

International incidence of childhood cancer, 2001–10: a population-based registry study



Eva Steliarova-Foucher, Murielle Colombet, Lynn A G Ries, Florencia Moreno, Anastasia Dolya, Freddie Bray, Peter Hesselning, Hee Young Shin, Charles A Stiller, and the IICC-3 contributors*



Summary

Background Cancer is a major cause of death in children worldwide, and the recorded incidence tends to increase with time. Internationally comparable data on childhood cancer incidence in the past two decades are scarce. This study aimed to provide internationally comparable local data on the incidence of childhood cancer to promote research of causes and implementation of childhood cancer control.

Methods This population-based registry study, devised by the International Agency for Research on Cancer in collaboration with the International Association of Cancer Registries, collected data on all malignancies and non-malignant neoplasms of the CNS diagnosed before age 20 years in populations covered by high-quality cancer registries with complete data for 2001–10. Incidence rates per million person-years for the 0–14 years and 0–19 years age groups were age-adjusted using the world standard population to provide age-standardised incidence rates (WSRs), using the age-specific incidence rates (ASR) for individual age groups (0–4 years, 5–9 years, 10–14 years, and 15–19 years). All rates were reported for 19 geographical areas or ethnicities by sex, age group, and cancer type. The regional WSRs for children aged 0–14 years were compared with comparable data obtained in the 1980s.

Findings Of 532 invited cancer registries, 153 registries from 62 countries, departments, and territories met quality standards, and contributed data for the entire decade of 2001–10. 385 509 incident cases in children aged 0–19 years occurring in 2·64 billion person-years were included. The overall WSR was 140·6 per million person-years in children aged 0–14 years (based on 284 649 cases), and the most common cancers were leukaemia (WSR 46·4), followed by CNS tumours (WSR 28·2), and lymphomas (WSR 15·2). In children aged 15–19 years (based on 100 860 cases), the ASR was 185·3 per million person-years, the most common being lymphomas (ASR 41·8) and the group of epithelial tumours and melanoma (ASR 39·5). Incidence varied considerably between and within the described regions, and by cancer type, sex, age, and racial and ethnic group. Since the 1980s, the global WSR of registered cancers in children aged 0–14 years has increased from 124·0 (95% CI 123·3–124·7) to 140·6 (140·1–141·1) per million person-years.

Interpretation This unique global source of childhood cancer incidence will be used for aetiological research and to inform public health policy, potentially contributing towards attaining several targets of the Sustainable Development Goals. The observed geographical, racial and ethnic, age, sex, and temporal variations require constant monitoring and research.

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Introduction

Cancers rarely occur before age 20 years, and when they do, they raise a range of medical, psychological, ethical, and societal concerns. These distinct types of neoplasms require specific treatment paths. Furthermore, the extent of the cancer burden in this young population is unknown in many low-income and middle-income countries (LMICs), where data on cancer incidence are not collected. Even in the presence of population-based cancer registries, collection of information about childhood cancers is often neglected because they

represent a small proportion of all cancers, additional data sources might be required, and the resulting statistics must be subjected to meticulous quality control because they are more sensitive to imprecision or missing information.

Since the publication of the *International Incidence of Childhood Cancer, volume 1* (IICC-1) in 1988¹ and IICC-2 in 1998,² no internationally comparable data on incidence patterns of childhood cancer have been published. To address this problem, the International Agency for Research on Cancer (IARC), in collaboration with the

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*Contributors listed in the appendix

Section of Cancer Surveillance, International Agency for Research on Cancer, Lyon, France

(E Steliarova-Foucher PhD, M Colombet MSc, A Dolya MSc, F Bray PhD); Division of Cancer Control and Population Sciences, National Cancer Institute, Bethesda, MD, USA (L A G Ries MS); Paediatric Cancer Registry, National Cancer Institute, Buenos Aires, Argentina (F Moreno PhD); Department of Paediatrics and Child Health, Stellenbosch University, Tygerberg Children's Hospital, Tygerberg, South Africa (P Hesselning PhD); Seoul National University Children's Hospital, Institute of Cancer Research, Seoul, South Korea (H Y Shin PhD); and National Cancer Registration and Analysis Service, Public Health England, Oxford, UK (C A Stiller MSc)

Correspondence to:

Dr Eva Steliarova-Foucher, Section of Cancer Surveillance, International Agency for Research on Cancer, World Health Organization, Lyon 69372, CEDEX 08, France steliarova@iarc.fr

See Online for appendix

For more on the IARC see <http://www.iarc.fr/>

Research in context

Evidence before this study

The worldwide incidence of cancer in children aged 0–14 years was reported in 1988 and 1998 in two volumes of the *International Incidence of Childhood Cancer* (IICC), which described incidence patterns in the 1970s and 1980s, respectively. We searched MEDLINE and participating registries for studies on the incidence of childhood cancer worldwide, with the search terms “childhood cancer”, “registry”, “incidence”, and “population”, in February, 2016, without language or publication date restrictions. We found that no globally comparable data on cancer types affecting children have been published since IICC-2, and no comparison of incidence patterns in children aged 15–19 years has been attempted on a global scale.

Added value of this study

This study provides an overview of the incidence of cancer in 2001–10 for children aged 0–19 years, based on data collected in 153 population-based cancer registries in 62 countries, departments, and territories on five continents. In addition to the sex, age, and tumour-specific incidence rates for

19 world regions or populations for 2001–10, we report an increase in the incidence of neoplasms since the 1980s in children aged 0–14 years. This study updates information on cancer incidence in children published almost 20 years ago and adds the first global overview of cancer incidence in young people aged 15–19 years. It comprehensively summarises the most recent and globally comparable data and presents information per sex, age group, geographical region, and tumour type.

Implications of all the available evidence

The wealth of information provided by this study constitutes a solid baseline to assess needs and define priorities in the area of paediatric oncology, supporting Sustainable Development Goal 3 to ensure healthy lives and promote wellbeing for all at all ages. The identified variations in cancer incidence according to region, sex, age group, and tumour type will provide a springboard for specific aetiological research. The shortage of high-quality local information should stimulate formation of new and more accurate data sources.

For more on the IACR see
<http://www.iacr.com/fr/>

For more on the IICC-3 see
<http://iicc.iacr.fr/>

International Association of Cancer Registries (IACR), has coordinated a study to assess the incidence of childhood cancer worldwide, the complete results of which will be published in IICC-3. The target age range for IICC-3 is 0–19 years, compared with 0–14 years in IICC-1 and IICC-2. Inclusion of the 15–19 years age group was motivated by the shortage of internationally comparable data in this transition age between childhood and adulthood. An age range of 0–19 years was also chosen in previous US and European studies of childhood cancer incidence and survival.^{3,4}

Here we provide an overview of the findings of the IICC-3 study, based on a selection of quality-assured datasets from around the world encompassing information for the complete decade of 2001–10. The objective of this study was to counteract the difficulties in collecting childhood cancer data from relevant data sources and provide globally comparable estimates of cancer occurrence in children to aid further research and adapt policy measures.

Methods

Study design and data sources

In this epidemiological, population-based registry study, we invited all population-based cancer registries that we identified among the membership of the IACR and from published literature in MEDLINE or other published sources, such as annual reports, in any language. The search terms used for the literature search were “population”, “childhood cancer”, “incidence”, and “registry”. Of the invited registries, we selected only those that had submitted their data and met standard data quality criteria. We included only registries covering

the entire decade of 2001–10, which eliminated some high-quality registries that covered non-overlapping time periods. Analyses for the 0–14 years age group were based on a larger database (the paediatric dataset) than those for the 0–19 years age group (general dataset), because the paediatric dataset included data from the paediatric cancer registries not collecting data in children older than 15 years and these registries tended to have a wider coverage than the cancer registries used in the general dataset (appendix pp 2–4).

All malignancies and non-malignant tumours of the CNS diagnosed before age 20 years (before age 15 years in most of the paediatric cancer registries) in the covered populations in 2001–10 were eligible for inclusion. In the USA, non-malignant tumours were only registered from 2004 onwards, so they were excluded. The submission from each registry contained cancer and population data, information about registration coverage, practices, and data sources, and a short narrative describing registry operations. Cancer data included individual records of cases with codes for the following: sex; age; date of birth; date of incidence; and tumour sequence (ie, the numerical order of occurrence of the neoplasm), site, morphology, behaviour, laterality, and most valid basis of diagnosis. The 3rd edition of the *International Classification of Diseases for Oncology*⁵ was required for coding of the tumour site, morphology, behaviour, multiple primary tumours, and basis of diagnosis. Non-conforming coding systems were converted and tumours were classified centrally according to the 3rd edition of the *International Classification of Childhood Cancer* (ICCC-3).⁶ We requested each contributing cancer registry to provide total population

counts for each ethnic or racial group, calendar year, sex, and year of age in the registration area of their registry.

Procedures

All submitted datasets were processed and assessed individually in a rigorous peer-review process. The assessment criteria included a range of indicators, including the following: minimum number of cases; proportion of cases confirmed from cancer tissue examination (microscopic verification); proportion of cases registered from death certificate only; proportion of cases with morphology not otherwise specified (ie, cases in the unspecified subgroups of ICCC-3 or those coded by unspecified morphology codes such as 8000, 8800, 9800); proportion of cases with an unlikely combination of site and morphology, or an unlikely age and tumour type; proportion of rare entities (ie, neoplasms that occurred with a frequency of <0.05% in large datasets); proportion of dates of birth and incidence that were incomplete; proportion of ages that were imprecise; overall incidence rates; proportion of cases and incidence by sex, age group, and tumour type; the stability of rates over time; proportion of multiple primary diagnoses; and consistency of population data. All questionable records and raised issues were fed back to the contributing registries with a request for correction or a response. This iterative process resulted in a marked improvement of the overall quality of the data included in the analyses. The improvements included completion of missing information, additional years of data, inclusion of information from missed data sources, correction of coding errors, improvements in calculation of age, inclusion of non-malignant CNS tumours, replacement of population data, and explanation of the questioned patterns.

Statistical analysis

We calculated age-specific incidence rates (ASR) for four 5-year age groups (0–4 years, 5–9 years, 10–14 years, and 15–19 years) as the quotient of the number of cases and the number of person-years in the respective categories of sex, geographical area, and racial or ethnic group for the applicable time period, expressed per million person-years. We defined person-years as the sum of the population counts in a specific geographical area surveyed by a registry in each year from 2001 to 2010, categorised by sex and age and, if relevant, race or ethnicity. In the absence of all required details, we estimated the covered population counts by linear interpolation (six registries) or extrapolation (three registries) from the data provided by the registries, based on an assumption of regular population growth, before calculating rates. The purpose of this study was to make comparisons among regions and time periods, so all reported incidence rates for the 0–14 years and 0–19 years age groups were adjusted via direct standardisation. We calculated

age-standardised rates (WSRs) as the weighted average of the three age-specific rates (to calculate rates for 0–14 years) or four age-specific rates (for 0–19 years), using the weights of the world standard population⁷ (appendix p 5). We calculated incidence sex ratios by dividing the incidence in male individuals with that in female individuals.

The results are presented for 19 geographical regions or populations and a combined total; all the defined regions are either UN-defined regions⁸ or an aggregate of UN-defined regions, with the exception of North America, for which we present data separately for Canada and the USA, and have split data for the USA into five racial or ethnic groups. Within the region of south Asia, eligible data were only available from India. The described region definition was also driven by the data availability and the sizes of the resulting respective regional datasets. We calculated the proportion of the covered population of the world and each region by dividing the included person-years by the total person-years in 2001–10 by age group, as estimated by the UN in 2015.⁹

To compare the incidence rates derived from the paediatric dataset for 2001–10 with those published in IICC-2,² we assigned IICC-2 registry data to the same geographical regions as used in this IICC-3 study. We calculated the WSRs and their 95% CIs for children aged 0–14 years. Although the target period of IICC-2 was the 1980s, the time periods of the contributing registries differed both in length and starting years, because they either followed on from the IICC-1 period or they could not provide other years of data.

Statistical analyses were done using Stata/IC (version 12.1).

Role of the funding source

The funder of the study had no role in study design, data collection, data analysis, data interpretation, or writing of the report. MC, AD, and ES-F had full access to all the raw data used in the study. The corresponding author had final responsibility for the decision to submit for publication.

Results

Of the 532 invited cancer registries, 420 submitted their data and 309 met standard data quality criteria. Of those, 153 quality-assured registries from 62 countries, departments, and territories contributed data for the entire decade (appendix pp 3–4). Of the 153 included registries, 72 were initially unacceptable, but improved through the iterative quality assurance procedure. Approximately 11.4% of the world population aged 0–14 years (contributing 18 376 710 144 person-years) was covered by the registries included in our study, ranging from 1.7% in south Asia (India) to 99.4% in North America (table 1). The coverage was slightly less, at 8.9%, for young people aged 15–19 years (contributing

	Population covered (%)		Person-years (1000s)		Cases		Microscopically verified (%)		Morphology NOS (%)		Non-malignant (%)	
	Age 0-14 years	Age 15-19 years	Age 0-14 years	Age 15-19 years	Age 0-14 years	Age 15-19 years	Age 0-14 years	Age 15-19 years	Age 0-14 years	Age 15-19 years	Age 0-14 years	Age 15-19 years
North Africa	3.0%	3.4%	18 606	6980	2011	975	90.6%	94.9%	13.4%	10.4%	0.5%	0.4%
Sub-Saharan Africa	5.1%	0.8%	167 696	6780	9249	988	86.3%	72.4%	8.3%	27.1%	0.8%	0%
Central America and Caribbean	8.1%	8.5%	49 803	16 030	6223	2304	92.9%	90.1%	9.6%	12.4%	0.9%	1.2%
South America	13.2%	3.9%	140 549	13 709	18 251	2241	95.2%	94.2%	7.0%	10.9%	2.4%	0.6%
North America	99.4%	99.9%	661 232	235 385	100 475	48 354	92.7%	96.4%	2.5%	3.1%	0.1%	0.2%
East Asia	4.4%	4.0%	133 678	53 486	17 009	7857	91.3%	93.6%	10.0%	10.4%	1.4%	1.3%
South Asia*	1.7%	2.2%	65 713	25 901	5662	2339	95.5%	93.3%	12.8%	16.2%	0.0%	0.0%
Southeast Asia	4.6%	5.2%	76 729	28 944	8984	3267	87.5%	84.9%	20.9%	23.7%	0.4%	0.2%
West Asia	11.4%	11.2%	78 603	23 316	10 896	4463	96.0%	96.7%	4.6%	5.8%	2.1%	3.0%
Eastern Europe	27.9%	21.5%	129 536	48 077	17 720	8456	94.3%	93.7%	6.9%	8.8%	4.4%	1.6%
Northern Europe	86.8%	31.7%	151 598	20 081	22 437	4109	92.0%	94.9%	3.8%	5.8%	11.1%	9.1%
Southern Europe	37.7%	25.9%	85 322	21 679	14 270	5281	93.0%	95.9%	4.4%	5.6%	7.7%	5.3%
Western Europe	94.0%	23.9%	285 149	26 482	43 987	6027	94.9%	96.7%	1.6%	2.6%	10.8%	7.4%
Oceania	60.4%	66.2%	50 585	17 492	7475	4199	94.5%	97.6%	3.1%	2.5%	3.0%	0.6%
Total†	11.4%	8.9%	2094 800	544 340	284 649	100 860	93.0%	95.1%	4.9%	6.1%	3.7%	1.6%

Data for the 0-14 years age group is based on the paediatric dataset, and data for the 15-19 years age group is based on the general dataset. NOS=not otherwise specified. *Comprising data from India only. †Coverage of the world population within the indicated age group.

Table 1: Coverage of the world population aged 0-19 years, person-years, numbers of cancer cases, and quality indicators of data included in the analyses, overall and by region, 2001-10

	Paediatric dataset (age 0-14 years)			General dataset (age 0-19 years)		
	Cases	Person-years (millions)	Incidence per million	Cases	Person-years (millions)	Incidence per million
Sex (WSR)						
Boys	156 721	1072	151.4	169 531	1039	163.2
Girls	127 917	1023	129.4	141 695	987	143.6
Age group, all cases (ASR)						
0-4 years	127 096	676	187.9	93 559	475	197.1
5-9 years	74 175	690	107.6	54 370	487	111.6
10-14 years	83 378	729	114.4	62 438	519	120.3
15-19 years	100 860	544	185.3
Age group, all cases (WSR)						
0-14 years*	284 649	2095	140.6	210 367	1481	147.2
0-19 years†	311 227	2025	155.8
Age group, malignant cases only (WSR)						
0-14 years	274 096	2095	135.6	206 027	1481	144.3
0-19 years	305 233	2025	152.8

ASR=age-specific rate. WSR=age-standardised rate (world standard). *Includes cases with unknown sex (11 in the paediatric dataset and one in the general dataset). †Includes one case with unknown sex.

Table 2: Numbers of cancer cases, person-years, and overall cancer incidence, by sex, age group, and malignant status

6 105 345 342 person-years), with the lowest being 0.8% in sub-Saharan Africa. Overall, 385 509 registered cases occurring in 2.64 billion person-years contributed to the analyses (tables 1, 2). Registration of non-malignant CNS tumours differed by registry, and was higher in the paediatric dataset than in the general dataset (table 1, 2). The overall WSRs were 140.6 per million person-years in children aged 0-14 years and 155.8 per million

person-years in those aged 0-19 years (table 2). The ASR in young people aged 15-19 years was 185.3 per million person-years.

Incidence rates were slightly higher in boys than in girls (incidence sex ratio was 1.17 in the 0-14 years age group and 1.14 in the 0-19 years age-group) and varied with age, region, and diagnostic group (table 2; appendix p 8). Across all regions, incidence was higher in boys than in girls (incidence sex ratio ranged from 1.1 to 1.4 in the 0-19 years age group; appendix p 8), except for the 15-19 years age group of Native Americans in the USA (incidence sex ratio of 0.9) and in east Asia (1.0), where girls had a higher incidence (ratios rounded to one decimal place; appendix p 8). The highest incidence sex ratio was observed in the age group of 5-9 years in south Asia (India; 1.7; appendix p 8). Sex-specific incidence varied by diagnostic group, with renal tumours and epithelial tumours more common in girls, with variations by age group (appendix p 8). Germ cell and gonadal tumours were also more common in girls than in boys in the 0-14 years age group (appendix p 8).

ASRs were higher in children aged 0-4 years and 15-19 years than in those aged 5-9 years and 10-14 years (table 2). ASRs also differed between regions, especially for the age groups with high ASRs (appendix pp 6, 9). In the 0-14 years group, overall WSRs varied from less than 100 per million person-years in sub-Saharan Africa, for Native American children in the USA, and in south Asia (India), to more than 150 per million person-years in some subpopulations of North America and Europe, and in Oceania (figure 1A; appendix p 6). In young people aged 15-19 years, the lowest ASR was observed in

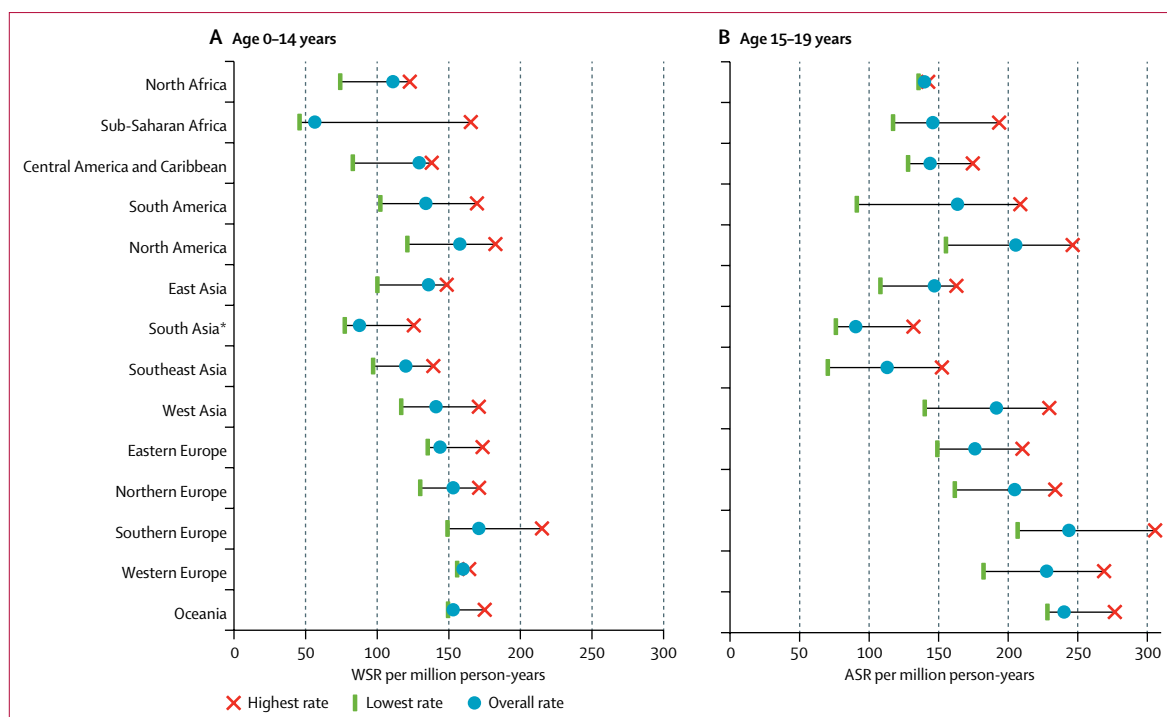


Figure 1: Variation in the overall incidence of childhood cancer by geographical region, 2001-10

Data are for children aged 0-14 years, from the paediatric dataset (A), and 15-19 years, from the general dataset (B). We only included registries with more than 100 cases when assessing the lowest and highest rates. ASR=age-specific rate. WSR=age-standardised rate (world standard population). *Comprising data from India only.

south Asia (India), whereas the highest ASRs were seen in some predominantly white populations of North America, Europe, and Oceania (figure 1B; appendix p 6).

The range of tumour types varied markedly with age group (figure 2). In children aged 0-4 years, leukaemia represented 36.1% (45 849 of 127 096) of all cases; however, the proportion of leukaemia cases was 15.4% (15 520 of 100 860) in young people aged 15-19 years. Conversely, lymphomas represented 5.3% (6766 of 127 096) of cases in children aged 0-4 years, and 22.5% (22 740 of 100 860) of cases in those aged 15-19 years. CNS tumours were the second most frequent tumour type after leukaemia in children aged 0-4 years (21 804 [17.2%] of 127 096), 5-9 years (19 471 [26.3%] of 74 175), and 10-14 years (16 680 [20.0%] of 83 378; figure 2). Epithelial tumours and melanoma represented 0.9% of all cases in children aged 0-4 years (1197 of 127 096), but were the second most common tumour group in young people aged 15-19 years (21 480 [21.3%] of 100 860).

The most common sympathetic nervous system tumour across all age groups was neuroblastoma. It was most frequent in children aged 0-4 years (15 888 [12.5%] of 127 096) and very rare in those aged 15-19 years (218 [0.2%] of 100 860; figure 2). Of note is the relatively high sympathetic nervous system tumour WSR of 10.2 per million person-years in black children in the USA, which contrasts with the WSR of 2.7 per million person-years in the mostly black population of Sub-Saharan Africa (table 3). Renal tumours were common in

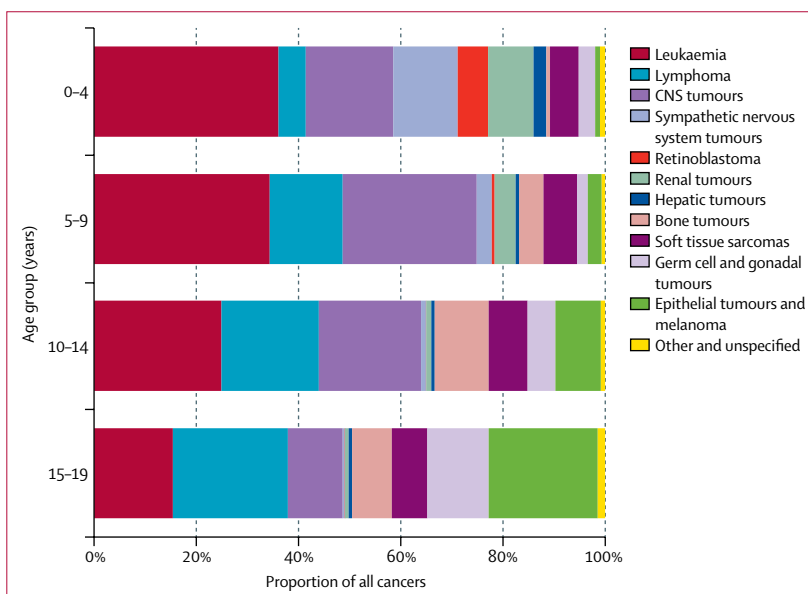


Figure 2: Proportional distribution of cancer type by age group, 2001-10, all regions combined

Tumours classified by *International Classification of Childhood Cancer, volume 3*.⁶ Statistics for children younger than 15 years are based on the paediatric dataset and the statistics for those aged 15-19 years are based on the general dataset.

children aged 0-4 years (11 297 [8.9%] of 127 096), and their relative frequency decreased in older age groups, to 0.7% (756 of 100 860) in young people aged 15-19 years (figure 2). Bone tumours constituted 4.7% (13 320

	Leukaemia		Lymphomas		CNS tumours		Sympathetic nervous system		Retino-blastoma		Renal tumours		Hepatic tumours		Bone tumours		Soft tissue sarcomas		Germ cell and gonadal tumours		Epithelial tumours and melanoma		Other and unspecified		
	n	WSR	n	WSR	n	WSR	n	WSR	n	WSR	n	WSR	n	WSR	n	WSR	n	WSR	n	WSR	n	WSR	n	WSR	n
North Africa	506	28.2	473	25.6	304	16.5	142	8.8	33	2.2	104	6.4	16	1.0	111	5.3	132	7.2	40	2.2	86	4.1	64	3.5	
Sub-Saharan Africa	2060	12.5	1664	9.8	1041	6.3	411	2.7	563	3.7	1031	6.7	165	1.0	385	2.1	1262	7.6	265	1.6	230	1.3	172	1.0	
Central America and Caribbean	2145	45.5	920	18.0	1023	21.1	277	6.6	155	3.8	292	6.7	88	2.0	294	5.2	337	6.9	239	4.7	283	5.1	170	3.5	
South America	6734	49.8	2319	16.2	3272	23.7	969	7.8	655	5.4	845	6.7	265	2.1	891	5.8	1062	7.7	634	4.6	425	2.8	180	1.3	
Canada	2780	55.5	949	16.2	1683	31.8	674	14.9	215	4.9	435	9.3	142	3.0	368	6.0	545	10.1	274	5.0	373	6.1	113	2.2	
USA																									
Native Americans	297	31.5	76	7.5	140	14.4	49	5.7	15	1.8	49	5.7	19	2.2	41	3.7	45	4.6	21	2.0	34	3.2	4	0.4	
Asian and Pacific Islanders	1294	43.9	378	12.1	596	19.9	214	7.7	107	3.9	117	4.2	99	3.5	133	4.1	177	5.8	200	6.4	125	3.8	11	0.4	
Black	2994	31.3	1442	13.4	2541	26.2	881	10.2	444	5.3	968	10.9	154	1.8	535	4.8	1093	10.8	383	3.8	423	3.7	65	0.7	
White Hispanic	7366	65.4	1885	16.2	3249	28.6	933	8.6	619	5.7	811	7.4	342	3.1	773	6.4	1169	10.2	755	6.5	578	4.8	79	0.7	
White non-Hispanic	17709	54.5	5905	15.8	13009	38.2	4452	15.0	1219	4.2	3029	10.0	823	2.7	2518	6.4	3757	10.9	1769	5.1	2781	7.1	230	0.7	
East Asia	5855	47.4	1730	12.7	2674	20.6	1236	12.0	506	5.1	542	5.1	377	3.5	958	6.2	925	7.2	1263	9.4	645	4.0	298	2.6	
South Asia*	2176	34.1	825	12.0	825	12.4	180	3.2	165	3.1	229	4.1	84	1.5	316	4.3	286	4.5	168	2.5	254	3.5	154	2.4	
Southeast Asia	3325	52.7	745	9.5	997	12.9	312	4.4	404	6.0	378	5.4	227	3.2	462	5.6	391	5.2	437	5.7	402	5.0	304	4.1	
West Asia	3459	45.2	1807	22.6	1918	24.5	727	10.1	326	4.6	534	7.3	126	1.7	532	6.4	631	8.2	342	4.4	413	5.0	81	1.0	
Eastern Europe	5265	44.3	2164	15.6	3793	30.0	1323	12.5	424	4.1	1062	9.8	248	2.2	895	6.0	1071	8.6	529	4.1	686	4.5	260	2.1	
Northern Europe	6987	49.1	2258	13.9	5821	38.9	1327	10.1	576	4.5	1235	9.3	277	2.1	942	5.5	1445	9.7	733	4.9	748	4.4	88	0.6	
Southern Europe	4200	51.1	1924	21.6	3156	37.6	1111	14.5	344	4.6	700	9.1	177	2.3	731	7.9	851	10.1	444	5.2	548	5.9	84	1.0	
Western Europe	13630	50.7	4992	16.3	10628	38.0	3162	13.0	1082	4.5	2447	9.8	490	1.9	2083	6.5	2695	9.7	1412	5.0	1291	4.1	75	0.3	
Oceania	2703	56.4	817	15.3	1285	26.2	500	11.4	187	4.3	384	8.5	115	2.6	352	6.4	449	9.0	292	5.9	354	6.2	37	0.8	
Total	92085	46.4	33273	15.2	57955	28.2	4234	10.4	18323	4.5	15192	8.2	4234	2.3	13320	5.7	18323	8.9	10200	4.9	10679	4.6	2469	1.2	

Regions are represented by the contributing cancer registries (appendix pp 3–4). Tumours classified by *International Classification of Childhood Cancer, volume 3*.²⁶ Data are based on the paediatric dataset. WSR=age-standardised rate (world standard) per million person-years. *Comprising data from India only.

Table 3: Numbers of cases and age-standardised incidence of tumours in children aged 0–14 years, 2001–10, by main diagnostic group

	Leukaemia		Lymphomas		CNS tumours		Sympathetic nervous system		Retino-blastoma		Renal tumours		Hepatic tumours		Bone tumours		Soft tissue sarcomas		Germ cell and gonadal tumours		Epithelial tumours and melanoma		Other and unspecified		
	n	ASR	n	ASR	n	ASR	n	ASR	n	ASR	n	ASR	n	ASR	n	ASR	n	ASR	n	ASR	n	ASR	n	ASR	n
North Africa	184	26.4	258	37.0	97	13.9	6	0.9	1	0.1	7	1.0	3	0.4	100	14.3	85	12.2	40	5.7	154	22.1	40	5.7	
Sub-Saharan Africa	129	19.0	183	27.0	41	6.0	2	0.3	0	0.0	16	2.4	21	3.1	101	14.9	208	30.7	58	8.6	126	18.6	103	15.2	
Central America and Caribbean	366	22.8	479	29.9	226	14.1	11	0.7	2	0.1	22	1.4	21	1.3	209	13.0	128	8.0	229	14.3	501	31.3	110	6.9	
South America	464	33.8	424	30.9	196	14.3	6	0.4	1	0.1	10	0.7	14	1.0	226	16.5	147	10.7	303	22.1	395	28.8	55	4.0	
Canada	570	27.2	1210	57.8	519	24.8	15	0.7	1	0.0	34	1.6	20	1.0	310	14.8	296	14.2	557	26.6	978	46.8	87	4.2	
USA																									
Native Americans	85	24.1	51	14.4	48	13.6	2	0.6	1	0.3	6	1.7	2	0.6	28	7.9	29	8.2	53	15.0	84	23.8	4	1.1	
Asian and Pacific Islanders	252	24.1	297	28.5	127	12.2	8	0.8	0	0.0	10	1.0	14	1.3	107	10.3	111	10.6	208	19.9	292	28.0	7	0.7	
Black	776	22.1	1385	39.4	542	15.4	29	0.8	0	0.0	86	2.4	44	1.3	417	11.9	592	16.8	421	12.0	758	21.6	43	1.2	
White	1508	43.0	1306	37.3	558	15.9	19	0.5	0	0.0	38	1.1	50	1.4	487	13.9	489	14.0	1277	36.4	1104	31.5	43	1.2	
Hispanic	3845	29.5	7178	55.1	3176	24.4	101	0.8	2	0.0	199	1.5	154	1.2	2014	15.5	2080	16.0	3636	27.9	7422	57.0	152	1.2	
non-Hispanic																									
East Asia	1683	31.5	1037	19.4	811	15.2	36	0.7	0	0.0	53	1.0	90	1.7	659	12.3	512	9.6	1014	19.0	1786	33.4	176	3.3	
South Asia*	565	21.8	345	13.3	263	10.2	7	0.3	1	0.0	18	0.7	12	0.5	298	11.5	171	6.6	178	6.9	365	14.1	116	4.5	
Southeast Asia	767	26.5	418	14.4	215	7.4	10	0.3	0	0.0	28	1.0	63	2.2	421	14.5	180	6.2	333	11.5	640	22.1	192	6.6	
West Asia	786	33.7	1222	52.4	513	22.0	23	1.0	0	0.0	26	1.1	25	1.1	448	19.2	297	12.7	338	14.5	736	31.6	49	2.1	
Eastern Europe	1048	21.8	2047	42.6	1001	20.8	41	0.9	0	0.0	80	1.7	51	1.1	637	13.2	477	9.9	1181	24.6	1693	35.2	200	4.2	
Northern Europe	512	25.5	954	47.5	726	36.2	17	0.8	0	0.0	26	1.3	18	0.9	303	15.1	244	12.2	494	24.6	788	39.2	27	1.3	
Southern Europe	650	30.0	1562	72.1	655	30.2	27	1.2	0	0.0	36	1.7	23	1.1	336	15.5	349	16.1	508	23.4	1081	49.9	54	2.5	
Western Europe	753	28.4	1466	55.4	814	30.7	19	0.7	0	0.0	38	1.4	30	1.1	449	17.0	357	13.5	777	29.3	1297	49.0	27	1.0	
Oceania	577	33.0	918	52.5	294	16.8	12	0.7	0	0.0	23	1.3	20	1.1	301	17.2	259	14.8	500	28.6	1280	73.2	15	0.9	
Total	15520	28.5	22740	41.8	10822	19.9	391	0.7	9	0.0	756	1.4	675	1.2	7851	14.4	7011	12.9	12105	22.2	21480	39.5	1500	2.8	

Regions are represented by the contributing cancer registries (appendix pp 3-4). Tumours classified by International Classification of Childhood Cancer, volume 3.³⁴ Data are based on the general dataset. ASR=age-specific rate per million person-years. *Comprising data from India only.

Table 4: Numbers of cases and age-specific incidence of tumours in young people aged 15-19 years, 2001-10, by main diagnostic group

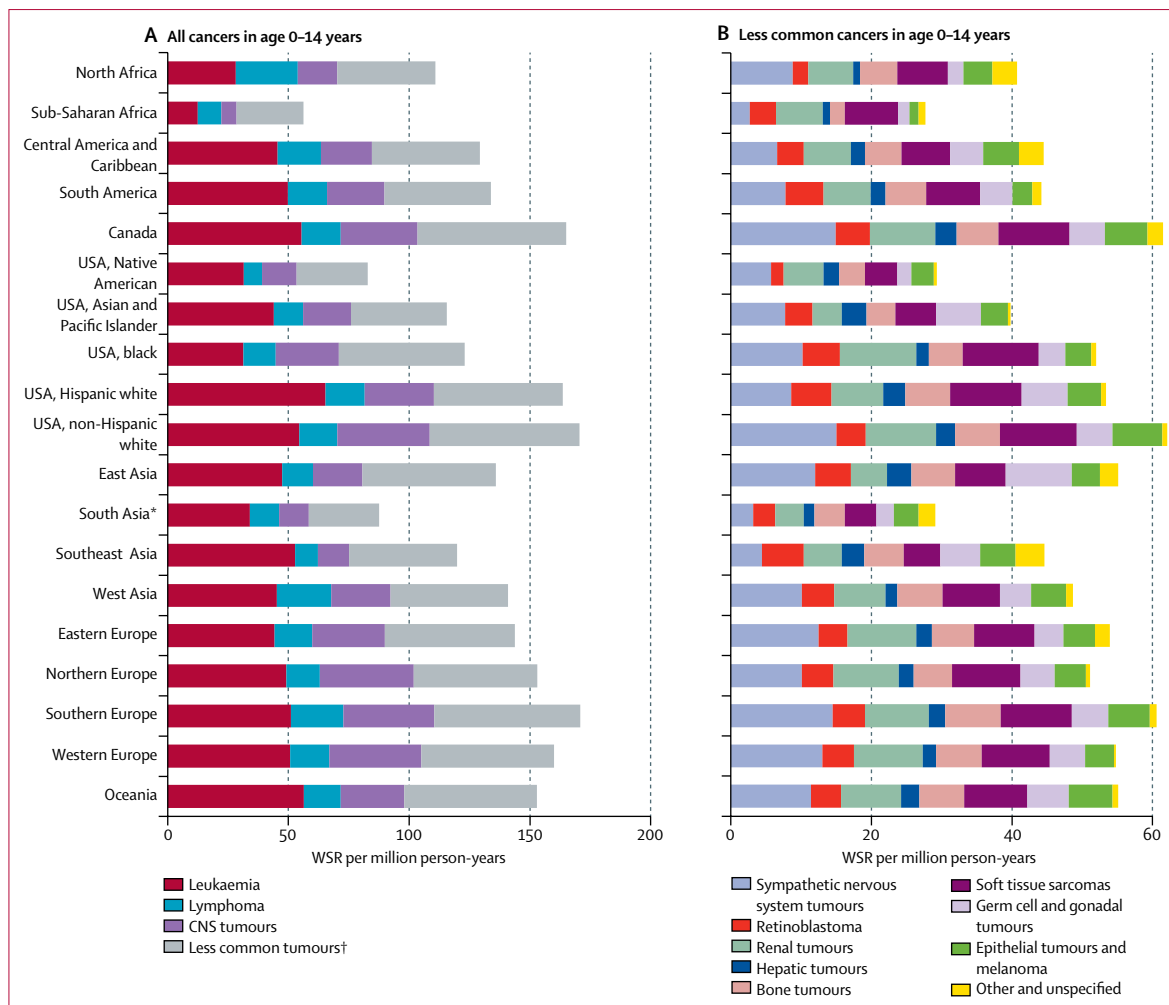


Figure 3: Incidence of cancer in children aged 0-14 years, 2001-10, by region
 Incidence of all tumours (A) and less common cancers (B). Tumours classified by *International Classification of Childhood Cancer, volume 3*.⁶ Data are based on the paediatric dataset. WSR=age-standardised rate (world standard population). *Comprising data from India only. †Defined in (B).

of 284649) of all cancers in children aged 0-14 years, and 7.8% (7851 of 100860) in those aged 15-19 years (figure 2; appendix p 10). Soft tissue sarcomas were present in a similar proportion of cases in children aged 0-14 years and those aged 15-19 years in most regions (table 3, 4; figure 2; appendix p 10). In sub-Saharan Africa, 46% (579 of 1262) of all soft tissue sarcomas in children aged 0-14 years were Kaposi's sarcoma compared with 57% (119 of 208) in those aged 0-19 years, whereas in all the other regions Kaposi's sarcoma represented less than 1% of all soft tissue sarcomas in children younger than 15 years and a maximum of 2% (14 of 592) in black children in the USA (data for all regions not shown). Germ cell and gonadal tumours were rare in children younger than 15 years (10200 [3.6%] of 284649; table 3), and more common in those aged 15-19 years (12105 [12.0%] of 100860; table 4). The group of epithelial neoplasms and melanoma comprises several

specific types of carcinomas (adrenocortical, thyroid, nasopharyngeal, skin), all other types of carcinoma (except those occurring in kidney, liver, and gonads), and melanoma. This group constituted 3.8% (10679 of 284649) of all tumours in children aged 0-14 years and 21.3% (21480 of 100860) in those aged 15-19 years (appendix p 10). The group of other and unspecified tumours comprised 0.9% (2469 of 284649) of all cases in children aged 0-14 years and 1.5% (1500 of 100860) of all cases in those aged 15-19 years. In 22 registries no tumours were classified in this category; other and unspecified tumours comprised 5% or more of all tumours in ten registries, including one registry for which 12% (68 of 561) of all tumours were classified as other and unspecified.

In children aged 0-14 years, the leading cancers in all regions combined were leukaemia, followed by CNS tumours, and then lymphomas, and this ranking was

observed in most world regions (table 3). The exception to this pattern was seen in the two African regions (appendix p 10). Incidence rates of all tumour types varied considerably between the regions (figure 3). The ratios of highest to lowest regional WSR were higher than 5.0 for CNS tumours (6.2), germ cell and gonadal tumours (5.9), neuroblastoma (5.6), epithelial tumours and melanoma (5.5), and leukaemia (5.2).

In young people aged 15–19 years, lymphomas were the most common cancers in all regions combined, followed by epithelial tumours and melanoma (table 4; figure 4). However, leukaemia was the most common neoplasm in this age group in South America, Native American and white Hispanic children in the USA, south Asia (India), and southeast Asia (table 4; figure 4). In Oceania, white non-Hispanic children in the USA, in east Asia, and in Central America and the Caribbean the most common cancer type was the group of epithelial cancers and melanoma (table 4; figure 4). Geographical variations in incidence, expressed as the ratio of the highest to the lowest ASR across regions, were 5.0 or higher for hepatic tumours (7.8), germ cell tumours (6.4), CNS tumours (6.0), lymphomas (5.4), epithelial tumours and melanoma (5.2), and soft tissue sarcomas (5.0).

From the 1980s to 2001–10, the overall WSR for all tumours in children aged 0–14 years increased from 124.0 per million person-years (95% CI 123.3–124.7) in the 1980s to 140.6 per million person-years (140.1–141.1) in 2001–10. The increase was seen in all regions except sub-Saharan Africa, and was smallest in Central America and the Caribbean (WSR 125.3 [95% CI 121.2–129.4] in 1980–89; 129.2 [95% CI 125.9–132.5] in 2001–10) and highest in southeast Asia (92.0 [95% CI 89.2–94.8]; 119.8 [95% CI 117.3–122.3]). In sub-Saharan Africa, a decrease was noted between the 1980s (81.0 [95% CI 76.8–85.2]) and 2001–10 (56.3 [95% CI 55.1–57.5]; figure 5; appendix p 7).

Discussion

Using data provided by 153 high-quality cancer registries, we report internationally comparable incidence rates of cancer in children aged 0–19 years during 2001–10. The incidence of cancer in children aged 0–14 years was 140.6 per million person-years, and in those aged 0–19 years was 155.8 per million person-years. We report considerable variations in incidence by diagnostic group,⁶ sex, age group, and world region. We have also shown that the overall WSR in children aged 0–14 years has increased between the preceding comparable publication covering the 1980s,¹⁰ and 2001–10. To our knowledge, this report presents the best available information on cancer incidence for the given period and age group.

We chose WSR as a relative estimate of cancer burden, but it should not be used for estimating numbers of cases in a population, because the value of WSR depends on the choice of the standard population. The world standard population⁷ has been used in previous international

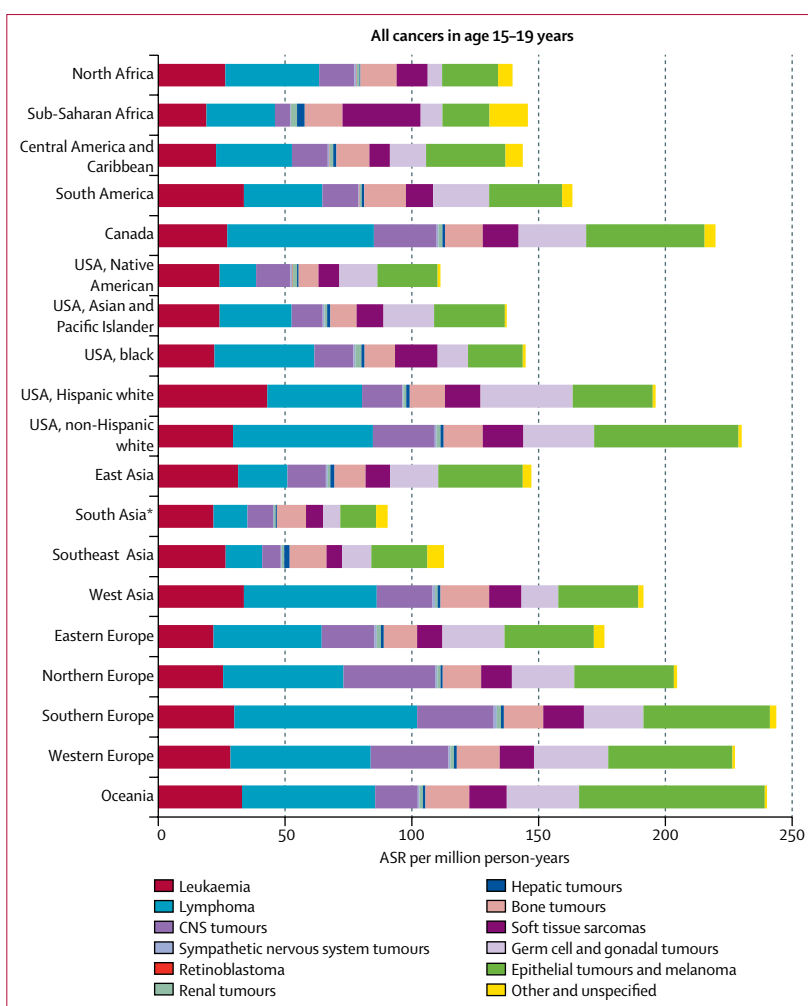


Figure 4: Incidence of cancer in young people aged 15–19 years, 2001–10, by region
Tumours classified by *International Classification of Childhood Cancer, volume 3*.⁶ Data are based on the general dataset. ASR=age-specific rate. *Comprising data from India only.

publications;^{1,2} choosing the same standard thus ensures comparability between populations and periods.

Although 20% of the cancers occurring in young people aged 15–19 years were epithelial tumours (the most prevalent histology types in adults) or melanoma, this age group also had high proportions of lymphomas, leukaemias, germ cell tumours, and sarcomas. Therefore, ICCC-3⁶ seems to be well adapted to reporting incidence in this age group, because the named tumour groups represent the main diagnostic categories in ICCC-3. We found that the incidence of cancer in young people aged 15–19 years was 185.3 per million person-years, based on 100 860 cases.

Because not all the contributing paediatric cancer registries could provide data for young people aged 15–19 years, the analyses for children aged 0–14 years were based on a different dataset (paediatric dataset) than were the analyses for children aged 0–19 years or those aged 15–19 years (general dataset). The two resulting datasets

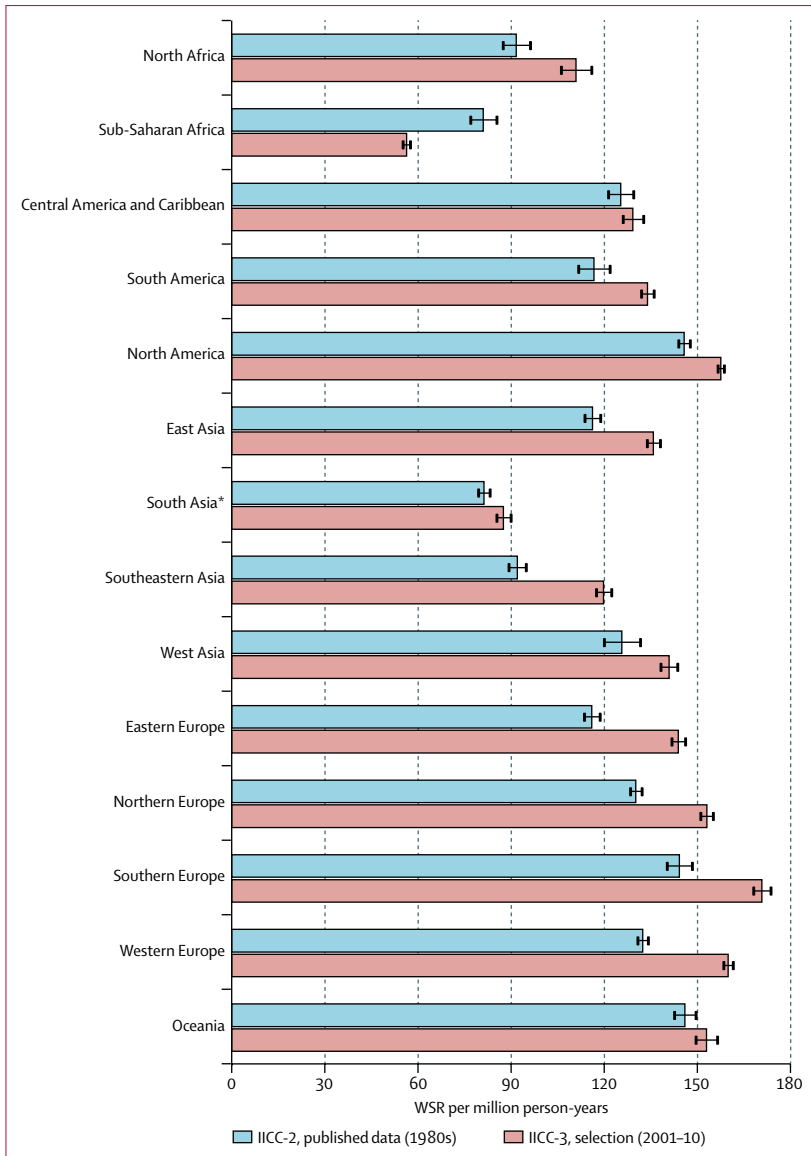


Figure 5: Comparison of incidence of neoplasms in children aged 0–14 years in 1980–89 and 2001–10, by region
 Data from the two time periods were standardised and peer reviewed using similar methods. Error bars show 95% CI. IICC-2=International Incidence of Childhood Cancer, volume 2.² IICC-3=paediatric dataset (defined in table 2 and appendix pp 3–4). WSR=age-standardised rate (world standard population). *Comprising data from India only.

gave different estimates of incidence for the (common) age group of 0–14 years, and this incidence was 5% lower in the paediatric than in the general dataset. This difference can be explained by variations in population coverage between the registries, and possibly distinctive data sources used by the two types of registries (ie, those that included data for young people aged 15–19 years and those that did not), as was shown previously in an analysis of data on childhood cancer in Europe.⁴ Although the results presented from both datasets are based on the best data available for each age group, they also emphasise the importance of quality assurance, particularly when dealing with a rare disease such as childhood cancer.

We selected the contributing registries because they provided quality-assured data for the entire decade of 2001–10. Inevitably, the reported rates were influenced by this selection; however, they provide the best and unique comparable global incidence estimates for the given period because they are not affected by intermittent contributions for parts of the target period.

A particular strength of our study comes from separate presentation of the incidence patterns for the five racial and ethnic groups distinguished in the US data, despite the difficulties in classifying the US population into racial and ethnic groups due to population mixing, migration, and self-declaration of ethnicity.¹¹ Unfortunately, the ethnic differences within multi-ethnic populations of Europe, Canada, and some other regions could not be readily studied. In other countries (South Africa, Israel, Australia) it was possible to gather incidence data for specific ethnicities within the population. For the purpose of simplicity these data were not included in this paper; however, they are available online on the IICC-3 website.

Our study improved the overall data quality in all participating cancer registries, particularly in 72 of 153 whose datasets were not acceptable at the beginning of the study. The study raised the registries' awareness of additional data sources (haematology, ophthalmology, orthopaedics, dermatologically, neurology, endocrinology, and paediatric clinics) and of specific quality control. We required a high level of quality and completeness for our study, because small errors or omissions would have had a large impact on the resulting incidence rates.

Approximately 30 cancer registries dropped out of the study, irrespective of the quality of their data. Such loss is especially regrettable in LMICs, where data are sparse and health policy has to be based on information from other countries. Data collected on more than 40 million person-years in 12 cancer registries (Canada [New Brunswick, Prince Edward Island], Denmark, Finland, Germany [Berlin, Brandenburg, Bremen, Mecklenburg-Vorpommern, Sachsen-Anhalt, and the Free States Saxony and Thuringia], and Singapore) could not be used because these registries were bound by requirements of a specific administrative procedure to approve provision of their data.

Although the cancer registries follow international guidelines, we observed wide variation in registration techniques, access to data sources, case eligibility criteria, application and interpretation of registration standards, and coding systems. Non-malignant tumours represented 0–10% of the total cases and affected international comparability. Non-malignant tumours were eligible for registration more commonly in Europe than in other parts of the world. Other practices reduced data quality (eg, restricted or no access to pathology reports), completeness (eg, restricted or no access to identifiable death certificates), or timeliness (eg, third-party data encryption). In some registries cases were coded as

microscopically verified only if the pathology report could be reviewed by registry staff. In others, precise morphology codes were not available for a large proportion of microscopically verified diagnoses. Registries are often under-resourced, and large investments for a small proportion of cases in general cancer registries might not be seen as a priority; however, these variations highlight the need for support to enable production of high-quality and comparable data.

A high proportion of specified diagnoses cannot be ensured without available specialised diagnostic facilities. The category of unspecified morphology might have included misclassified specific cancers, but in the absence of precise documentation it might have also included cases that were not malignant or cancers that did not occur in children.

Although the differences in sex ratios by diagnostic group and age might have to a large extent reflected true differences in disease occurrence,¹² the differences by geographical region could have more readily reflected sociocultural customs whereby boys are favoured over girls to seek medical attention when sick. The potential for such selective treatment in India and in some African populations was supported by our registry contacts in those regions when we inquired about a possibility of inequality in seeking health care.

The registries with no access to a national database of all causes of death might not have registered the cancers that were only discovered at the time of death, even though the proportion of missed cases might be low in childhood populations. In some countries death certification is non-existent whereas in others access to these records is hindered by data protection laws.

Factors external to the registry operations should also be considered. Higher socioeconomic level of development, better treatment facilities, and unobstructed access to data sources are linked to more advanced registration systems and better data quality. The registries operating in some LMICs were particularly affected by missing or infrequent official population estimates, and these data needed to be estimated by interpolation or extrapolation, influencing the reported rates. The low incidence rates observed in low-resource regions might in part have resulted from underdiagnosis,¹³ but our analysis could not quantify its extent. Treatment abandonment, common in LMICs,¹⁴ might also have led to under-registration if identified patients who refused treatment were not sufficiently well described (eg, missing dates or place of residence). Under-ascertainment of diagnosed cases might have resulted from administrative restrictions of access to medical files, political or social instability, competing needs, and inadequate political will, causing a dearth of resources, shortage or volatility of registry personnel, loss of perennial expertise, and missing or broken links with relevant data sources. Overestimates of incidence might have occurred in areas with superior treatment facilities if the place of

permanent residence could not be correctly determined for registered patients, and national coverage would neutralise such artefactual regional differences within a country.¹⁵ Accurate incidence rates are also difficult to obtain in ethnic minorities, such as the Native American population in the USA, possibly because of imprecise classification of the at-risk population, as well as patients with cancer.¹⁶ Cancer statistics in high-income countries might be influenced by overdiagnosis of some cancers detected by non-invasive imaging and screening tests, including neuroblastoma, thyroid cancer, melanoma, and kidney cancer.¹⁷

Although a complete assessment of time trends in incidence falls outside the scope of this report, our data showed that the overall incidence of registered neoplasms in childhood increased between the 1980s and the 2000s. Our comparison is certainly influenced by the composition of the two data pools (different registries have contributed to the two compared periods); however, the increase in the overall rates is clear. The reasons might include improved diagnosis (eg, imaging of brain tumours in high-income countries and less under-diagnosis in some parts of world). Cancer registration is also developing constantly, and more effective ascertainment techniques might have been used since the 1980s. Future detailed analyses of time trends by tumour group and subpopulations will bring more clarity to the interpretation of these secular changes.

The reduction of cancer incidence in sub-Saharan Africa probably had two main causes. First, this IICC-3 study included data from the national childhood cancer registry of South Africa, which contributed 70% of all cases for this region and reported very low incidence rates. Of the five datasets classified as sub-Saharan African in IICC-2, only one (from Namibia) reported a lower overall incidence rate than the South African registry in IICC-3. The possible reasons for the low rates in South Africa are discussed elsewhere.¹⁸ Second, the implementation of antiretroviral therapy in some areas affected by HIV¹⁹ has contributed to a decrease in incidence of Kaposi's sarcoma. This cancer type represented more than a third of the total cancer incidence in Kampala, Uganda, in the IICC-2 (WSR 182.7 per million person-years). The incidence of Kaposi's sarcoma in Kampala halved between the IICC-2 and this IICC-3 study, and a decreasing trend between the 1990s and the 2000s was also noted in a Kampala registry study.²⁰

Leukaemia, the most common cancer in children worldwide, had the largest impact on total cancer incidence. The strikingly low incidence of leukaemia in the African regions, notably in sub-Saharan Africa, might give some clues as to the cause of this cancer. Even though strong underdiagnosis of leukaemia is suspected in sub-Saharan Africa,²¹ leukaemia was also less commonly diagnosed in black children in the USA than in the other US ethnic groups; however, whether

leukaemia might be underdiagnosed as a result of reduced access to health care associated with lower economic status in this group is unclear.²² In children aged 0–14 years, the highest leukaemia rates were in Hispanic white children in the USA (40% of total incidence). The Native American component of Hispanic ancestry was a presumed risk factor, based on the observations that known risk alleles at loci identified in genome-wide association studies of European-ancestry populations in *CDKN2A*, *PIP4K2A*, *CEBPE*, and *ARID5B* were all significantly associated with Native American ancestry.²³ Comparatively high incidence rates and the largest proportion of leukaemia among all cancers were reported in southeast Asia. A link to the massive use of pesticides in this world region²⁴ to protect crops and increase yields should be examined in specific studies, since exposure to pesticides has been associated with leukaemia risk.^{25,26}

The male predominance among patients with lymphoma is probably a result of innate sex differences in susceptibility, but with increasing age other factors might play a role, such as the increased risk of HIV infection in boys compared with girls.²⁷ The highest incidence of lymphomas worldwide has been reported in the Mediterranean region (as shown in our study and others²⁸), and in HIV-infected populations,²⁹ with B-cell non-Hodgkin lymphoma an AIDS-defining malignancy. Burkitt's lymphoma is endemic and common in some sub-Saharan African childhood populations,³⁰ in which HIV infection can further accentuate its incidence.^{2,31} Hodgkin's lymphoma is rarer in populations of south and east Asia than in other parts of the world; the undergoing socioeconomic changes might gradually result in a similarly high incidence to that observed in Western populations.^{32,33}

The highest incidence of all CNS tumours was noted in high-income countries, which is clearly related to the wide availability of diagnostic facilities. The lower incidence rates in LMICs most probably (and proportionately) reflect poor access to neuroimaging facilities (eg, low number of CT or MRI scans, long waiting lists, and prohibitive costs of diagnostics tests).^{34,35} This poor access causes delay in diagnosis and possibly underdiagnosis of brain tumours.³⁶ Comparability of incidence of CNS tumours across the continents would be greatly improved if all registries attempted to collect information on non-malignant CNS tumours, at least in children.

This study describes the global cancer incidence patterns in children younger than 20 years for 2001–10, providing an update to comparable information that is now almost 20 years old.² Despite possible artefacts influencing data availability, quality, and comparability, the size of the studied populations and the observed differences in our study suggest that our data are sufficiently robust for international comparisons of childhood cancer occurrence, and provide useful pointers

for further studies. The collected data can be used for further research, including continued development of childhood-specific registration standards and guidelines, detailed studies in subpopulations and by tumour subtype, and global estimates of cancer indicators. This report constitutes a springboard for attaining several targets of the Sustainable Development Goal aiming at ensuring healthy lives and promoting wellbeing of all at all ages. High-quality data over time is vital to devising cancer control mechanisms in childhood populations worldwide. Local data that are internationally comparable are therefore indispensable for aetiological research and for effective health policy actions. Sharing of data internationally should not impose an unaffordable burden on data acquisition and validation in coordinated studies. To secure data availability and quality, constant support of cancer registration is required at local, national, and international levels.

Contributors

ES-F, LAGR, FM, PH, HYS, and CAS designed the study. ES-F, AD, and MC, in collaboration with LAGR, FM, PH, HYS, and CAS acquired the data for the study. MC did the analyses. ES-F and MC produced the tables and figures. AD and ES-F did the literature search. ES-F drafted the manuscript, which was reviewed, modified, and approved by ES-F, LAGR, FM, AD, FB, PH, HYS, and CAS.

Declaration of interests

We declare no competing interests.

Acknowledgments

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