RESEARCH ARTICLE

Paediatric Retinoblastoma in India: Evidence from the National Cancer Registry Programme

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Abstract

**Background:** Globally, retinoblastoma is the most common primary intraocular malignancy occurring in children. This paper documents the recent incidence rates of retinoblastoma by age and sex groups from the Population Based Cancer Registries (PBCRs) of Bangalore, Mumbai, Chennai, Delhi and Kolkata using the data from the National Cancer Registry Programme. **Materials and Methods:** Relative proportions, sex ratio, method of diagnosis, and incidence rates (crude and age standardized) for each PBCR and pooled rates of the five PBCRs were calculated for the years 2005/06 to 2009/10. Standard errors and 95% confidence limits of ASIRs by sex group in each PBCR were calculated using the Poisson distribution. Standardised rate ratios of ASIR by sex group and rate ratios at risk were also calculated. **Results:** The maximum retinoblastoma cases were in the 0-4 age group, accounting for 78% (females) and 81% (males) of pooled cases from five PBCRs. The pooled crude incidence rate in the 0-14 age group was 3.5 and the pooled ASIR was 4.4 per million. The pooled ASIR in the 0-4, 5-9 and 10-14 age group were 9.6, 2.0 and 0.1 respectively. The M/F ratio in Chennai (1.9) and Bangalore PBCRs (2.0) was much higher than the other PBCRs. Among the PBCRs, the highest incidence rate in 0-4 age group was found in males in Chennai (21.7 per million), and females in Kolkata (18.9 per million). There was a distinct variation in incidence rates in the PBCRs in different geographic regions of India.

**Keywords:** Retinoblastoma-population based cancer registry-incidence rate

Introduction

Globally, retinoblastoma is the most common primary intraocular malignancy occurring in children. The incidence of retinoblastoma reported in the Cancer Incidence in Five continents Volume X is lower in the developed nations as compared to the developing countries, and the age standardized incidence rates (ASIR) ranges from 1 per 1,000,000 population in 0 -14 years male and female children in many parts of the world to 8 per 1,000,000 in Blantyre, Malawi in Africa (Forman et al., 2013). There have been earlier studies that have analysed the incidence of retinoblastoma from specific regions in India, that pertain to the last two decades (Nandakumar et al., 1996; Yeole et al., 2002; Swaminathan et al., 2008). Retinoblastoma is a rare disease, and calculation of incidence rates requires reliable data from a defined population at risk. Using the data from the National Cancer Registry Programme (NCRP), India this paper explains the recent incidence rates of retinoblastoma by age and sex groups from the Population Based Cancer Registries (PBCRs) of Bangalore, Mumbai, Chennai, Delhi and Kolkata.

Materials and Methods

The NCRP in India was established in 1982 and has expanded to include 29 PBCRs till date. The NCRP

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collects information on incident cases of all cancers, diagnosed among residents (resident for at least a year or in children less than a year old, whose parents were resident for at least a year) in a defined population, from multiple sources and records it through a software data entry module. Information on deaths due to cancer are also collected from hospitals and the Civil registration system and matched with incident cases. Detection of duplicate registrations and internal validity checks are carried out by the software programme of the NCRP on an ongoing basis. The cancers are classified using ICD-O-3 (International Classification of diseases-Oncology, 3rd edition) morphology and topography codes and by anatomical site using ICD-10 (International Classification of Diseases-10,10th revision). The NCRP produces periodic reports on ASIRs (standardized to world population), patterns of cancers, and time trends of incidence rates to explain the magnitude of cancer burden and highlight the priority areas of cancer research and control.

Cases registered with the respective PBCRs since its commencement [Mumbai, Chennai, Bangalore-1982, Delhi-1988, Kolkata-2005] till 31st December of the last reported year (2010 for all registries and 2009 for Delhi and Kolkata) were extracted from the NCRP database. The inclusion criteria included all records with ICD-O-3 morphology codes of 9510 to 9513. Variables of individual cases like name, age, sex, duration of stay at usual residence, date of first diagnosis, date of registration at the reporting institution, morphology, topography and sequence were extracted. Internal checks on the validity of the data were performed that included missing fields, age specified as >14 and consistency checks between topography and morphological coding. Records with discrepancies were corrected wherever possible by verifying the original records in the PBCRs. The latest completed data from 2006-2010 (Bangalore, Chennai, Mumbai) and 2005-2009 (Delhi, Kolkata) were extracted from the NCRP database.

Results

There were 214 registered retinoblastoma cases in age group 0-14 years from Bangalore, Chennai and Mumbai PBCRs (2006-2010) and Delhi and Kolkata PBCRs (2005-2009). Microscopic verification of histopathological diagnosis of retinoblastoma was available in 97.7% of pooled cases in PBCRs and 2.3% cases were diagnosed clinically or by other methods. In the HBCRs, microscopic verification of diagnosis (77%), and xray and ultrasonogram (23%) were the methods of diagnosis.

The maximum numbers were in the 0-4 age group in both sexes, accounting for 78% (females) and 81% (males) of pooled cases from PBCRs. Children aged 5-9 years accounted for 15 to 20% of cases, and thus, 98 to 100 % of the cases were under 10 years.

Population estimates in India are based on decadal national censuses conducted since 1961. Estimates of population by five yearly age groups have been projected based on difference distribution method for population projections (Takiar and Shobana, 2009). The 1991 and 2001 census were used for population estimates for the time period of 2006-2010. Relative proportions, sex ratio, method of diagnosis, and incidence rates (crude and age standardized) for each PBCR and pooled rates of the five PBCRs were calculated for the years 2005/06 to 2009/10 to provide recent information. ASIRs was calculated by the direct method, using the world standard population, expressed per million population. Standard error and 95% confidence limits was also determined. Rate Ratio at risk was calculated from cumulative risk and rate estimations.

Data from the Hospital based Cancer Registries (HBCRs) of Bangalore and Chennai (2006 to 2010), and Mumbai (2006-2008), were used to calculate relative proportions, sex ratio, and method of diagnosis.

Table 1A. Numbers, Relative Proportion by Age Group and Sex, Male Female Ratio of Retinoblastoma Cases in 0-14 Age Group from Selected PBCRs (2006-2010)

<table>
<thead>
<tr>
<th>Age Group</th>
<th>Bangalore</th>
<th>Chennai</th>
<th>Delhi*</th>
<th>Mumbai</th>
<th>Kolkata*</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>M</td>
<td>F</td>
<td>M</td>
<td>F</td>
<td>M</td>
</tr>
<tr>
<td>00-04</td>
<td>7 (70.0)</td>
<td>16 (76.2)</td>
<td>9 (81.8)</td>
<td>30 (66.7)</td>
<td>13 (100.0)</td>
</tr>
<tr>
<td>05-09</td>
<td>3 (30.0)</td>
<td>0 (0.0)</td>
<td>2 (18.2)</td>
<td>14 (31.1)</td>
<td>0 (0.0)</td>
</tr>
<tr>
<td>10-14</td>
<td>0 (0.0)</td>
<td>0 (0.0)</td>
<td>2 (2.6)</td>
<td>1 (2.2)</td>
<td>0 (0.0)</td>
</tr>
<tr>
<td>00-14</td>
<td>10 (100.0)</td>
<td>21 (100.0)</td>
<td>11 (100.0)</td>
<td>76 (100.0)</td>
<td>45 (100.0)</td>
</tr>
<tr>
<td>M/F</td>
<td>2</td>
<td>1.9</td>
<td>1.7</td>
<td>1.6</td>
<td>1.1</td>
</tr>
</tbody>
</table>

*Data 2005-2009

Table 1B. Numbers, Relative Proportion (%) by Age Group and Sex, Male Female Ratio of Retinoblastoma Cases in 0-14 Age Group in Selected HBCRs (2006-2010)

<table>
<thead>
<tr>
<th>Age group</th>
<th>Bangalore</th>
<th>Chennai</th>
<th>Mumbai*</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>M</td>
<td>F</td>
<td>M</td>
</tr>
<tr>
<td>00-04</td>
<td>13 (72.2)</td>
<td>9 (81.8)</td>
<td>13 (81.3)</td>
</tr>
<tr>
<td>05-09</td>
<td>5 (27.8)</td>
<td>2 (18.2)</td>
<td>3 (18.7)</td>
</tr>
<tr>
<td>10-14</td>
<td>0 (0.0)</td>
<td>0 (0.0)</td>
<td>0 (0.0)</td>
</tr>
<tr>
<td>00-14</td>
<td>18 (100.0)</td>
<td>11 (100.0)</td>
<td>16 (100.0)</td>
</tr>
<tr>
<td>M/F</td>
<td>0.5</td>
<td>0.7</td>
<td>1.7</td>
</tr>
</tbody>
</table>

*Data 2006-2008
cases occurred in 0-9 age group. The male to female (M/F) ratio varied from 1.1 (Kolkata) to 2.0 (Bangalore) (Table 1A). In the HBCRs, maximum numbers occurred in 0-4 age group (83%), and the M/F ratio was <1 in Bangalore and Chennai and 1.7 in Mumbai (Table 1B).

The pooled (all PBCRs) crude incidence rate in 0-14 age group was 3.5 and the pooled ASIR was 4.4 per million. The pooled ASIR in the 0-4, 5-9 and 10-14 age group was 9.6, 2.0 and 0.1 respectively. The pooled ASIR in 0-14 age group by sex was 5.2 (males) and 3.5 (females). (Table 2).

The ASIR in 0-14 age group males ranged from 2 per million in Mumbai to 10.4 per million population in Chennai. In 0-14 age group females, the ASIR ranged from 1.3 in Mumbai to 7.8 in Kolkata (Table 2). The ASIRs for males was higher than females in all PBCRs except Kolkata. The Rate Ratio of ASIR by sex was greater than 1 in all PBCRs except Kolkata (Table 2).

Discussion

Childhood cancers in age group 0-14 years ranged from 0.5 % to 5.8 % of all cancers in the PBCRs in India (National Cancer Registry Programme,2012). The four PBCRs and all the HBCRs have contributed to the NCRP since the 1980s except Kolkata PBCR (since 2005), and have established datasets for analysis with microscopically verified diagnosis of 97.7% in PBCRs and 77% in HBCRs. In both the PBCRs and HBCRs, the maximum numbers of cases occurred in the 0-4 age group, a pattern reported in earlier studies in India (Yeole et al., 2002; Swaminathan et al., 2008; Hazarika et al.,2014) and in developed nations (MacCarthy et al., 2009; Mitra et al., 2012). The M/F ratio in Chennai (1.9) and Bangalore PBCRs (2.0) was much higher than the other PBCRs. Globally the M/F ratio of retinoblastoma has been reported as unity,(Parkin et al.,1998; MacCarthy et al., 2009; Mitra et al., 2012) and in the past, Chennai had reported a M/F ratio of 1.0 (Swaminathan et al., 2008) and Mumbai of 1.4 (Yeole et al., 2002).

The ASIR in 0-14 age group males ranged from 2 per million in Mumbai to 10.4 per million population in Chennai. In 0-14 age group females, the ASIR ranged from 1.3 in Mumbai to 7.8 in Kolkata (Table 2). The ASIRs for males was higher than females in all PBCRs except Kolkata. The Rate Ratio of ASIR by sex was greater than 1 in all PBCRs (p<0.05) except Kolkata (Table 2).
Retinoblastoma was among the top five childhood cancers reported by the Chennai PBCR (1990-2001) with ASIR of 7.6 (males) and 7.9 (females)(Swaminathan et al., 2008). The latest data from the Population based Cancer Registries (2009-2011) in India indicate that ASIR of retinoblastoma could range from 6.7 per million population among 0-14 age group female children in Kolkata to 12.3 per million among 0-14 age group male children in Chennai.

(National Cancer Registry Programme, 2012). In this study, Chennai had high rates in 0-14 males (10.4) and lower rate in females (5.6). The rates of Mumbai PBCR are two times lower than reported in the late 1980s (Yeole et al., 2002). The rates reported in Bangalore PBCR is lower than the incidence rate of 3.1 (both sexes) in 1982-1989 (Nandakumar et al., 1996). Kolkata is second to Chennai in incidence in males. The ASIR of 0-14 male children in Chennai (10.4) was higher than the maximum reported ASIR of microscopically verified retinoblastoma in Malawi, Africa (8.0 per million in males) in the latest release of Cancer Incidence in five continents Vol X (2003-2007) (Forman et al., 2013).

The incidence rates in both sexes were highest in the 0-4 age group and decreased as age increased, being almost nil in 10-14 age group, a pattern observed globally (Table 3). The incidence rate in 0-4 age group, ranged from 5.1 (Bangalore) to 21.7 (Chennai) per million among males, and 3.4 (Mumbai) to 18.9 (Kolkata) among female children. The incidence rates in 0-4 male children were greater than the rates in female children in all PBCRs, except for Kolkata (Table 2). Previously, Mumbai PBCR had shown higher rates in 0-4 males (9.4) as compared to female children (5.9) (Yeole et al., 2002), while Chennai had revealed similar incidence rates in 0-4 male (15.9) and female (16.1) children (Swaminathan et al., 2008). Similar rates in both sexes in 0-4 age group has been reported from Great Britain and Korea (MacCarthy et al., 2009; Park et al., 2014). The Male Female Incidence Rate Ratio with 95% confidence limits (Table 2) revealed that the incidence rates were statistically significantly higher in males as compared to females when standardized to the world population in all PBCRs except Kolkata.

It was also observed that the trend in incidence rates in 0-4 years children (both sexes) showed a non-significant decrease over time in Bangalore, Chennai and Delhi, and statistically significant decrease in Mumbai (data not presented), which may be related to issues of low coverage. Trend analysis in developed nations show varied patterns- an increase in incidence in Great Britain (1983-2002) (MacCarthy et al., 2009), a decrease in incidence rates in Canada (1992-2006) among males (Mitra et al., 2012), and stable rates in the USA in both sexes (1975-2004) (Broaddus et al., 2009). In extremely rare cancers with low numbers, the trends observed may be due to random changes in the annual rates. Population expansion, changes in demographic composition and demarcation of geographic area, decrease in infant mortality rate, increased general awareness on cancer, improved diagnostic and treatment services, improved registration practices and case ascertainment, increase in reporting centres could be reasons for any apparent changes in incidence rates over time.

An increasing trend in incidence of unilateral cases in 0 to 4 years, especially among below 1 year children over 1963-2002 was reported in Great Britain. The authors postulated that this could be due to improved ascertainment of cases or a common exposure to an environmental factor (MacCarthy et al., 2009). Our present analysis revealed a distinct variation in incidence rates in the PBCRs of different geographic regions, though trends in incidence rates shown a non-significant decrease in three of the PBCRs. It is unknown if this variation in incidence between the PBCRs is related to environmental factors. Globally, risk factors like exposure to sunlight (Hooper, 1999; Jemal et al., 2000), viruses (Orjuela et al., 2000), low level radiation (Stiller, 1993) and parental dietary factors (Orjuela et al., 2005; Bunin et al., 2013) have been explored but the evidence has been inconclusive.

Retinoblastoma is a genetic disease with biallelic inactivation or loss of the retinoblastoma 1 gene-13q14.2. It is known to occur as heritable (mutations occurring in the germline and present in all somatic cells), and non-heritable or sporadic forms (with mutations arising locally within the developing retina). Sex differences have not been reported in the heritable or non-heritable forms. It is also known that all bilateral cases are heritable (26-40%) and unilateral cases could be familial/heritable (10-15%) and sporadic (50-70%) (MacCarthy et al., 2009; Broaddus et al., 2009). On pooled analysis, 52.3% cases were unilateral, 6.5% were reported as bilateral in the PBCRs, while in the HBCRS 71% were unilateral and 12.5% were bilateral cases. The Mumbai PBCR had reported 17% of bilateral cases in an earlier study (Yeole et al., 2002). Hospital based cancer registries in developing countries like Oman (Khandekar et al., 2004) and Saudi Arabia (Khandekar et al., 2014) (where consanguinity is well known), and Malaysia (Subramaniam et al., 2014) have reported 25-36% (bilateral) and 63-75% (unilateral) retinoblastoma. Hospital based studies in India (Sahu et al., 1998; Harini et al., 2001) have reported consanguinity in 9-17% of children with retinoblastoma, an association that needs to be further studied.

The present study findings have to be interpreted carefully as they represent low number of cases over the latest five years of available data of a relatively rare childhood cancer. The database did not have complete information on laterality, sequence of eye affected, heritability, follow-up or survival, but has definitely opened up questions on trends in incidence, variation in age and sex group, and genotype-phenotype relationship.

It has been shown that children with positive family history of retinoblastoma who underwent regular comprehensive screening of their eyes since birth had early diagnosis (as young as 8 months), and treatment modalities that could save the eye, leading to better ocular and patient survival (Abramson et al., 2003). This aspect could be true of unilateral retinoblastoma cases as well if detected early. In a hospital based study of infants in Iran, the median age at diagnosis of retinoblastoma was 6.96 months and the overall 5 year survival was 77% (Mehdiabadi GB et al., 2014), which was mainly attributed to early diagnosis and treatment interventions.
Thus early detection needs to be emphasized through raising awareness of the medical fraternity (primary care physicians, obstetricians, pediatricians) to ask about family history of retinoblastoma, to conduct screening examination of the eye of the newborn and serially during a child’s preventive health care visits, and facilitate prompt referral when there are risk factors like family history, leuokocoria and strabismus. Public health programmes for raising awareness and screening of eyes of children as a component of existent child health programmes have to be considered, especially as 15% of cases were 5 to 9 years, which could indicate late diagnosis.

Current treatment modalities of systemic chemotherapy and focal treatment have contributed significantly to disability limitation and patient survival. There has been a considerable body of knowledge on the genetics of the retinoblastoma gene and its protein, and there is potential for developing targeted molecular therapies(Sachdeva and O’Brien, 2012). Survival studies of retinoblastoma have shown less than 50% absolute survival in Chennai PBCR (Swaminathan et al., 2008), suggesting that there is need for concerted efforts in screening, treatment and follow up of RB children with retinoblastoma in India. Genetic testing could inform the type of mutation (germline or somatic), risk of unilateral or bilateral disease, choice of treatment protocols and prognosis of the condition. A diagnostic model has been proposed for developing countries (Joseph et al., 2006), though genetic and clinical screening is still inaccessible and unaffordable for the affected children and families.

The retinoblastoma gene is a tumour suppressor gene and its mutations are related to the initiation and progression of cancers (Benedict et al., 1990). There is evidence that survivors of hereditary retinoblastoma are at increased risk as compared to non-heritable retinoblastoma cases, to develop non-ocular second primary tumors like sarcomas, brain tumour, melanoma, lung and bladder cancers(Moll et al.,1996; Kleinerman et al., 2000; 2007; MacCarthy et al., 2013). The risk is further increased if patients had been treated with radiotherapy for their retinoblastoma (Yu et al., 2009).

Projection of burden of retinoblastoma in India based on data of the NCRP revealed that there will be 1000 male and female children affected with retinoblastoma every year. As retinoblastoma is a unique disease due to its early age of occurrence, genetic aetiology, and risk of second cancers, there is tremendous scope for epidemiological and molecular genetics research.

References


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