Obesity and Sarcopenia in Survivors of Childhood Acute Lymphoblastic Leukemia

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Objective: To describe the prevalence of obesity and sarcopenia among survivors of childhood acute lymphoblastic leukemia (ALL) using DEXA scan, and study associated risk factors. Methods: This case control study was conducted between July, 2013 and June, 2014 at a tertiary care cancer centre in India. Study participants included 65 survivors of childhood ALL who were <18 years of age at diagnosis, treated between years 1996 and 2008, and were at least two years since completion of therapy. The controls included 50 matched siblings. Dual energy X-ray absorptiometry (DEXA) was used to study the body composition (body fat percentage, BF% and lean body mass, LBM) of the participants and controls. McCarthy’s body fat reference data were applied and logistic regression analysis was used to study various risk factors. Results: At a median (range) follow-up of 5 (7.2-17.2) years, BF% (DEXA) identified a significantly higher prevalence of obesity of 21.5% (14/65) and sarcopenic obesity (14%) among survivors as compared to the controls (0/50, P<0.001), while the prevalence of sarcopenia as detected by LBM was similar at 60% (39/65) and 56% (28/50), respectively. On multivariate analysis, age at evaluation, high-risk disease and cranial irradiation were independently associated with high likelihood of obesity, while none of the factors predicted sarcopenia. Conclusion: High prevalence of obesity and sarcopenic obesity were observed among survivors of childhood ALL. Keywords: Body composition, Body fat, DEXA, Lean body mass, Metabolic syndrome.

Obesity is recognized as a common chronic health problem in childhood cancer survivors, and cardiovascular disease has been shown to occur at an earlier age in survivors of childhood cancer [1-4]. As acute lymphoblastic leukemia (ALL) is the commonest childhood cancer, we aimed to study the prevalence of obesity and sarcopenia among survivors of childhood ALL using DEXA scan, as compared to sibling controls, and evaluate association with risk factors.

METHODS

This study was conducted between July, 2013 and June, 2014 at the pediatric hemato-oncology department of our comprehensive cancer center. The study participants (cases) included survivors of childhood ALL who were less than 18 years of age at diagnosis, treated between 1996 and 2008, in first complete remission, and were at least two years after completion of therapy. Of the total 207 ALL patients that were treated, 122 were eligible for the study and we were able to contact and obtain consent from 65 children and young adult survivors (52 males). The treatment protocol was based on a Berlin Frankfurt Munster (BFM) backbone that included a four-drug induction and prednisolone as the glucocorticoid [5]. Their results were compared with 50 healthy sibling controls that were matched for age (±1 year) and sex. The controls were determined to be healthy and eligible for the study if there was no evidence of any medical illness on detailed history and physical examination. The study was approved by the institutional ethics committee and conducted after obtaining informed consent from the study participant or parents (in case of minor).

Details of method of measuring anthropometric indices (weight, height, BMI) and physical activity quotient (PAQ) have been previously published [5]. Three-compartment body composition was assessed using DEXA and Hologic Explorer (S/N 91531) software version 13.3.3. The DEXA machine directly generates absolute values as well as Z scores for bone mineral content, and lean body mass (LBM) measures, as the National Health and Nutrition Examination Survey (NHANES) reference data is integrated into the software.

Study participants’ body mass index (BMI) status was determined by using WHO BMI growth charts and
categorized as normal weight (BMI >5th to <85th percentile), overweight (>85th to <95th percentile), or obese (BMI ≥95th percentile) [6,7]. High adiposity was defined as body fat percentage (BF%) levels higher than the 85th percentile of McCarthy BF% reference data (NHANES) for each age-sex group (overweight ≥85th percentile and obesity ≥95th percentile) [8,9]. Sarcopenia was defined as LBM <5th percentile of reference data (NHANES). Sarcopenic obesity was defined as participants fulfilling criteria for sarcopenia and obesity by BF% and LBM [8]. Body fat mass index (BFMI) and lean body mass index (LBMI) were calculated from DEXA-measured body-composition data as BF or LBM in kg per m² height.

Statistical analyses: The body compositions of two groups such as weight, height, BMI, BF % etc. were compared using independent t-test and chi-square test. Analysis of variance (ANOVA) was used to compare delta BMI change among thin, average and overweight patients at diagnosis. Univariate and multivariate logistic regressions were used to study the influence of various demographic and disease-related factors for obesity and sarcopenia among the survivors. The results were interpreted using odds ratio (OR) and 95% confidence interval (CI). The Statistical Package for Social Sciences for Window’s software (IBM SPSS Statistics version 23.0) was used for all analyses. Significance was set at P<0.05.

RESULTS

At the time of evaluation, the 65 study participants [median (range) age, 15 (7.7-27.5) years] were median (range) 4.3 (2-14.8) years from treatment completion, and had a median (range) follow-up of 7.2 (5-17.2) years.

The values for body composition for the survivors and the control groups are provided in Table I. The mean (SD) BF% was significantly higher among ALL survivors as compared to the controls [35.2 (7.4) vs 30.2 (8.0); P=0.001], and a similar trend was observed when analyzed by gender. Using BF%, obesity was observed among 21.5% (14/65) of study participants and none of the controls (P=0.00) and overweight among 55% (36/65) and 48% (24/50) of cases and controls respectively (P=0.7). On the other hand BMI under-estimated obesity (6%, 4/65, P=0.02) and overweight (26%, 17/65, P=0.01) among the study participants. It also detected fewer control participants to be overweight (6%, 3/50, P=0.01).

We further looked at WHO BMI z-scores at diagnosis and noted that 10/65 (15%) were underweight/thin, 9/65 (14%) were overweight, and none was obese. While at evaluation, these proportions changed to 26.1% (n=17) overweight and 6.1% (n=4) obese, respectively. No participant was underweight/thin. Delta BMI change was calculated from baseline BMI percentile, which revealed that the highest mean (SD) delta change occurred among thin [2067 (882)] followed by normal [273 (50)] and overweight [45 (30)] patients, (P<0.001).

The absolute values for LBM and LBMI for the study participants’ were lower than the controls but this was not statistically significant (P=0.38, 0.68) (Table I). The prevalence of sarcopenia was similar in both the groups (39/65, 60%; 28/50, 56%), respectively (P=0.10). However, the female survivors (11/13, 85%) were more sarcopenic compared to their male counterparts (28/52, 54%) (P=0.05). Sarcopenic obesity was observed among 14% (9/65) of study participants and none of the controls (P=0.00). Of these 9 patients, 1 was thin and none were overweight or obese based on baseline BMI, this difference was not statistically significant (P=0.29).

Distribution of demographic, disease and treatment exposure among participants who were obese or sarcopenic is displayed in Table II. The study participants had a 28 times higher odds of being obese compared to the controls [OR (95% CI) 28 (1.7 to 490); P=0.002]. On univariate logistic regression analysis female gender, age at diagnosis less than 10 years, T immunophenotype, high NCI risk, receiving cranial irradiation, more than 5 years since therapy, younger age at evaluation, being sarcopenic, and PAQ>2 were significantly associated with obesity. On multi-variate analysis, only female gender [OR (95% CI) 7.3 (1.1-50.1); P=0.04], high NCI risk category [OR (95% CI) 6.7 (1.1-43.5); P=0.04], cranial

| Table I Body Composition by DEXA Scan and Anthropometry of Survivors of Childhood Acute Lymphoblastic Leukemia and Sibling Controls (N=115) |
|-----------------|-----------------|---------|---|
| Characteristics                                      | ALL survivors | Controls | P value |
| Weight, kg                                            | 50.9 (16)      | 48.5 (16.3) | 0.42 |
| Height, cm                                             | 154.6 (15.3)  | 156 (14.9) | 0.61 |
| BMI, kg/cm²                                            | 20.3 (3.7)     | 19.3 (3.8) | 0.03 |
| Body fat, %                                            | 35.2 (7.4)     | 30.2 (8.0) | <0.001 |
| BFMI                                                    | 7.26 (2.4)     | 5.8 (2.3) | <0.001 |
| Lean body mass                                         | 30355.9 (9886.1) | 32092.7 (10446.4) | 0.38 |
| LBMI                                                   | 12.3 (2.1)     | 12.5 (2.5) | 0.68 |
| Trunk/Leg fat %                                        | 0.9 (0.6)      | 0.8 (0.5) | 1.0 |
| Trunk lean mass                                        | 14861 (5000.9) | 15835.93 (5509.8) | 0.33 |

All values in mean (SD). DEXA: Dual energy X-ray absorptiometry; ALL: Acute lymphoblastic leukemia; BMI: Body mass index at evaluation; BF%: Body fat percentage; BFMI: Body fat mass index; LBMI: Lean body mass index.
irradiation [OR (95% CI) 9.9 (1.2-83.3); \( P = 0.04 \)], and younger age [OR (95% CI) 10.2 (1.1-91.4); \( P = 0.04 \)], had a significant association with obesity.

Among 31 children exposed to cranial RT, doses of 12.6 Gy were administered to 27 of whom one was obese. Four children received cranial RT at doses of 18 Gy and three of them were obese. None of the four obese survivors suffered from endocrinopathies or short stature to account for obesity. None of the baseline characteristics were associated with higher prevalence of sarcopenia.

### DISCUSSION

In the present analysis, using DEXA we observed that more than one-fifth of the survivors of childhood ALL were obese and half of them were overweight at a median follow-up of 7 years. A wide variation in prevalence rates of obesity (18-80%) has been reported among childhood cancer survivors in studies using DEXA scans. The St Jude life time cohort study [10] reported obesity in 63% male and 85% female ALL survivors at a mean follow up of 25 years, while Barr, et al. [11] from Canada reported obesity and overweight rates of 12% and 18% respectively at a median follow-up of 21 years. The range in findings may be attributed to differences in definitions of obesity, treatment protocol and gluco-corticoid doses, duration of follow-up, ethnicity, and social factors as well as the prevalence of obesity in the normal population of the region. Comparison with a control population helped obviate many of these factors. The main advantage of using sibling controls was to avoid confounding biases due to constitutional and environmental factors.

Published studies form India have used weight- and height-based indices and reported lower prevalence rates of obesity (2.5-12%) and overweight (19-20%) [12,13]. In order to be comparable with these data, we adopted the BMI criteria and observed similar rates of obesity (6%) and overweight (24%) among our ALL survivors. Hence, BMI underestimated the prevalence of adiposity as it is unable to identify normal and underweight individuals with high body fat. Blijorg, et al. [14] used total fat percentage as the gold standard, and reported that 42% of male survivors and 65% of female survivors were misclassified as non-obese using BMI [9].

It is noteworthy that the prevalence of obesity and overweight was not influenced by the study participants’ nutritional status at diagnosis, since the thin and normally nourished children treated for ALL were equally predisposed. However, we observed that the delta change in BMI was highest amongst those who were under-weight/thin at diagnosis, and they probably require close monitoring during follow-up.

Various investigators have described muscle mass loss during the treatment for ALL and its progression throughout therapy and after treatment completion [15,16]. This is attributed mainly to the degradation and decreased synthesis of myosin heavy chains, and steroid use, which causes increased glycogen and lipid levels in muscle cells. The prevalence of sarcopenia was equally high amongst our control population, which possibly indicates that ethnically, our population has lower muscle mass compared to Western counterparts. We did; however, observe a high prevalence of sarcopenic obesity.
(14%) among the survivors that was not seen among the controls. This assumes importance in the context of current literature that highlights the combination of the two to be more detrimental and an important contributor to the development of metabolic syndrome [16].

The observed gender difference has previously also been reported [10,11,17]. Hyperleptinemia, which occurs in girls during puberty, has been linked to body fat and has been described as a possible mechanism for obesity. Although many investigators have reported the association of cranial RT with obesity, it remains a controversial point [17]. Disturbances influencing the satiety centre and dysfunction of hypothalamic-pituitary axis have been found to cause obesity as well. However, with modern treatment protocols, wherein smaller doses of radiation are delivered with better techniques, recent papers have revealed no association of cranial radiation with the incidence of obesity in survivors of childhood ALL [18]. The increased incidence of obesity among children with NCI high risk disease status may be attributed to the use of higher doses of glucocorticosteroid therapy, poor physical activity and use of cranial radiation in this subset.

The results of our analysis should be interpreted in light of the small sample size and skewed gender ratio, in addition to the fact that influence of unknown psychosocial factors on the controls could not be completely excluded. However, our use of DEXA to accurately identify adiposity and sarcopenia as well as use of matched controls strengthens our findings.

The findings of the present study highlight the high prevalence of obesity and sarcopenic obesity in our population of survivors of childhood ALL. Since these are believed to be forerunners of cardio-metabolic syndrome our results emphasize the need for early recognition and aggressive preventive strategies. Larger interventional studies may identify strategies that have an impact on reducing obesity in this sub-population of children.

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**REFERENCES**

1. Tamilselvan St, Scott JX, Sneha L, Divyalakshmi J. Metabolic syndrome in childhood cancer survivors. J Clin Oncol. 2017;35:145.
2. Chueh HW, Yoo JH. Metabolic syndrome induced by anticancer treatment in childhood cancer survivors. Ann Endocrinol Metab. 2017;22:82-9.
3. Barnea D, Raghunathan N, Friedman DN, Tonorezos ES. Obesity and metabolic disease after childhood cancer. Oncology (Williston Park). 2015;29:849-55.
4. Commission on ending childhood obesity: Taking action on childhood obesity report. 2018;WHO/NMH/PND/ECHO/18.1.
5. Jain S, Jain S, Kapoor G, Virmani A, Bajpai R. No impact of disease and its treatment on bone mineral density in survivors of childhood acute lymphoblastic leukemia. Pediatr Blood Cancer. 2017;64.
6. World Health Organization. World Health Organization Child Growth Standards. 2006. Accessed March 5, 2012. https://www.who.int/childgrowth/en/
7. Krebs NF, Himes JH, Jacobson D, et al. Assessment of child and adolescent overweight and obesity. Pediatrics. 2007;120:S193-228.
8. McCarthy HD, Cole TJ, Fry T, Jebb SA, Prentice AM. Body fat reference curves for children. Int J Obes (Lond). 2006;30:598-602.
9. Ogden CL, Li Y, Freedman DS, Borrud LG, Flegal KM. Smoothed percentage body fat percentiles for U.S. children and adolescents, 1999-2004. Natl Health Stat Report. 2011;1-7.
10. Karlage RE, Wilson CL, Zhang N, et al. Validity of anthropometric measurements for characterizing obesity among adult survivors of childhood cancer: A report from the St. Jude Lifetime Cohort Study. Cancer. 2015;121:2036-43.
11. Marriott CJ, Beaumont LF, Farncombe TH, et al. Body composition in long-term survivors of acute lymphoblastic leukemia diagnosed in childhood and adolescence: A focus on sarcopenic obesity. Cancer. 2018;124:1225-31.
12. Prasad M, Arora B, Chinnaswamy G, et al. Nutritional status in survivors of childhood cancer: Experience from Tata Memorial Hospital, Mumbai. Indian J Cancer. 2015;52:219-23.
13. Mohapatra S, Bansal D, Bhalia AK, et al. Is there an increased risk of metabolic syndrome among childhood acute lymphoblastic leukemia survivors? A developing country experience. Pediatr Hematol Oncol. 2016;33:136-49.
14. Blijdorp K, van den Heuvel-Eibrink MM, Pieters R, et al. Obesity is underestimated using body mass index and waist-hip ratio in long-term adult survivors of childhood cancer. PLoS One. 2012;7:e43269.
15. Orgel E, Mueske NM, Sposto R, et al. Limitations of body mass index to assess body composition due to sarcopenic obesity during leukemia therapy. Leuk Lymphoma. 2018;59:138-45.
16. Rayar M, Webber CE, Nayiager T, Sala A, Barr RD. Sarcopenia in children with acute lymphoblastic leukemia. J PediatrHematol Oncol. 2013;35:98-102.
17. Garmey EG, Liu Q, Sklar CA, et al. Longitudinal changes in obesity and body mass index among adult survivors of childhood acute lymphoblastic leukemia: A report from the Childhood Cancer Survivor Study. J Clin Oncol. 2008;26:4639-45.
18. Withycombe JS, Post-White JE, Meza JL, et al. Weight patterns in children with higher risk ALL: A report from the Children’s Oncology Group (COG) for CCG 1961. Pediatr Blood Cancer. 2009;53:1249-54.