



Clinical characteristics and survival patterns of subsequent sarcoma, breast cancer, and melanoma after childhood cancer in the DCOG-LATER cohort

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Abstract

Purpose Childhood cancer survivors are at increased risk of developing subsequent malignant neoplasms (SMNs). We compared survival and clinical characteristics of survivors with SMNs (sarcoma, breast cancer, or melanoma) and a population-based sample of similar first malignant neoplasm (FMN) patients.

Methods We assembled three case series of solid SMNs observed in a cohort of 5-year Dutch childhood cancer survivors diagnosed 1963–2001 and followed until 2014: sarcoma ($n=45$), female breast cancer ($n=41$), and melanoma ($n=17$). Each SMN patient was sex-, age-, and calendar year-matched to 10 FMN patients in the population-based Netherlands Cancer Registry. We compared clinical and histopathological characteristics by Fisher's exact tests and survival by multivariable Cox regression and competing risk regression analyses.

Results Among sarcoma-SMN patients, overall survival [hazard ratio (HR) 1.88, 95% confidence interval (CI) 1.23–2.87] and sarcoma-specific mortality (HR 1.91, 95% CI 1.16–3.13) were significantly worse compared to sarcoma-FMN patients (foremost for soft-tissue sarcoma), with 15-year survival rates of 30.8% and 61.6%, respectively. Overall survival did not significantly differ for breast-SMN versus breast-FMN patients (HR 1.14, 95% CI 0.54–2.37), nor for melanoma-SMN versus melanoma-FMN patients (HR 0.71, 95% CI 0.10–5.00). No significant differences in tumor characteristics were observed between breast-SMN and breast-FMN patients. Breast-SMN patients were treated more often with mastectomy without radiotherapy/chemotherapy compared to breast-FMN patients (17.1% vs. 5.6%).

Conclusions Survival of sarcoma-SMN patients is worse than sarcoma-FMN patients. Although survival and tumor characteristics appear similar for breast-SMN and breast-FMN patients, treatment differs; breast-SMN patients less often receive breast-conserving therapy. Larger studies are necessary to substantiate these exploratory findings.

Keywords Childhood cancer survivors · Subsequent malignant neoplasm · Survival · Epidemiology · Long-term complications

Flora E. van Leeuwen and Cecile M. Ronckers have contributed equally to this work.

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Introduction

Survivors of childhood cancer are at increased risk of developing long-term health problems even decades after treatment [1–4]. Subsequent malignant neoplasms (SMNs) are among the most serious adverse conditions that childhood cancer survivors may develop and represent a major cause of morbidity and mortality [5–9].

Several studies have addressed differences in prognostic, clinical, and/or histopathological characteristics of specific types of solid SMNs after childhood, adolescent, and young

adult cancers and comparable first malignant neoplasms (FMNs) in the general population [10–19]. Since the majority of these type of studies focused on breast-SMN, most were conducted in Hodgkin lymphoma or other adolescent and young adult (AYA) cancer survivors. Only few studies have specifically focused on young cancer survivors [12, 14, 17] and few have compared cause-specific survival between patients with specific types of SMNs following cancer at a young age and comparable FMN patients [14, 16].

Therefore, we examined clinical characteristics, overall survival, and cause-specific survival of patients in the Dutch Childhood Cancer Oncology Group—Long-Term Effects After Childhood Cancer (DCOG-LATER) cohort who developed SMNs of sarcoma, breast, or melanoma and compared those outcomes with matched patients having similar FMNs in the general population, based on the Netherlands Cancer Registry (NCR). We hypothesized that survival of SMNs may be different from survival of FMNs for several reasons: earlier detection due to surveillance efforts among survivors (favorable), variability of tumor characteristics due to different pathogenesis (favorable or unfavorable), limited treatment options due to previous childhood cancer treatment (unfavorable), and a higher frequency of other potentially lethal long-term chronic health conditions (unfavorable).

Methods

Subsequent solid malignant neoplasms in the DCOG-LATER cohort

The DCOG-LATER cohort includes 6,165 five-year childhood cancer survivors diagnosed before the age of 18 years in one of the seven Dutch pediatric oncology and stem cell transplant centers between 1 January 1963 and 31 December 2001. Detailed information on childhood cancer diagnosis and treatment was collected and incident SMNs were obtained via various methods, as previously described [8]. This particular study included cohort members with SMNs (referred to as SMN patients hereafter) diagnosed from 1989 through 2014 that were verified by the NCR (population-based ≥ 1989) [20]. We designed three case series of most prevalent invasive subsequent solid SMNs observed in the DCOG-LATER cohort: sarcoma, female breast cancer, and melanoma. Eligible morphology/topography codes are presented in Supplementary Table 1. In an earlier report on the current cohort, sarcoma, female breast cancer, and melanoma were shown to represent 20%, 20%, and 4%, respectively, of the total number of solid SMNs [8]. To avoid misclassification and bias, we excluded patients for whom the respective SMN of inclusion was a third tumor preceded by an intervening second malignancy treated with radiotherapy and/or chemotherapy.

Population-based first malignant neoplasms (FMN)

Each of the SMN patients in the three case series (sarcoma, female breast cancer, and melanoma) from the DCOG-LATER cohort was matched to 10 comparison patients with FMNs from the population-based NCR (referred to as FMN patients hereafter), based on sex, age at cancer diagnosis, calendar year of cancer diagnosis ± 3 years, and for sarcomas also on primary site (bone/soft tissue). When > 10 comparison patients fulfilled these criteria, the 10 patients closest to incidence date of the corresponding case were selected. When less than the 10 comparison FMN patients from the cancer registry fulfilled these criteria for a specific index SMN patient, the matching criterion for age was widened until 10 eligible comparison FMN patients were identified. That is, the FMN comparison group was supplemented with cancer patients closest in age at their respective FMN cancer diagnosis to the index SMN case diagnosis, but within the same diagnosis calendar year defined by the index SMN case. Sampled comparison patients were excluded from serving as FMN patients if they had a history of cancer, with the exception of basal cell carcinoma.

Data on prognostic, clinical, and histopathological characteristics

The following SMN/FMN characteristics were obtained from the NCR: morphology, stage, grade, treatment (for all three cancer sites yes/no indicators for surgery, chemotherapy, and radiotherapy and for breast cancer additionally yes/no indicators for mastectomy, breast-conserving surgery, and hormonal therapy), and vital status. For patients with breast cancer, information on receptor status was also retrieved. No genetic data were available. Patients were followed until date of death, date of emigration, or through 12/31/2016, whichever occurred first. Coded cause of death information for deceased SMN/FMN patients was obtained by linkage with Statistics Netherlands. We categorized cause of death into cancer of interest-related deaths and death due to other causes. Data collection for the study was exempted from review by the Institutional Review Boards of all participating centers and all data were processed and analyzed anonymously.

Statistical analyses

We compared clinical and histopathological characteristics for SMN and FMN patients by Fisher's exact tests. Overall survival was calculated using Kaplan–Meier methods and represented by Kaplan–Meier curves. For cause-specific mortality, cumulative incidences functions (probability of

death due to the specific cause) were estimated. When evaluating cancer of interest-related deaths, deaths due to other causes were treated as a competing risk and vice versa. We used Cox proportional hazard regression analyses to test differences in overall survival between the SMN and FMN groups in univariable models with patient group (SMN vs. FMN) as an independent variable and in multivariable models with additional factors hypothesized to influence a difference in survival between the SMN and FMN groups. For sarcoma, morphology was one of the factors we adjusted for in the analyses. For any sarcoma, we grouped morphology into three groups based on similarity in survival estimates reported in literature: (1) osteosarcoma/rhabdomyosarcoma; (2) chondrosarcoma; (3) other or unspecified. Proportionality of hazards for each variable was evaluated by adding interaction terms with follow-up time. Fine and Gray competing risk models were used to test differences between SMN and FMN patients with respect to mortality due to the cancer of interest and mortality from other causes [21]. All analyses were performed with Stata 14 software (StataCorp. 2015. *Stata Statistical Software: Release 14*. College Station, TX: StataCorp LP).

Results

Sarcoma—comparison of clinical and histopathological characteristics

Of 46 eligible patients with subsequent sarcoma (sarcoma-SMN) after childhood cancer, one patient was excluded because of a SMN diagnosis (acute lymphoblastic leukemia (ALL); treatment with radiotherapy and chemotherapy) prior to developing a sarcoma as third malignancy. We also excluded one comparison patient in the first primary sarcoma (sarcoma-FMN) group, who was erroneously selected for inclusion, despite a history of a prior cancer. In total, we included 45 sarcoma-SMN patients and 449 sarcoma-FMN patients. Among sarcoma-SMN patients, 30.4% also had a sarcoma as childhood cancer diagnosis, 8.7% had a retinoblastoma as childhood cancer diagnosis, and 60.9% had previous radiotherapy treatment to the sarcoma-SMN region (Supplementary Table 2).

Several differences were observed in the morphology of sarcoma-SMNs and sarcoma-FMNs. Twenty-seven percent of sarcoma-SMN patients had osteosarcoma, versus 12.9% of sarcoma-FMN patients (Table 1) ($p = 0.022$, not shown). In contrast, none of the sarcoma-SMN patients had Ewing sarcoma, while this was the morphology of 13.1% of sarcoma-FMN patients. Fibrosarcoma was much less common among sarcoma-SMN patients (2.2%) than among sarcoma-FMN patients (16.5%) ($p = 0.008$, not shown), with dermatofibrosarcomas protuberans being the

largest contributor of fibrosarcomas among FMN patients (62 cases, 13.8%). Remarkably, no dermatofibrosarcomas protuberans occurred among sarcoma-SMN patients. Among the minority with a known tumor grade (34.4%), sarcoma-SMN patients were more likely to have a high tumor grade than sarcoma-FMN patients (SMN 60.0% vs. FMN 26.0% poor/high grade and SMN 10.0% vs. FMN 2.0% undifferentiated/anaplastic grade for sarcoma-SMN and sarcoma-FMN patients, respectively; $p = 0.002$). We did not observe significant differences in stage distribution or treatments received.

Sarcoma—overall and cause-specific survival

Median follow-up time was 3.6 years (max 34.0 years) for sarcoma-SMN patients and 7.5 years (max 28.0 years) for sarcoma-FMN patients. At the end of follow-up, 62.2% of sarcoma-SMN patients were deceased compared to 33.2% of sarcoma-FMN patients (Table 1). Because dermatofibrosarcoma protuberans behaves in many respects as a benign tumor and has very high survival rates [22], we excluded those patients ($n = 62$, all sarcoma-FMN patients) from all survival analyses. Furthermore, we excluded Ewing sarcoma ($n = 59$, all sarcoma-FMN patients) and Kaposi sarcoma ($n = 19$, 18 sarcoma-FMN patients and 1 sarcoma-SMN patient) from all survival analyses, because those sarcoma types are rare to occur as SMN. Survival rates after 5, 10, and 15 years were 44.9%, 38.5%, and 30.8%, respectively, among sarcoma-SMN patients, considerably lower than those among sarcoma-FMN patients with 67.4%, 63.1%, and 61.6%, respectively (Table 2, Fig. 1a). Overall survival was significantly worse among sarcoma-SMN patients (multivariable HR 1.88; 95% CI 1.28–2.87) (Table 3, Fig. 1). Sarcoma-specific mortality was also significantly higher in sarcoma-SMN patients compared to sarcoma-FMN patients (15-year mortality 66.0% vs. 34.5% and multivariable HR 1.91; 95% CI 1.16–3.13, Tables 2 and 3). Mortality due to other causes after 5, 10, and 15 years was very low in all groups (<3%).

Stratified by type of sarcoma, survival differences between SMN and FMN patients seemed to be greater for soft-tissue sarcoma (multivariable HR 2.31; 95% CI 1.37–3.89) than for bone sarcoma (multivariable HR 1.21; 95% CI 0.57–2.53) (Supplementary Table 3, Supplementary Fig. 1). In analyses per specific sarcoma morphology type, we found no difference in survival between osteosarcoma-SMN and FMN patients (multivariable HR 1.02; 95% CI 0.45–2.34), but we found a statistically significant difference between synovial sarcoma SMN and FMN patients (multivariable HR 7.82; 95% CI 1.41–43.35) (not shown). For other specific sarcoma morphology types, numbers were too small to make meaningful comparisons.

Table 1 Characteristics of patients with sarcomas and melanomas as a subsequent malignant neoplasm (SMN) and matched patients with sarcomas and melanomas as a first malignant neoplasm (FMN)

	Sarcoma				<i>p</i> value	Melanoma				<i>p</i> value
	DCOG-LATER cohort SMN (<i>n</i> = 45)		Cancer registry FMN (<i>n</i> = 449) ^a			DCOG-LATER cohort SMN (<i>n</i> = 17)		Cancer registry FMN (<i>n</i> = 170)		
	<i>N</i>	%	<i>N</i>	%		<i>N</i>	%	<i>N</i>	%	
Age at diagnosis ^b					0.924					1.000
< 30 years	35	77.8	335	74.6		6	35.3	60	35.3	
30–39 years	8	17.8	83	18.5		6	35.3	60	35.3	
40+ years	2	4.4	31	6.9		5	29.4	50	29.4	
Sex ^b					1.000					1.000
Male	25	55.6	249	55.5		6	35.3	60	35.3	
Female	20	44.4	200	44.5		11	64.7	110	64.7	
Incidence year ^b					0.909					1.000
1989–1999	12	26.7	129	28.7		3	17.7	29	17.1	
2000–2009	25	55.6	249	55.5		8	47.1	81	47.7	
2010–2014	8	17.8	71	15.8		6	35.3	60	35.3	
Morphology sarcoma					NE					
Osteosarcoma	12	26.7	58	12.9						
Chondrosarcoma	4	8.9	57	12.7						
Ewing sarcoma	0	0.0	59	13.1						
Rhabdomyosarcoma	3	6.7	23	5.1						
Leiomyosarcoma	4	8.9	22	4.9						
Fibrosarcoma	1	2.2	74	16.5						
Malignant fibrous histiocytoma	2	4.4	12	2.7						
Malignant peripheral nerve sheath tumor	3	6.7	23	5.1						
Liposarcoma	3	6.7	19	4.2						
Synovial sarcoma	3	6.7	24	5.4						
Kaposi sarcoma	1	2.2	18	4.0						
Other or unspecified bone	1	2.2	19	4.2						
Other or unspecified soft tissue ^d	8	17.8	41	9.1						
Morphology melanoma										0.335
Nodular						1	5.9	20	11.8	
Superficial spreading						15	88.2	117	68.8	
Other or unspecified						1	5.9	33	19.4	
Stage ^e					0.977					1.000
I	12	57.1	106	50.2		13	76.5	120	73.6	
II	5	23.8	53	25.1		3	17.7	28	17.2	
III	0	0.0	7	3.3		1	5.9	15	9.2	
IV	4	19.1	45	21.3		0	0.0	0	0.0	
Unknown	24		238			0		7		
Grade ^e					0.002					NE
Well/low grade	4	20.0	79	52.7		0	0.0	0	0.0	
Moderate/intermediate	2	10.0	29	19.3		0	0.0	0	0.0	
Poor/high	12	60.0	39	26.0		0	0.0	0	0.0	
Undifferentiated/anaplastic	2	10.0	3	2.0		0	0.0	0	0.0	
Unknown	25		299			17		170		
Surgery					0.256					0.382
No	13	28.9	95	21.2		1	5.9	4	2.4	
Yes	32	71.1	354	78.8		16	94.1	166	97.7	
Radiotherapy					0.716					1.000

Table 1 (continued)

	Sarcoma				<i>p</i> value	Melanoma				<i>p</i> value
	DCOG-LATER cohort SMN (<i>n</i> = 45)		Cancer registry FMN (<i>n</i> = 449) ^a			DCOG-LATER cohort SMN (<i>n</i> = 17)		Cancer registry FMN (<i>n</i> = 170)		
	<i>N</i>	%	<i>N</i>	%		<i>N</i>	%	<i>N</i>	%	
No	33	73.3	341	76.0		17	100	168	98.8	
Yes	12	26.7	108	24.1		0	0.0	2	1.2	
Chemotherapy					0.632					NA
No	26	57.8	277	61.7		17	100	170	100	
Yes	19	42.2	172	38.3		0	0.0	0	0.0	
Treatment category					0.165					0.493
No treatment	6	13.3	23	5.1		1	5.9	4	2.4	
Surgery only	16	35.6	199	44.3		16	94.1	164	96.5	
Surgery + RT and/or CT	16	35.6	155	34.5		0	0.0	2	1.2	
RT and/or CT only	7	15.6	72	16.0		0	0.0	0	0.0	
Vital status at end of follow-up					<0.001					0.685
Alive	17	37.8	300	66.8		15	88.2	153	90.0	
Deceased	28	62.2	149	33.2		2	11.8	17	10.0	

CT chemotherapy, FMN first malignant neoplasm, NE not estimable, RT radiotherapy, SMN subsequent malignant neoplasm

^a1 control was excluded, because this patient appeared to have had a prior malignancy

^bMatching factors

^cIncluded 1 unspecified bone sarcoma among sarcoma-SMN patients

^dIncluded 2 each with unspecified soft-tissue sarcoma and giant cell sarcoma, 1 each with spindle cell sarcoma, round cell sarcoma, hemangiomasarcoma, and hemangiopericytoma among sarcoma-SMN patients

^ePercentages and *p* values for stage and grade are based on subjects with known information only

Breast cancer—comparison of clinical and histopathological characteristics

Of 42 eligible patients with subsequent female breast cancer (breast-SMN) after childhood cancer, one was excluded because of radiotherapy treatment for an intervening meningioma prior to breast cancer diagnosis. We included 41 breast-SMN patients and 410 matched first primary female breast cancer (breast-FMN) patients. Of all breast-SMN cases, 17.1% occurred after a Hodgkin lymphoma diagnosis and 40.0% after a sarcoma diagnosis (Supplementary Table 2). Almost one-third of the breast-SMN patients (31.7%) had no prior radiotherapy and also nearly one-third (31.7%) had prior radiotherapy to the region where the breast cancer developed (26.8% with chest radiotherapy and 4.9% with total body irradiation). More than half of the breast-SMN patients (53.7%) had prior anthracycline treatment (Supplementary Table 2).

Table 4 shows tumor and breast cancer treatment characteristics of the breast-SMN and breast-FMN patients. The proportion of patients with infiltrating ductal carcinoma was the same in the breast-SMN (87.8%) and FMN (87.3%) groups, as were the distributions of stage and grade. Patients in the breast-SMN group were more likely

to have been treated without surgery (SMN 9.8% vs. FMN 3.2%), but markedly more likely to have received mastectomy (SMN 63.4% vs. FMN 49.3%) than breast-FMN patients ($p = 0.024$). Breast-SMN patients were less likely, however, to have received radiotherapy (SMN 51.2% vs. FMN 71.7%, $p = 0.011$) and chemotherapy (SMN 58.5% vs. FMN 76.3%, $p = 0.011$) compared to breast-FMN patients. No significant differences were observed in hormone receptor status single or combined, i.e., triple negative, and receipt of hormonal therapy between breast-SMN and breast-FMN patients. When evaluating combinations of treatments, a higher proportion of breast-SMN patients received mastectomy without RT or CT compared to breast-FMN patients (SMN 17.1% vs. FMN 5.6%). Also, a lower proportion of breast-SMN patients was treated with breast-conserving surgery with radiotherapy and/or chemotherapy (SMN 26.8% vs. FMN 46.6%). In both groups, the majority of patients who did not have surgery for breast cancer had stage IV disease (SMN 75.0% vs. FMN 84.6%). For at least three of seven (42.9%) breast-SMN patients with mastectomy without radiotherapy or chemotherapy (\pm hormonal therapy), there was an indication that prior childhood cancer treatment limited breast cancer treatment options. One breast-SMN patient received a mastectomy

Table 2 Overall survival and cause-specific mortality of subsequent malignant neoplasm and matched first malignant neoplasm cohorts

	Sarcoma ^a		Breast cancer ^b		Melanoma	
	SMN (<i>n</i> = 45)	FMN (<i>n</i> = 449) ^c	SMN (<i>n</i> = 41)	FMN (<i>n</i> = 410)	SMN (<i>n</i> = 17)	FMN (<i>n</i> = 170)
	% (95% CI)	% (95% CI)	% (95% CI)	% (95% CI)	% (95% CI)	% (95% CI)
5-year overall survival	44.9 (29.9–58.9)	67.4 (61.8–72.3)	93.8 (77.4–98.4)	88.5 (84.7–91.4)	94.1 (65.0–99.2)	93.4 (88.4–96.3)
10-year overall survival	38.5 (23.7–53.2)	63.1 (57.3–68.3)	73.6 (52.0–86.6)	79.9 (74.8–84.1)	82.4 (42.6–95.7)	88.9 (82.3–93.2)
15-year overall survival	30.8 (16.5–46.3)	61.6 (55.6–67.1)	63.1 (34.9–81.8)	77.4 (71.5–82.2)	82.4 (42.6–95.7)	87.2 (79.5–92.2)
5-year mortality ^d						
Cancer of interest ^e	51.6 (35.7–65.3)	30.8 (25.6–36.1)	6.2 (1.1–17.9)	10.8 (7.8–14.3)	0.0	5.5 (2.7–9.7)
Other causes ^f	2.4 (0.2–11.0)	0.7 (0.1–2.2)	0.0	0.0	0.0	0.6 (0.1–3.0)
10-year mortality ^d						
Cancer of interest ^e	58.2 (41.2–71.8)	33.9 (28.5–39.4)	22.7 (9.0–40.0)	17.2 (13.2–21.7)	12.5 (0.7–42.3)	10.0 (5.6–15.8)
Other causes ^f	2.4 (0.2–11.0)	1.9 (0.7–4.2)	3.8 (0.3–16.1)	0.7 (0.1–3.6)	0.0	0.6 (0.1–3.0)
15-year mortality ^d						
Cancer of interest ^e	66.0 (47.6–79.3)	34.5 (29.0–40.0)	33.2 (12.2–56.1)	19.8 (15.0–25.1)	12.5 (0.7–42.3)	10.0 (5.6–15.8)
Other causes ^f	2.4 (0.2–11.0)	1.9 (0.7–4.2)	3.8 (0.3–16.1)	0.7 (0.1–3.6)	0.0	2.3 (0.4–8.1)

CI confidence interval, FMN first malignant neoplasm, SMN subsequent malignant neoplasm

^aPatients with dermatofibrosarcoma protuberans (*n* = 62 FMN), Ewing sarcoma (*n* = 54 FMN), and Kaposi sarcoma (*n* = 18 FMN and *n* = 1 SMN) were excluded

^bMetastatic breast cancers (stage IV; *n* = 3 breast-SMN patients and *n* = 22 breast-FMN patients) and breast cancers with unknown stage (*n* = 1 breast-SMN patients and *n* = 2 breast-FMN patients) were excluded

^c1 control was excluded, because this patient appeared to have had a prior malignancy

^dBecause of missing causes of death, the values do not always correspond with the overall survival numbers

^eMortality due to other causes was considered a competing risk

^fMortality due to the cancer of interest was considered a competing risk

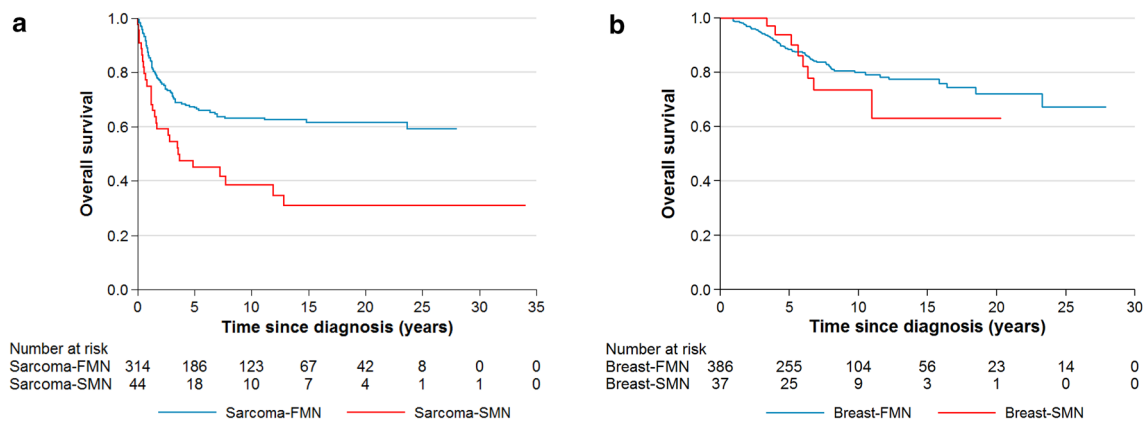


Fig. 1 Overall survival of patients with a subsequent malignant neoplasm (SMN) and matched patients with a first malignant neoplasm (FMN) for sarcoma (a) and non-metastatic breast cancer (stage I–III) (b). Patients

with dermatofibrosarcoma protuberans (*n* = 62 FMN), Ewing sarcoma (*n* = 54 FMN), and Kaposi sarcoma (*n* = 18 FMN and *n* = 1 SMN) were excluded in the sarcoma plot

because adjuvant radiotherapy was impossible due to previous chest radiotherapy treatment for Hodgkin lymphoma and two breast-SMN patients had an indication for adjuvant chemotherapy, but were not treated because the maximum cumulative anthracycline dose would be exceeded.

Breast cancer—overall and cause-specific survival

Median follow-up time was 6.3 years (range 0.5–20.3 years) for breast-SMN patients and 7.1 years (range 0.3–27.9 years) for breast-FMN patients. At end of

Table 3 Regression analyses to ascertain the contribution of tumor and treatment characteristics between sarcoma as a subsequent malignant neoplasm (SMN) and sarcoma as a first malignant neoplasm (FMN) with respect to overall and cause-specific mortality

	Overall mortality ^a		Sarcoma-specific mortality ^b		Mortality other causes ^b	
	HR	95% CI	HR	95% CI	HR	95% CI
Univariable SMN versus FMN	2.13	1.41–3.22	2.21	1.45–3.37	1.13	0.14–9.74
Multivariable						
Model 1: adjustment tumor characteristics ^c						
SMN versus FMN	1.83	1.21–2.78	1.90	1.22–2.96	1.06	0.13–8.47
Morphology						
Osteosarcoma/rhabdomyosarcoma	1.00 (ref)		1.00 (ref)		1.00 (ref)	
Chondrosarcoma	0.12	0.05–0.29	0.08	0.03–0.26	0.44	0.05–3.96
Other or unspecified	0.66	0.47–0.92	0.71	0.49–1.01	0.26	0.05–1.38
Model 2: adjustment treatment characteristics ^d						
SMN versus FMN	1.90	1.25–2.90	1.93	1.18–3.18	0.98	0.07–14.54
Surgery (yes vs. no)	2.40 ^e	0.44–13.05 ^e	0.28	0.18–0.45	0.38	0.03–5.44
Radiotherapy (yes vs. no)	1.44	1.01–2.06	1.66	1.13–2.44	0.47	0.05–4.23
Chemotherapy (yes vs. no)	1.88	1.29–2.73	2.12	1.38–3.27	0.44	0.04–5.49
Model 3: adjustment tumor + treatment characteristics ^f						
SMN versus FMN	1.88	1.23–2.87	1.91	1.16–3.13	0.77	0.05–10.97
Surgery (yes vs. no)	2.67 ^g	0.49–14.61 ^g	0.30	0.19–0.48	0.47	0.04–4.98
Radiotherapy (yes vs. no)	1.29	0.90–1.85	1.46	1.00–2.13	0.58	0.06–5.18
Chemotherapy (yes vs. no)	1.68	1.07–2.63	2.13	1.29–3.52	0.13	0.02–0.97
Morphology						
Osteosarcoma/rhabdomyosarcoma	1.00 (ref)		1.00 (ref)		1.00 (ref)	
Chondrosarcoma	0.26	0.10–0.69	0.23	0.07–0.77	0.14	0.01–1.26
Other or unspecified	1.08	0.72–1.64	1.33	0.85–2.06	0.10	0.02–0.60

Patients with dermatofibrosarcoma protuberans ($n = 62$ FMN), Ewing sarcoma ($n = 54$ FMN), and Kaposi sarcoma ($n = 18$ FMN and $n = 1$ SMN) were excluded

CI confidence interval, FMN first malignant neoplasm, HR hazard ratio, SMN subsequent malignant neoplasm

^aCox proportional hazards regression analyses including 28 deaths among sarcoma-SMN patients and 114 deaths among sarcoma-FMN patients

^bCompeting risk regression analyses. When evaluating sarcoma-specific mortality, mortality due to other causes was considered a competing risk and vice versa

^cAdjusted for morphology (osteosarcoma/Ewing sarcoma/rhabdomyosarcoma; chondrosarcoma; other or unspecified)

^dAdjusted for surgery (yes vs. no), radiotherapy (yes vs. no), and chemotherapy (yes vs. no)

^eProportional hazards assumption was violated for category “Surgery (yes vs. no).” At a follow-up duration of 6.41 years (median), the hazard ratio was 0.99 (95% CI: 0.40–2.44) with an increase of a factor 1.58[^](years of follow-up after 6.41 years)

^fAdjusted for morphology, surgery, radiotherapy, and chemotherapy

^gProportional hazards assumption was violated for category “Surgery (yes vs. no).” At a follow-up duration of 6.41 years (median), the hazard ratio was 2.67 (95% CI 0.49–14.61) with an increase of a factor 1.59[^](years of follow-up after 6.41 years)

follow-up, 12 (29.3%) breast-SMN patients were deceased compared to 89 (21.7%) breast-FMN patients (Table 4). For non-metastatic breast cancer (stage I–III), overall survival rates after 5 and 15 years were 93.8% (95% CI 77.4–98.4%) and 63.1% (95% CI 34.9–81.8%) among breast-SMN patients and 88.5% (95% CI 84.7–91.4%) and 77.4% (95% CI 71.5–82.2%) among breast-FMN patients, respectively (Table 2, Fig. 1b). Fifteen-year survival rates among all patients with breast cancer (including metastatic) were 47.6% (95% CI 21.9–69.5%) for breast-SMN and 73.2% (95% CI 67.3–78.1%) for breast-FMN patients.

For non-metastatic breast cancer, breast cancer-specific mortality at 15-year was 33.2% (95% CI 12.2–56.1%) for breast-SMN patients and 19.8% (95% CI 15.0–25.1%) for breast-FMN patients (Table 2). Mortality due to other causes was very low (0–4%). Overall survival did not significantly differ between breast-SMN patients and breast-FMN patients (multivariable HR 1.14; 95% CI 0.54–2.37) (Table 5). Similar trends were observed for breast cancer-specific mortality (multivariable HR 1.09; 95% CI 0.49–2.39).

Table 4 Characteristics of patients with female breast cancer as a subsequent malignant neoplasm (SMN) and matched patients with female breast cancer as a first malignant neoplasm (FMN)

	DCOG-LATER cohort SMN (<i>n</i> = 41)		Cancer registry FMN (<i>n</i> = 410)		<i>p</i> value
	<i>N</i>	%	<i>N</i>	%	
Age at diagnosis ^a					1.000
< 30 years	6	14.6	60	14.6	
30–39 years	23	56.1	230	56.1	
40–49 years	8	19.5	80	19.5	
50+ years	4	9.8	40	9.8	
Incidence year ^a					1.000
1989–1999	7	17.1	70	17.1	
2000–2009	20	48.8	200	48.8	
2010–2014	14	34.2	140	34.2	
Morphology breast cancer					0.379
Ductal	36	87.8	358	87.3	
Lobular	3	7.3	21	5.1	
Mixed ductal/lobular	2	4.9	12	2.9	
Other ^b	0	0.0	19	4.6	
Stage ^c					0.499
I	16	40.0	126	30.9	
II	15	37.5	197	48.3	
III	6	15.0	63	15.4	
IV	3	7.5	22	5.4	
Unknown	1		2		
Grade ^c					0.172
Well differentiated	2	6.9	37	12.2	
Moderately differentiated	15	51.7	103	33.9	
Poorly differentiated	12	41.4	164	54.0	
Unknown	12		106		
ER status ^c					0.202
Negative	5	16.1	83	28.1	
Positive	26	83.9	212	71.9	
Unknown	10		115		
PR status ^c					0.315
Negative	8	28.6	114	38.9	
Positive	20	71.4	179	61.1	
Unknown	13		117		
Her2 receptor status ^{c,e}					0.803
Negative	20	76.9	210	79.0	
Positive	6	23.1	56	21.1	
Unknown	15		144		
Receptor status ^c					0.393
Triple negative	2	7.7	55	20.8	
ER and PR negative, Her2 receptor positive	1	3.9	14	5.3	
ER and/or PR positive, Her2 receptor positive	5	19.2	40	15.2	
ER and/or PR positive, Her2 receptor negative	18	69.2	155	58.7	
Unknown	15		146		
Surgery ^c					0.024
No	4	9.8	13	3.2	
Mastectomy	26	63.4	202	49.3	
Breast-conserving surgery	11	26.8	193	47.1	
Surgery, type unknown	0	0.0	2	0.5	

Table 4 (continued)

	DCOG-LATER cohort SMN (<i>n</i> = 41)		Cancer registry FMN (<i>n</i> = 410)		<i>p</i> value
	<i>N</i>	%	<i>N</i>	%	
Radiotherapy					0.011
No	20	48.8	116	28.3	
Yes	21	51.2	294	71.7	
Chemotherapy					0.022
No	17	41.5	97	23.7	
Yes	24	58.5	313	76.3	
Hormonal therapy					0.870
No	23	56.1	219	53.4	
Yes	18	43.9	191	46.6	
Treatment category					0.011
No surgery ^d	4	9.8	13	3.2	
Mastectomy, no RT or CT (± HT)	7	17.1	23	5.6	
Mastectomy + RT and/or CT (± HT)	19	46.3	179	43.7	
Breast-conserving surgery, no RT or CT (± HT)	0	0.0	2	0.5	
Breast-conserving surgery + RT and/or CT (± HT)	11	26.8	191	46.6	
Surgery, type unknown	0	0.0	2	0.5	
Vital status at end of follow-up					0.325
Alive	29	70.7	321	78.3	
Deceased	12	29.3	89	21.7	

CT chemotherapy, ER estrogen receptor, FMN first malignant neoplasm, HT hormonal therapy, PR progesterone receptor, RT radiotherapy, SMN subsequent malignant neoplasm

^aMatching factors

^bIncludes seven medullary carcinomas, six mucinous adenocarcinomas, two tubular adenocarcinomas, one each of small cell neuroendocrine carcinoma, apocrine adenocarcinoma, medullary carcinoma with lymphoid stroma, and metaplastic carcinoma

^cPercentages and *p* values are based on subjects with known information only

^dAmong breast-SMN cases, two were treated with CT only and 1 each with HT only and no recorded treatment. Among breast-FMN cases, five were treated with CT only, two each with RT + CT, CT + HT, and no recorded treatment, and one each with RT + HT and HT only

^eHer2 receptor status has been systematically ascertained from September 2005 and patients with a positive Her2 receptor status have been treated with Herceptin according to Dutch treatment guidelines since then

Melanoma—comparison of clinical and histopathological characteristics

Of 18 eligible patients with subsequent melanoma (melanoma-SMN) after childhood cancer, one patient was excluded because of a history of radiotherapy and chemotherapy for an intervening ALL diagnosis. We included 17 melanoma-SMN patients and 170 patients with first primary melanoma (melanoma-FMN). More than forty percent of melanoma-SMN cases (41.2%) occurred after childhood leukemia and 23.5% of melanoma-SMN cases had prior radiotherapy to the melanoma region (Supplementary Table 2).

Among patients with melanoma-SMN, 88.2% were diagnosed with superficial spreading melanoma, compared to 68.8% among melanoma-FMN patients. No significant differences between melanoma-SMN and melanoma-FMN

patients were observed for stage distribution and melanoma treatment (Table 1).

Melanoma—overall and cause-specific survival

Median follow-up time was 9.4 years (range 0.9–20.9 years) for melanoma-SMN patients and 9.2 years (range 0.4–20.9 years) for melanoma-FMN patients. At end of follow-up, 11.8% of melanoma-SMN patients were deceased compared to 10.0% of melanoma-FMN patients (Table 1). Fifteen-year survival exceeded 80% in both patients with melanoma-SMN (82%) and melanoma-FMN (87%) (Table 2). Melanoma-specific mortality at 15 year since diagnosis was 12.5% for melanoma-SMN and 10.0% for melanoma-FMN patients. No significant differences were observed between melanoma-SMN and melanoma-FMN patients in overall survival (multivariable

Table 5 Regression analyses to ascertain the contribution of tumor and treatment characteristics between female breast cancer as a subsequent malignant neoplasm (SMN) and female breast cancer as a first malignant neoplasm (FMN) with respect to overall and cause-specific mortality

	Overall mortality ^a		Breast cancer-specific mortality ^b	
	HR	95% CI	HR	95% CI
Univariable SMN versus FMN	1.45	0.79–2.65	1.43	0.79–2.59
Multivariable				
Model 1: adjustment tumor characteristic ^c				
SMN versus FMN	1.32	0.70–2.47	1.30	0.65–2.59
Stage (III/IV vs. I/II)	4.96	3.30–7.45	5.55	3.62–8.53
Model 2: adjustment treatment characteristics ^d				
SMN versus FMN	1.23	0.61–2.47	1.19	0.61–2.32
Surgery type (mastectomy vs. breast-conserving or type unknown)	3.15	1.86–5.34	3.45	1.85–6.41
Radiotherapy (yes vs. no)	2.14	1.23–3.72	2.35	1.24–4.47
Chemotherapy (yes vs. no)	0.79	0.47–1.35	0.86	0.47–1.56
Model 3: adjustment tumor + treatment characteristics ^e				
SMN versus FMN	1.14	0.54–2.37	1.09	0.49–2.39
Stage (III/IV vs. I/II)	2.92	1.62–5.26	3.09	1.39–6.85
Surgery type (mastectomy vs. breast-conserving or type unknown)	1.90	1.02–3.53	2.03	0.87–4.78
Radiotherapy (yes vs. no)	1.30	0.66–2.53	1.40	0.54–3.62
Chemotherapy (yes vs. no)	0.77	0.44–1.33	0.84	0.44–1.60

CI confidence interval, FMN first malignant neoplasm, HR hazard ratio, SMN subsequent malignant neoplasm

^aCox proportional hazards regression analyses including 12 deaths among breast-SMN patients and 89 deaths among breast-FMN patients

^bCompeting risk regression analyses. Mortality due to other causes was considered a competing risk and vice versa. Univariable HR for mortality due to other causes: 10.00 (95% CI 0.64–156.62)

^cAdjusted for stage (III/IV vs. I/II). One breast-SMN patient and two breast-FMN patients with unknown stage were excluded

^dAdjusted for surgery type (mastectomy vs. breast-conserving or type unknown), radiotherapy (yes vs. no), and chemotherapy (yes vs. no). One breast-SMN patient and two breast-FMN patients without surgery were excluded

^eAdjusted for stage, surgery type, radiotherapy, and chemotherapy. Two breast-SMN patients and four first-SMN patients were excluded because of missing stage or no surgical treatment

HR 0.80; 95% CI 0.10–6.21) and melanoma-specific survival (univariable HR 0.71; 95% CI 0.10–5.00) (Supplementary Table 4).

Discussion

In this study, survival and characteristics of three types of solid SMNs from a large cohort of childhood cancer survivors were compared to FMNs of the same type in the general population. Survival of patients with sarcomas after childhood cancer was poor and significantly worse than survival of patients with sarcomas as FMNs. Although we observed that breast-SMN patients more often received a mastectomy without radiotherapy or chemotherapy compared to breast-FMN patients, survival was not different between breast-SMN and breast-FMN patients.

This is one of the few studies to compare SMNs among childhood cancer survivors with FMNs occurring in an age- and diagnosis-era matched population-based sample of cancer patients. We found that overall survival was worse for sarcoma-SMN patients than for sarcoma-FMN patients, both for bone sarcoma and soft-tissue sarcoma. This worse survival is in line with results from two registry-based studies in the United States comparing sarcoma-SMN with sarcoma-FMN patients across patients of all ages [14, 23] and with two studies comparing sarcoma-SMN patients with prior irradiation to sarcoma-FMN patients [24, 25]. We observed a considerable shift in type of sarcoma seen after childhood cancer as SMN vs. those seen as FMN, in particular a shift from Ewing sarcoma and dermatofibrosarcoma in the FMN group to osteosarcoma in the SMN group. The group of sarcoma-SMN patients may harbor more patients with a genetic predisposition than sarcoma-FMN patients, especially those patients with sarcoma-SMNs occurring after retinoblastoma or Li–Fraumeni syndrome-associated tumors [26–28]. The stage distribution was not different between the groups in our study. Robison et al. found that sarcoma-SMN patients who had received prior radiotherapy to the sarcoma region seemed to have a more advanced stage than sarcoma-FMN patients, although this difference was not statistically significant [23]. Similar to what Robison et al. reported, grade seemed to be worse among sarcoma-SMN patients in our study. However, in both studies, the majority of grades were unknown, so no firm conclusions can be drawn [23]. Bjerkehagen et al. showed that the poorer prognosis of sarcomas occurring in previous radiation fields compared to sporadic sarcomas could be explained by more unfavorable factors, such as central tumor site and incomplete surgical remission [24]. Treatment modalities administered for sarcoma treatment did not differ between sarcoma-SMN and sarcoma-FMN patients in our study. Although this may indicate that treatment generally was not influenced by previous childhood cancer treatment, a note of caution is warranted since administration of specific, effective agents (in particular anthracyclines) may

differ between the groups. This level of detail is not available for the cancer registry sample of FMN patients. Also, the differences with respect to sarcoma subtypes between the groups have somewhat limited the clinical impact of our comparisons.

Breast-SMN and breast-FMN patients generally had comparable tumor characteristics. No differences between the two groups were observed with regard to stage and grade distribution, in accordance with two small studies among HL survivors [13, 29]. In contrast to our findings, two large registry-based studies in AYA cancer survivors did report that AYA cancer/HL survivors with breast SMNs were significantly more likely to have estrogen- and progestogen-negative breast cancer compared to breast-FMN patients [16, 17]. The percentage of triple negative breast cancer in our study was lower than that reported in a retrospective case series of breast cancer patients with a past history of chest radiotherapy (7.7% vs. 29.2%) [30]. An important observation in our study was the higher proportion of breast-SMN patients who had received a mastectomy compared to breast-FMN patients, in particular the proportion of mastectomy without radiotherapy or chemotherapy (17% vs. 6%), as reported earlier [10, 13, 16, 17, 29, 31]. Prior chest radiotherapy treatment for childhood cancer may have limited the adjuvant treatment options after breast-conserving surgery and therefore mastectomy may have been the treatment of choice. In one breast-SMN patient with a mastectomy without radiotherapy or chemotherapy, we found evidence that indeed prior childhood cancer radiotherapy led to the choice for mastectomy over breast-conserving surgery. We found no significant difference in overall survival between breast-SMN and breast-FMN patients. However, patients with breast-SMN seemed to have slightly higher late cumulative mortality rates (10-year and 15-year) than breast-FMN patients, although these differences were not statistically significant. Although the high percentage of breast cancer as the cause of death among deceased patients is suggestive of breast-SMN patients experiencing more often late recurrences/metastases than breast-FMN patients, numbers were too small to interpret. Previous registry-based follow-up studies in Hodgkin lymphoma survivors or in childhood and young adult cancer survivors did find a significantly increased mortality risk in breast-SMN patients compared to breast-FMN patients [14, 16, 17, 29, 31]. Three smaller single-institution studies among (Hodgkin) lymphoma survivors showed mixed results, with one also showing significantly worse survival in breast-SMN patients [18], while the other study did not find a difference between breast-SMN and breast-FMN patients [10, 13]. We observed that the large majority of breast-SMN patients died because of the breast cancer, in contrast to Milano et al. who found that many of the breast-SMN patients after Hodgkin lymphoma died of other cancers, heart disease, or other causes [16].

The high risks of other fatal late adverse effects among patients with Hodgkin lymphoma treated with high dose, large volume radiotherapy, and/or high doses of chemotherapy [32] can probably not be extrapolated to that in a mixed cohort of childhood cancer survivors with a large proportion of breast cancers in non-irradiated survivors. Compared to Hodgkin lymphoma survivors, the group of childhood cancer survivors harbors a subpopulation of individuals with *TP53*-related or other genetic predisposition profiles that predispose to breast cancer. Therefore, it is possible that the related breast tumors are biologically different as well, although this a mere hypothesis at this point. Because breast cancers in the breast-SMN group were diagnosed at fairly young ages, and thus matched breast-FMN group represents a sample of young patients, it is likely that the breast-FMN group includes several women with a genetic predisposition, such as BRCA mutation carriers. Overall survival among young individuals with melanoma is very good, with few fatalities among SMN and FMN cases alike. Melanoma-SMN patients harbor no worse prognosis profile vs. melanoma-FMN patients and there are no strong predictors of mortality. Keegan et al. did find a slight significantly worse overall survival in melanoma-SMN compared to melanoma-FMN [14]. Previous studies have shown that childhood cancer survivors are at increased risk of skin cancers, mainly basal cell carcinomas, occurring mainly in skin surface areas that had received radiation [33, 34]. Survivors who received radiotherapy should be educated to be on alert for local skin changes in order to timely seek medical attention for skin lesions that may be potentially malignant.

Strengths of our study are the availability of clinical, histopathological, and treatment data of solid SMNs from a large, nationwide cohort of childhood cancer survivors and a matched population-based comparison group of FMNs. Furthermore, we were able to not only compare survival, but also causes of death, which enabled us to evaluate whether survival differences were caused by differences in cancer-specific mortality or mortality from other causes. A limitation was the relatively small numbers of site-specific SMNs included in our study, which may have limited the power to detect differences in clinical characteristics and survival between the cancer groups. Furthermore, for several tumor characteristics (e.g., sarcoma stage and grade and breast cancer receptor status information), there was a relatively high percentage of the data missing, in particular for cases diagnosed in the distant past, when cancer registration (and in fact clinical practice) did not include such detail. This omission limits impact of these findings.

In conclusion, our exploratory analyses reveal that survival of sarcoma-SMN patients is poor, with less than one in three surviving 15 years after sarcoma-SMN diagnosis, and is worse than survival of sarcoma-FMN patients. This observation seems partly related to differences in sarcoma

subtypes. In our study, breast cancer survival and breast tumor characteristics are not different between breast-SMN and breast-FMN patients, but treatment differs as breast-SMN patients receive more often a mastectomy than breast-FMN patients. Survival of melanoma-SMN is favorable. Further studies including more cases should elaborate on possible explanations of a worse survival of (soft tissue) sarcoma-SMN patients and on treatment strategies and underlying deliberations among breast-SMN patients.

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Compliance with ethical standards

Conflict of interest The authors declare that they have no conflict of interest.

Research involving human rights Data collection for the study was exempted from review by the Institutional Review Boards of all participating centers and all data were processed and analyzed anonymously.

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
References

- Bhakta N, Liu Q, Ness KK, Baassiri M, Eissa H, Yeo F, Chemaitilly W, Ehrhardt MJ, Bass J, Bishop MW, Shelton K, Lu L, Huang S, Li Z, Caron E, Lanctot J, Howell C, Folse T, Joshi V, Green DM, Mulrooney DA, Armstrong GT, Krull KR, Brinkman TM, Khan RB, Srivastava DK, Hudson MM, Yasui Y, Robison LL (2017) The cumulative burden of surviving childhood cancer: an initial report from the St Jude Lifetime Cohort Study (SJLIFE). *Lancet* 390(10112):2569–2582
- Geenen MM, Cardous-Ubbink MC, Kremer LC, van den Bos C, van der Pal HJ, Heinen RC, Jaspers MW, Koning CC, Oldenburger F, Langeveld NE, Hart AA, Bakker PJ, Caron HN, van Leeuwen FE (2007) Medical assessment of adverse health outcomes in long-term survivors of childhood cancer. *JAMA* 297(24):2705–2715
- Hudson MM, Ness KK, Gurney JG, Mulrooney DA, Chemaitilly W, Krull KR, Green DM, Armstrong GT, Nottage KA, Jones KE, Sklar CA, Srivastava DK, Robison LL (2013) Clinical ascertainment of health outcomes among adults treated for childhood cancer. *JAMA* 309(22):2371–2381
- Oeffinger KC, Mertens AC, Sklar CA, Kawashima T, Hudson MM, Meadows AT, Friedman DL, Marina N, Hobbie W, Kadan-Lottick NS, Schwartz CL, Leisenring W, Robison LL, Childhood Cancer Survivor S (2006) Chronic health conditions in adult survivors of childhood cancer. *N Engl J Med* 355(15):1572–1582
- Armstrong GT, Liu Q, Yasui Y, Neglia JP, Leisenring W, Robison LL, Mertens AC (2009) Late mortality among 5-year survivors of childhood cancer: a summary from the Childhood Cancer Survivor Study. *J Clin Oncol* 27(14):2328–2338
- Fidler MM, Reulen RC, Winter DL, Kelly J, Jenkinson HC, Skinner R, Frobisher C, Hawkins MM, British Childhood Cancer Survivor Study Steering G (2016) Long term cause specific mortality among 34 489 five year survivors of childhood cancer in Great Britain: population based cohort study. *BMJ* 354:i4351
- Reulen RC, Winter DL, Frobisher C, Lancashire ER, Stiller CA, Jenney ME, Skinner R, Stevens MC, Hawkins MM, British Childhood Cancer Survivor Study Steering G (2010) Long-term cause-specific mortality among survivors of childhood cancer. *JAMA* 304(2):172–179
- Teepen JC, van Leeuwen FE, Tissing WJ, van Dulmen-den Broeder E, van den Heuvel-Eibrink MM, van der Pal HJ, Loonen JJ, Bresters D, Versluys B, Neggers S, Jaspers MWM, Hauptmann M, Heiden-van der Loo M, Visser O, Kremer LCM, Ronckers CM (2017) Long-Term Risk of Subsequent Malignant Neoplasms After Treatment of Childhood Cancer in the DCOG LATER Study Cohort: Role of Chemotherapy. *J Clin Oncol* 35(20):2288–2298
- Turcotte LM, Liu Q, Yasui Y, Arnold MA, Hammond S, Howell RM, Smith SA, Weathers RE, Henderson TO, Gibson TM, Leisenring W, Armstrong GT, Robison LL, Neglia JP (2017) Temporal trends in treatment and subsequent neoplasm risk among 5-year survivors of childhood cancer, 1970–2015. *JAMA* 317(8):814–824
- Alm El-Din MA, Hughes KS, Raad RA, Goldberg SI, Aisenberg AC, Niemierko A, Taghian AG (2009) Clinical outcome of breast cancer occurring after treatment for Hodgkin's lymphoma: case-control analysis. *Radiat Oncol* 4:19
- Chowdhry AK, Fung C, Chowdhry VK, Bergsma D, Dhakal S, Constine LS, Milano MT (2018) A population-based study of prognosis and survival in patients with second primary thyroid cancer after Hodgkin lymphoma. *Leuk Lymphoma* 59(5):1180–1187
- Goldfarb M, Freyer DR (2014) Comparison of secondary and primary thyroid cancer in adolescents and young adults. *Cancer* 120(8):1155–1161
- Janov AJ, Tulecke M, O'Neill A, Lester S, Mauch PM, Harris J, Schnitt SJ, Shapiro CL (2001) Clinical and pathologic features of breast cancers in women treated for Hodgkin's disease: a case-control study. *Breast J* 7(1):46–52

14. Keegan THM, Bleyer A, Rosenberg AS, Li Q, Goldfarb M (2017) Second primary malignant neoplasms and survival in adolescent and young adult cancer survivors. *JAMA Oncol* 3(11):1554–1557
15. Milano MT, Li H, Constine LS, Travis LB (2011) Survival after second primary lung cancer: a population-based study of 187 Hodgkin lymphoma patients. *Cancer* 117(24):5538–5547
16. Milano MT, Li H, Gail MH, Constine LS, Travis LB (2010) Long-term survival among patients with Hodgkin's lymphoma who developed breast cancer: a population-based study. *J Clin Oncol* 28(34):5088–5096
17. Sadler C, Goldfarb M (2015) Comparison of primary and secondary breast cancers in adolescents and young adults. *Cancer* 121(8):1295–1302
18. Sanna G, Lorizzo K, Rotmensz N, Bagnardi V, Cinieri S, Colletoni M, Nole F, Goldhirsch A (2007) Breast cancer in Hodgkin's disease and non-Hodgkin's lymphoma survivors. *Ann Oncol* 18(2):288–292
19. Youn P, Li H, Milano MT, Stovall M, Constine LS, Travis LB (2013) Long-term survival among Hodgkin's lymphoma patients with gastrointestinal cancer: a population-based study. *Ann Oncol* 24(1):202–208
20. Forman D, Bray F, Brewster DH, Gombe Mbalawa C, Kohler B, Piñeros M, Steliarova-Foucher E, Swaminathan R, Ferlay J (2014) Cancer incidence in five continents. Vol. X. IARC Scientific Publications No. 164. Lyon: International Agency for Research on Cancer
21. Fine JP, Gray RJ (1999) A proportional hazards model for the subdistribution of a competing risk. *J Am Stat Assoc* 94(446):496–509
22. Criscione VD, Weinstock MA (2007) Descriptive epidemiology of dermatofibrosarcoma protuberans in the United States, 1973 to 2002. *J Am Acad Dermatol* 56(6):968–973
23. Robinson E, Bar-Deroma R, Rennert G, Neugut AI (1992) A comparison of the clinical characteristics of second primary and single primary sarcoma: a population based study. *J Surg Oncol* 50(4):263–266
24. Bjerkeheggen B, Smastuen MC, Hall KS, Skjeldal S, Smeland S, Fossa SD (2012) Why do patients with radiation-induced sarcomas have a poor sarcoma-related survival? *Br J Cancer* 106(2):297–306
25. Gladdy RA, Qin LX, Moraco N, Edgar MA, Antonescu CR, Alektiar KM, Brennan MF, Singer S (2010) Do radiation-associated soft tissue sarcomas have the same prognosis as sporadic soft tissue sarcomas? *J Clin Oncol* 28(12):2064–2069
26. Bright CJ, Hawkins MM, Winter DL, Alessi D, Allodji RS, Bagnasco F, Bardi E, Bautz A, Byrne J, Feijen EAM, Fidler MM, Garwicz S, Grabow D, Gudmundsdottir T, Guha J, Haddy N, Jankovic M, Kaatsch P, Kaiser M, Kuehni CE, Linge H, Ofstaa H, Ronckers CM, Skinner R, Teepen JC, Terenziani M, Vu-Bezin G, Wesenberg F, Wiebe T, Sacerdote C, Jakab Z, Haupt R, Lahteenmaki P, Zaletel LZ, Kuonen R, Winther JF, de Vathaire F, Kremer LC, Hjorth L, Reulen RC, PanCareSurFup C (2018) Risk of Soft-Tissue Sarcoma Among 69 460 Five-Year Survivors of Childhood Cancer in Europe. *J Natl Cancer Inst* 110(6):649–660
27. Kleinerman RA, Tucker MA, Abramson DH, Seddon JM, Tarone RE, Fraumeni JF Jr (2007) Risk of soft tissue sarcomas by individual subtype in survivors of hereditary retinoblastoma. *J Natl Cancer Inst* 99(1):24–31
28. Li FP, Fraumeni JF Jr (1969) Soft-tissue sarcomas, breast cancer, and other neoplasms. A familial syndrome? *Ann Intern Med* 71(4):747–752
29. Veit-Rubin N, Rapiti E, Usel M, Benhamou S, Vinh-Hung V, Vlastos G, Bouchardy C (2012) Risk, characteristics, and prognosis of breast cancer after Hodgkin's lymphoma. *Oncologist* 17(6):783–791
30. Demoor-Goldschmidt C, Supiot S, Mahe MA, Oberlin O, Allodji R, Haddy N, Helfre S, Vigneron C, Brillaud-Meflah V, Bernier V, Laprie A, Ducassou A, Claude L, Diallo I, de Vathaire F (2018) Clinical and histological features of second breast cancers following radiotherapy for childhood and young adult malignancy. *Br J Radiol* 91(1086):20170824
31. Elkin EB, Klem ML, Gonzales AM, Ishill NM, Hodgson D, Ng AK, Marks LB, Weidhaas J, Freedman GM, Miller RC, Constine LS, Myrehaug S, Yahalom J (2011) Characteristics and outcomes of breast cancer in women with and without a history of radiation for Hodgkin's lymphoma: a multi-institutional, matched cohort study. *J Clin Oncol* 29(18):2466–2473
32. van Leeuwen FE, Ng AK (2017) Late sequelae in Hodgkin lymphoma survivors. *Hematol Oncol* 35(Suppl 1):60–66
33. Perkins JL, Liu Y, Mitby PA, Neglia JP, Hammond S, Stovall M, Meadows AT, Hutchinson R, Dreyer ZE, Robison LL, Mertens AC (2005) Nonmelanoma skin cancer in survivors of childhood and adolescent cancer: a report from the childhood cancer survivor study. *J Clin Oncol* 23(16):3733–3741
34. Teepen JC, Kok JL, Kremer LC, Tissing WJE, van den Heuvel-Eibrink MM, Loonen JJ, Bresters D, van der Pal HJ, Versluis B, van Dulmen-den Broeder E, Nijsten T, Hauptmann M, Hollema N, Dolsma WV, van Leeuwen FE, Ronckers CM, Group D-LS (2019) Long-term risk of skin cancer among childhood cancer survivors: A DCOG-LATER Cohort Study. *J Natl Cancer Inst*. <https://doi.org/10.1093/jnci/djy212>

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