies are essential to predict further management.

Keywords: Central nervous system, Neoplasms, Meningioma, Astrocytoma, Kenya.

Introduction
Tumors of CNS are rare and account for less than 1–2% of all malignancies among the cancer patients. The Globcan study shows 64.9% of cancer deaths occur in less developed regions of the world and the CNS tumors with their high morbidity and mortality makes them the most dreaded form of cancer. CNS tumors diagnosis is often difficult as general pathologist lack experience and non-neoplastic processes mimic tumors and primary needs to be differentiated from secondaries.¹,² The most prevalent primary CNS tumors in adults are astrocytic tumors followed by meningiomas. Numerous classifications have been put forth to classify tumors since 1979, which was based on light microscopy. The recent WHO classification of CNS tumors (2016) has moved beyond the light microscopy, immunohistochemistry and ultrastructural studies into the realm of incorporating molecular parameters into the classification of CNS tumors, thereby integrating both genotypic and phenotypic parameters adding a level of objectivity. Overlapping genotype like IDH mutation in both oligodendroglioma and diffuse astrocytoma (multiple susceptibility foci for gliomas seen) makes a compelling example for the new classification.³⁴

Like earlier classification, the 2016 Classification acknowledges WHO grade tumour determinations is still based on histology and genotyping each category creates new diagnostic challenges in different setups and hence integrating both genotype and phenotype is still necessary.⁵ The major restructuring has been done in diffuse gliomas, medulloblastomas, embryonal tumors and ependymoma with incorporation of genetically defined entities.⁶⁷ In meningiomas, there were no major revisions except for introduction of brain invasion as criterion for diagnosis of atypical meningiomas, WHO grade II. Brain invasion is now used for grading, rather than staging and brain invasion alone suffices for a diagnosis of atypical meningioma similar to a histological criterion of mitotic count of 4 or more.⁸⁹

Majority of setups all over the world is devoid of molecular or IHC studies and is dependent on histology. The present study was conducted to identify the prevalence of primary CNS tumors based on light microscopy findings in an independent laboratory setup and this could be one of the few studies from this part of this region in published literature.

Materials and Methods
This retrospective study was conducted over a period of 6 months (March 2016 to September 2016) in an independent laboratory in Nairobi, Kenya. Data of all primary intracranial tumors including patient’s age, sex and histopathology was collected. Inclusion criteria included all primary CNS neoplasms including pituitary neoplasm’s above 19 years age group. Exclusion criteria included CNS tumors in pediatric age groups, incomplete data, and inconclusive diagnosis, metastatic and non-neoplastic conditions of the CNS.

Specimens were grossed and processed according to standard procedures. All slides were stained with Haematoxylin and Eosin stain and reporting was done.

Statistical Analysis
Descriptive statistics – data was calculated as percentages.
Result

During this period, a total of 1726 specimens were received from various departments and among these 42 specimens involved the Central nervous system. Non-neoplastic lesions formed 8 of the 42 cases. On gross examination, specimen size varied from <0.5 cm to upto 4x4 cm size, particularly with larger tumors. Multiple bits were taken and processed and slides were stained with H&E. Amongst non-neoplastic lesions, 3 cases of tuberculoma, 2 cases of arachnoid cyst, 1 case each of epidermoid cyst, pyogenic abscess and infarct was noted.

The mean age of tumors was 49.41 years with an age range of 30-72 years. Tumors were more common during the 5th decade. Females composed 20/34 cases with a mean age of 48.85 years with an age range of 30-69 years. Amongst males 14/34 cases was seen with the mean age of 50.21 years, with a mean range of 31-72 years.

Meningiomas were the most common type of lesions followed by astrocytic tumors. Average age group among 15 cases of meningiomas was 51.13 years with a minimum age of 31 and maximum age of 72 years. Females formed the predominant cases of meningiomas (11/15). In both males and females, meningotheial meningiomas were the commonest findings. Other variants of meningioma included psammomatous meningioma (Fig. 1), secretory meningioma (Fig. 2), atypical meningioma and transitional meningioma.

Astrocytic tumors (8/34) were the next most common lesions. Average age was 56.75 years with an age range of 47-69 years. Glioblastoma multiforme (5/8) (Fig. 3, 4) was the most common type. Astrocytic tumors were again more common in females (5/8).

Other tumors found in the study were pituitary macroadenoma, Dysembryoplastic neuroectodermal tumor (Fig. 5), neurofibroma (Fig. 6), schwannoma and plasmacytoma.

Fig. 1: Psammomatous meningioma showing psammoma bodies- 10x photomicrograph HxE

Fig. 2: Secretory meningioma showing eosionophilic secretions Photomicrograph 10x HxE

Fig. 3: Glioblastoma multiforme showing palisading necrosis Photomicrograph 10x HxE

Fig. 4: Glioblastoma multiforme showing vascular proliferation and anaplasia Photomicrograph 10x HxE
Fig. 5: Dysembryoblastic neuroectodermal tumor Photomicrograph 10x H&E

Fig. 6: Neurofibroma showing spindle cells with wavy nuclei Photomicrograph 10x H&E

Discussion

Studies about the CNS tumors are rare in the African population. Gliomas are more common followed by meningiomas similar to various studies. In the neurology centers in the African continent, CNS tumors incidence vary from 4.4% to 12% of all admissions. All over, malignant CNS tumors are more common in the pediatric age group compared to adults and they are on the rise, probably due to better detection techniques or environmental exposures. However, in developing countries like Kenya, the high gap in neurosurgical care compared to developed countries, adverse socioeconomic factors, cultural beliefs and most importantly, an alarming shortage of neurooncologists contribute to a delayed presentation and an adverse prognosis. Malignant tumors are more common in males, while benign tumors are more common in females.

WHO 2016 CNS tumors classification is based on integrated diagnoses and along with genotyping, histopathology is necessary for classification. Genotype alone cannot be useful as mutations can overlap, for eg: oligodendroglia and diffuse astrocytoma both have IDH mutation. Additionally, tumor grading and tumors with overlapping phenotypic and genotypic criteria require histopathological confirmation too. This 2016 WHO CNS tumors classification is seen as a stepping stone towards a major incorporation of molecular data in future classifications, which are more treatment oriented thereby further removing wastebasket categories. This is expected to improve and facilitate improvements in life of patients with brain tumors. Most common tumor in the study was meningioma followed by astrocytomas. Meningiomas are tumors arising from the arachnoid “cap” cells of the arachnoid villi and they are the most common benign primary brain tumors affecting adults. They are mostly benign, may occur intracranially or spinal and are usually clinically silent, but have a potentially high impact on morbidity of the patients. In our study, meningiomas formed 15/34 primary brain tumors, and it was more common in females. Other studies have shown similar data with occasional study showing astrocytomas to be more common. Research has shown multiple genetic and environmental risk factors for meningiomas. Genetically, it is known to be associated with neurofibromatosis type II and multiple endocrine neoplasia type I. Environmental risk factors principally ionizing radiation. Inconclusive data exists on use of mobile phones, hair dyes, petrochemicals, allergens, smoking, hormonal uses, etc. Meningiomas carry a better prognosis in adults compared to pediatric age group. No cases of recurrent meningiomas were noted in the study, possibly due to short duration of study or total tumor resection leading to complete cure because of the benign nature of the lesion. These findings are consistent with various studies indicating high incidence of intracranial meningiomas in Africans.

Gliomas represent 81% of the primary malignant CNS tumors. Glioblastoma is supposed to be the most common histology and our study also shows glioblastomas to be the most common findings. Syndromes known to be associated with gliomas are neurofibromatosis type I and II, Lynch syndrome, tuberous sclerosis, etc. Other environmental risk factors are similar to meningioma.

Nerve sheath tumors were very few in our study. Other studies also have shown similar findings. Comparison of primary CNS tumors in various studies in relation to our study is as shown in table 1.
Table 1: Comparison of frequently reported CNS tumors in various studies

<table>
<thead>
<tr>
<th>Authors</th>
<th>Year</th>
<th>Country</th>
<th>Most frequently reported variant</th>
</tr>
</thead>
<tbody>
<tr>
<td>Jaiswal J et al(1)</td>
<td>2016</td>
<td>India</td>
<td>Meningiomas (23.2%)</td>
</tr>
<tr>
<td>Ostrom QT et al(22)</td>
<td>2013</td>
<td>United States of America</td>
<td>Meningioma (36%)</td>
</tr>
<tr>
<td>Sumathi V et al(8)</td>
<td>2016</td>
<td>India</td>
<td>Astrocytomas – 49%</td>
</tr>
<tr>
<td>Pidakala P et al(2)</td>
<td>2016</td>
<td>India</td>
<td>Meningiomas – 18.1%</td>
</tr>
<tr>
<td>Ibebuike K et al(18)</td>
<td>2013</td>
<td>Johanannesberg, South Africa</td>
<td>Meningiomas – 31.8%</td>
</tr>
<tr>
<td>Hema NA et al(21)</td>
<td>2016</td>
<td>India</td>
<td>Gliomas – 44%</td>
</tr>
<tr>
<td>Odowu OE(11)</td>
<td>2007</td>
<td>Nigeria</td>
<td>Meningiomas – 30%</td>
</tr>
<tr>
<td>Kuipers SE et al(24)</td>
<td>2013</td>
<td>Suriname, South America</td>
<td>Meningiomas – 27.1%</td>
</tr>
<tr>
<td>Present Study</td>
<td>2016</td>
<td>Kenya</td>
<td>Meningiomas – 44.11%</td>
</tr>
</tbody>
</table>

Conclusion

The incidence of CNS tumors is comparatively rare compared to other sites. High incidence is seen in 4th and 5th decade with a slight female predominance. Most common primary CNS tumors seen were meningiomas followed by astrocytomas. We believe, further epidemiological studies are essential in this part of the world to provide further insight into this high incidence and possible etiological factors.

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References


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