Epidemiologic Evaluation of Central Nervous System Sarcomas in Iran: A 2009-2014 Survey

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Introduction

Being derived from Greek word sarx meaning flesh, sarcomas originates from transformed mesenchymal cells with numerous subtypes and various distribution pattern and prognosis [1]. Mentioned versatility coupled with its rarity, are two main factors responsible for restricted reports of sarcomas description throughout history, which account for current limited knowledge on appropriate management of different types of sarcomas [2,3]. The American Cancer Society’s estimation of new bone and soft tissue sarcomas are 3500 and 12750 cases in the United States by the end of 2019 [4]. Considering the total estimate of new diagnosed cancer is 1762450 cases, bone and soft tissue sarcomas constitute about 0.2 and 0.7% of newly diagnosed malignancies respectively, making total cases of newly diagnosed malignancies. Although insignificant in number, these tumors may complicate patient’s overall health status compared with other malignancies, because of their numerous subtypes and variable features which may pose an everlasting challenge to both clinical and surgical treatment strategies. Existence of these complications and a relatively small available data regarding Central Nervous System’s (CNS) sarcomas, encouraged authors to conduct a retrospective study in a 6-year period to evaluate the specific epidemiological features of CNS sarcomas, including brain, spinal cord and meningeal layer sarcomas extracted from Iranian National Cancer Registry (INCR). Our study revealed CNS sarcoma’s trend to affect males more than females, affecting them most in their midlife and a higher prevalence of brain involvement compared to spinal cord and meningeal layers. Also, we have provided detailed morphological features of the tumors, as well as patient’s geographical distribution, with Northern parts of Iran have the lowest incidence rate (10.06%). Moreover, our analysis of crude rate revealed lower Age Specific Incidence Rate (ASIR) of CNS sarcomas than expected number of cases compared to standard world population by World Health Organization (WHO’s) ASIR as 0.035 person per 100000-years. Authors believe this report of epidemiological assessment of CNS sarcomas in Iran could act as a foundation for better understanding the underlying pathophysiological mechanism as it is the first survey in developing countries and selection of optimal treatment strategies based on improved understanding of these neoplasms.

Keywords: Sarcoma; Soft tissue sarcoma; Central nervous system; Epidemiologic study

Abstract

Sarcomas rise from mesenchymal layers and may transform to variable malignant lesions in multiple organs systems. Although they are mostly localized, many of them have predilection of distant sites away from their primary origins and subsequent metastasis to different anatomical locations of body. Soft Tissue Sarcomas (STSs) represent one percent of adult and about 7% of pediatric malignancies. Although insignificant in number, these tumors may complicate patient’s overall health status compared with other malignancies, because of their numerous subtypes and variable features which may pose an everlasting challenge to both clinical and surgical treatment strategies. Existence of these complications and a relatively small available data regarding Central Nervous System’s (CNS) sarcomas, encouraged authors to conduct a retrospective study in a 6-year period to evaluate the specific epidemiological features of CNS sarcomas, including brain, spinal cord and meningeal layer sarcomas extracted from Iranian National Cancer Registry (INCR). Our study revealed CNS sarcoma’s trend to affect males more than females, affecting them most in their midlife and a higher prevalence of brain involvement compared to spinal cord and meningeal layers. Also, we have provided detailed morphological features of the tumors, as well as patient’s geographical distribution, with Northern parts of Iran have the lowest incidence rate (10.06%). Moreover, our analysis of crude rate revealed lower Age Specific Incidence Rate (ASIR) of CNS sarcomas than expected number of cases compared to standard world population by World Health Organization (WHO’s) ASIR as 0.035 person per 100000-years. Authors believe this report of epidemiological assessment of CNS sarcomas in Iran could act as a foundation for better understanding the underlying pathophysiological mechanism as it is the first survey in developing countries and selection of optimal treatment strategies based on improved understanding of these neoplasms.

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Methods

In this retrospective study, authors have analyzed patients with confirmed histopathological study of CNS sarcomas in Iranian population. Patients with CNS sarcomas were identified from Iranian associated sarcomas in young individuals [9,10]. Undifferentiated sarcoma, fibrosarcoma and malignant fibrous histiocytoma were among the most commonly found subtypes of CNS sarcomas in the past decade [11]. In addition to previous reports of incidence rate of CNS sarcoma of 3 per 10 million person-years, primary Intracranial Sarcomas (IS) which were first described in 1929, have incidence rate from 0.1 to 4.3% based on multiple types of studies with variable definition of primary IS [12,13]. Inclusion of reticulum cell sarcoma, circumscribed sarcoma of cerebellum and hemangiopericytoma and report’s inconsistency in unified definition of primary IS have contributed to this high value of incidence despite its rarity [14]. Since epidemiologic and descriptive studies are the cornerstones of enhanced understanding of these group of malignancies, authors decided to describe the epidemiologic aspect of CNS sarcomas and the prevalence of these challenging tumors in brain and spinal column and their subtypes in a 6-year period in Iran from 2009 to 2014.
National Cancer Registry (INCR) from April 2009 to December 2014. Also, authors extracted population data of from Iranian National Cancer Registry. Patient’s demographic data including age, sex, living environment and city, primary tumor location and histopathological subtype based on World Health Organization (WHO) classification system of sarcoma with confirmed diagnosis, were gathered and recorded. Patient’s specific tumor data documented in INCR have been filed based on histopathological and radiological confirmation, with roughly 16700 documented sarcoma patients were analyzed in the same mentioned 6-year period. Due to recent changes of STSs classification system and ongoing updates, authors have reported both macroscopic and microscopic tumor description as tumor’s morphology and its classification, respectively. Since some tumors have higher incident rate in specific geographical areas, we have analyzed the geographical distribution pattern of our patients to find a possible relevant “hot-zone” for CNS sarcomas in Iran, given that it may lead to extended research on potential environmental hazards and carcinogens. Also, authors performed statistical analysis using SPSS program version 22 to perform Age Specific Incidence Rate (ASIR) and its annual trend.

Results

In this study, authors have thoroughly examined the INCR’s information in a period between 2009 to 2014, and analyzed 298 patients with confirmed diagnosis of CNS sarcomas, either of brain (primary IS) or spinal cord, out of the 16700 sarcoma patients. Our analysis revealed 180 males and 118 females with male to female ratio of 1.52 and a periodic prevalence of 1.78%. Specific details of CNS sarcoma prevalence among both sexes is described in Table 1.

Also, authors have classified patients based on their presenting age on 3 main groups as 0-14 years as age range group I, 15-64 years as age range group II and 65-100 years as age range group III. Based on the results, 31 patients (10.4%) were categorized as group I, 227 patients (76.17%) as group II and 40 patients (13.42%) as group III. Detailed age specifications are demonstrated on Table 2.

Authors deep further in statistical analysis and performed Age Specific Incidence Rate (ASIR) analysis of CNS sarcomas in complete study period demonstrated in Table 3. Our analysis revealed ASIR of CNS sarcoma as 0.396 per 100000 person years compared to 0.431 per 100000 person years of expected number cases based on Standard World Population from WHO, that indicate lower ASIR of our study compared to the latest with a new age range classification, as children and adolescent (0-19 years), young adults (20-34 years), old adults (35-59 years) and elderly (more than 65 years) based on WHO definitions (Table 4, Figure 1).

Given annual trend above state that elderly population are at the highest risk of CNS sarcoma development, while children and adolescents stands at the opposite edge with the lowest probability. On the other hand, clinical and surgical evaluations revealed the anatomical location of CNS sarcomas, reported as located in brain as 184 cases (61.7%), followed by 101 cases (33.8%) in the spine and 13 cases (4.3%) located primarily on their meningeal layers. Authors have classified the studied lesions from histopathological point of view and have provided their grade based on World Health Organization (WHO)'s sarcoma scoring system (ICD-O). Tissue examination of CNS sarcomas revealed 10 cases (3.3%) as grade 1 or adipocytic tumors, 14 cases (4.6%), as grade 2 or fibroblastic/myofibroblastic tumors, 15 cases as grade 3 or fibrohistiocytic tumors (5%), 11 cases (3.6%) as grade 4 or smooth muscle tumors and other 248 cases (83.2%) as grade 9 or tumors of uncertain differentiation.

| Year (CNS sarcoma patients) | Brain | Spine |
|---------------------------|-------|-------|
|                           | Male (percentage) | Female (percentage) | M:F ratio | Male (percentage) | Female (percentage) | M:F ratio |
| 2009 (8)                  | 6 (75%)       | 1 (12.5%)    | 6         | 1 (12.5%)       | 0                 | 1         |
| 2010 (20)                 | 8 (40%)       | 10 (50%)     | 0.8       | 1 (5%)          | 1 (5%)            | 1         |
| 2011 (83)                 | 38 (45.7%)    | 25 (30.1%)   | 1.52      | 12 (14.4%)      | 8 (9.6%)          | 1.5       |
| 2012 (72)                 | 25 (34.7%)    | 14 (19.4%)   | 1.78      | 20 (27.7%)      | 13 (18%)          | 1.53      |
| 2013 (65)                 | 27 (41.5%)    | 16 (24.6%)   | 1.68      | 16 (24.6%)      | 6 (9.2%)          | 2.66      |
| 2014 (50)                 | 18 (36%)      | 15 (30%)     | 1.2       | 6 (12%)         | 11 (22%)          | 0.54      |
| Total (298)               | 122 (40.9%)   | 81 (27.1%)   | 1.5       | 56 (18.7%)      | 39 (13%)          | 1.43      |

CNS: Central Nervous System, M: Male, F: Female.
Table 5 contains details of morphological features of excised and studied lesions of our study sample. Also, our analysis regarding most encountered histopathological subtype of CNS sarcoma in each age group category which defined above, age group 1 (0-14 years) mainly consisted of sarcoma with mostly unspecified grade (22.5%), followed by desmoplastic small round cell tumor with unspecified grade and gliosarcoma with mostly unspecified grade (12.9% each). Also, gliosarcoma with mostly unspecified grade ranked first (18.9%) in age group 2 (15-64 years), followed by chordoma with unspecified grade (11.8%) and desmoplastic small round cell tumor with mostly unspecified grade (11%). At the latest age group as group 3 (65-100 years), gliosarcoma with mostly unspecified grade (40%) consist the major CNS sarcoma type, followed by hemangiopericytoma malignant with mostly grade 3 (17.5%) and chordoma with unspecified grade (10%).

According to previous statements of possibility of geographical relevance to development of STSs, especially CNS sarcomas, we have assessed the distribution of our patients based on their living area and country’s territories on Table 6.

**Discussion**

Based on histopathological classifications, subtypes of bone and soft tissue sarcomas are osteosarcoma, chondroblastoma, poorly differentiated round or spindle cell tumors, chordoma and neural sheet tumors [15]. Also, soft tissue sarcoma subtypes include liposarcoma, fibrosarcoma, leiomyosarcoma, rhabdomyosarcoma, angiosarcoma, gastrointestinal stromal tumor and undifferentiated sarcomas, along with other less frequent subtypes [16]. Although no exact mechanism or etiology is found to be responsible for development of sarcomas, several environmental and genetic mutations risk factors are considered to play important role in transformation of mesenchymal cells to different sarcoma subtypes. To give an instance, chromosomal translocation of chromosome 11 and chromosome 22 which results in EWS gene is associated with Ewing sarcoma [17]. Also, duplication of a part of chromosome 12 and resultant CDK4 gene and MDM2 gene
play a significant role in development liposarcoma [18]. Furthermore, CSF1 gene which follows chromosomal translocation between chromosome 1 and chromosome 2, is related to increased risk of development of giant cell tumors of soft tissue [19]. Primary CNS sarcomas are primitive non-meningothelial tumors which represent about 0.2% of all intracranial malignancies, with equal trend to affect both sexes at any age [20-22]. Despite the trivial risk, sarcoma’s burden on affected patients demands our fortified understanding of its natural history and pathogenesis to reach acceptable standards towards appropriate management of sarcomas individually. As stated above, mesenchymal origin of sarcomas provides emersion of wide variety of subtypes of both bone and soft tissue sarcomas [23]. Although rare, wide variety of presentation and subtypes have made Soft Tissue Sarcomas (STSs) a challenging malignancy with somehow unknown etiologic background.

In a report on 13 Children, Adolescent and Young Adults (CAYA) from 1990 to 2015, it is found that the affected patients with primary IS have mean age of 7 years, while another report on CAYA patient groups from 1990 to 2001, revealed the mean age of diagnosis of primary IS as 16.9 years [24,25]. In addition to the mentioned data of former report from United States of mean age of diagnosis, this 25-year observation stated unclassified sarcoma, followed by chondrosarcoma and rhabdomyosarcoma [24]. Also, Benesch et al conducted a cooperative study in Austria from 1988 to 2009 on 19 CAYA age group cases diagnosed with CNS sarcomas with mean age of 4.8 and found 14 patients with intracranial sarcoma followed by the rest as spinal sarcomas in Canada [26]. Given the above results of mentioned researches, this study includes a considerable greater population of patients with age group 1 (0-14) which is comparable to CAYA age group (0-19), with similar first major subtype as unspecified sarcoma (22%) to other two studies, followed by desmoplastic small round cell and gliosarcoma (12.9% each) which demonstrated different sequence compared to previous surveys. Away from CAYA age group, Tihan et al reported a 20-year survey on 43 cases diagnosed with primary CNS sarcomas with mean age of 40.3 years (3-75 years). In this study which was performed in United States and published in 2007, 16 cases were diagnosed with hemangiopericytoma, followed by 15 cases with chondrosarcoma and 3 with solitary fibrous tumor [27]. In addition, another 40-year report on 18 cases aged between 3-63 years (mean age 28) affected with primary IS revealed 15 cases with cerebral sarcoma, 2 with cerebellum involvement and one with spinal sarcoma [28]. Their histopathological evaluation exhibit fibrosarcoma, malignant fibrous histiocytoma and undifferentiated sarcoma as the most common tumor types in descending manner. Although Oliveira et al did not report detailed subtypes based on patient’s age group, regarding their mean age of 28 years, our age group 2 (15-64) most common sarcoma types exhibit different sequence, as gliosarcoma (18.9%) chordoma 11.8% and desmoplastic small round cell tumor (11%). Although former surveys that were conducted on developed countries, demonstrates different sequence of common CNS sarcoma types based on their histopathologic features, authors believe that our

| Tumor type              | Number | Tumor type              | Number | Tumor type              | Number |
|-------------------------|--------|-------------------------|--------|-------------------------|--------|
| Gliosarcoma             | 63     | Small cell sarcoma, round cell | 4      | Alveolar soft part sarcoma | 1      |
| Desmoplastic small round cell tumor | 33     | Rhabdomyosarcoma         | 4      | Liposarcoma NOS          | 1      |
| Chordoma                | 31     | Leiomyosarcoma           | 3      | Myxosarcoma              | 1      |
| MPNST                   | 23     | Giant cell sarcoma       | 3      | Fibrosarcoma             | 1      |
| Hemangiopericytoma malignant | 22     | Malignant fibrous histiocytoma | 3      | Endometrial stromal sarcoma | 1      |
| Neurilemoma sarcoma      | 13     | MPNST with rhabdomyoblastic differentiation | 2      | Mixed malignant tumor    | 1      |
| Malignant tumor spindle cell type | 12     | Malignant histiocytosis   | 2      | Myxoid liposarcoma       | 1      |
| Ewing’s sarcoma          | 10     | Mixed type rhabdomyosarcoma | 2      | Embryonal rhabdomyosarcoma | 1      |
| Spindle cells sarcoma    | 9      | Synovial sarcoma         | 2      | Glionus tumor malignant  | 1      |
| Meningeal sarcomatosis   | 9      | Cerebellar sarcoma       | 2      | Fibrous mesotheloma malignant, Sarcomatoid mesotheloma | 1      |
| Chondrosarcoma           | 5      | Osteosarcoma             | 2      |                         |        |
| Malignant rhabdoid tumor | 5      | Epitheloid sarcoma       | 1      |                         |        |
| Myeloid sarcoma          | 5      | Mast cell sarcoma        | 1      |                         |        |
| Schwannoma               | 5      | Paraganglioma malignant  | 1      |                         |        |
| Chondroid chordoma       | 5      | Synovial sarcoma biphasic| 1      |                         |        |
| Sarcoma                  | 4      | Kaposi sarcoma           | 1      |                         |        |

NOS: Not Otherwise Specified; MPNST: Malignant Peripheral Nerve Sheath Tumor.

| Region of Iran          | Number of affected patients (Percentage) |
|-------------------------|----------------------------------------|
| Central parts           | 115 (38.59%)                           |
| Southern parts          | 58 (19.46%)                            |
| Eastern parts           | 53 (17.78%)                            |
| Western parts           | 42 (14.09%)                            |
| Northern parts          | 30 (10.06%)                            |
| Total                   | 298 (100%)                             |

Table 5: Morphological features of CNS sarcomas.

Table 6: CNS sarcomas distribution based on country’s territories.
results may act as a considerable representative of epidemiological and histopathological features of developing countries with different values compared to developed ones. Despite scant available data on tumorigenesis of primary IS, studies suggest activation of receptor tyrosine kinase and angiogenesis pathways in most of the pediatric ISs [29,30]. Also, overexpression of platelet-derived growth factor receptor was found to be associated with osteosarcoma, rhabdomyosarcoma and Ewing sarcoma [31]. Beside special molecular and genetic processes responsible for development of this cancer, sarcomas are linked to multiple environmental offenders, such as ionizing radiation, alkylating agents and vinyl chloride and arsenic exposure. Moreover, sarcomas are associated with certain syndromes, including Li-fraumeni and neurofibromatosis type 1 [32,33]. Regarding primary IS, similar to other space occupying lesions, its clinical and neurological manifestation is consistent with tumor’s location. Despite its natural tendency to originate from various cells in different locations, existing evidence suggest primary IS’s predilection towards supra-tentorial regions, mostly affecting temporal and parietal lobes [34]. Neurological symptoms range from asymptomatic and mild headache, to significant neurological deficit and altered level of consciousness. However, multiple histopathological subtypes were found in previous conducted studies, being mostly reported as undifferentiated sarcomas, followed by chordoma, chondrosarcoma and rhabdomyosarcoma, as well as other less frequent subtypes [35-37]. Our analysis revealed obvious male to female dominance regarding prevalence of CNS sarcomas, with three quarters of the patients being in second age group category from 15-64 years (76.17%), followed by age group III (13.42%) and I (10.4%), showing higher tendency of CNS sarcomas to affect people away from their extremes of their ages. Our ASIR analysis revealed lower ASIR of CNS Sarcomas than expected number of cases compared to standard world population by WHO’s ASIR as 0.035 person per 100000-years. Also, our evidence suggests CNS sarcoma’s trend to affect brain roughly 2 times more than spinal cord and isolated meningeal layers of less than 5%. Also, WHO classification of CNS sarcomas of our studied patients revealed significant grade 9 or tumors of uncertain differentiation compared to other subtypes. Gliosarcoma represents the most frequent detected CNS sarcoma, followed by desmoplastic small round cell tumor, chordoma, malignant peripheral nerve sheet tumor and hemangiopericytoma malignant as top 5 lesions based on frequency. Moreover, our detailed report of patient’s geographical distribution pattern may provide potential fields for environmental and public health activist to use this data as a primary reference in their future plans.

Conclusion

Coherent with various subtypes, multiple medical and surgical approaches have been suggested for patients diagnosed with primary IS. However, due to lack of unified viewpoint of its natural history and existence of criteria for selection of appropriate management approach, insufficiency of an optimal strategy towards primary IS, is perceived. This is the first report of epidemiological assessment of CNS sarcomas in Iran, which can act as a foundation for better understanding the underlying pathophysiological mechanism and selection of optimal treatment strategies based on improved understanding of these neoplasms.

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