Impact of Pediatric Hematopoietic Stem-Cell Transplantation on Craniofacial Growth

Alexandre Viana Frascino, Marcelo Fava, Maria Dulce Silveira Collassanti, Vicente Odone-Filho

INTRODUCTION

Hematopoietic stem-cell transplantation (HSCT) has become one of the most important approaches for pediatric patients with onco-hematological malignancies (1-3). HSCT is preceded by intensive myeloablative conditioning, isolated or combined total body irradiation, and chemotherapy (4). It is estimated that 10,000 transplants are performed annually worldwide (5).

Although HSCT for pediatric patients has been associated with increased survival rates, several long-term complications have been reported (6,7). Endocrine, cardiopulmonary, gastrointestinal, hepatic, renal, neurological, and skeletal impairments clinically manifest as low stature, disproportionality growth, osteoporosis, increased risk of bone fractures, diabetes mellitus, delayed sexual maturation, and cognitive deficit (8,9).

Impaired skeletal growth and development have been associated to reduce bone mineral density in the femur, vertebrae, and jaws (10-13). Craniofacial and dental development disturbances have been described to be more prevalent in children submitted to HSCT at a young age (<10 years) (14,16,24). Impaired skeletal growth in long-term childhood HSCT survivors negatively affects their quality of life (6,12,13). The present retrospective study aimed to quantitatively assess craniofacial growth in HSCT children in comparison with age-sex matched-paired controls.

METHODS

A case-controlled retrospective comparison of the craniofacial growth in 25 HSCT children and 25 matched-paired controls was conducted. Craniofacial growth was quantitatively assessed by linear and angular measurements in panoramic radiographic images using ImageJ (1). Stature growth and body weight were obtained through physical examination. Cancer diagnosis, myeloablative conditioning, and HSCT were retrieved from medical records.

RESULTS

Patients aged 12.2 years (±3.8; 16 male, 9 female). Radiographic images were obtained on an average of 2.43 (±2.0) years after HSCT. The main malignant diagnosis was acute lymphoblastic leukemia (56%), followed by acute myeloid leukemia (36%) and myelodysplastic syndromes (8%). Total body irradiation was associated with chemotherapy at 80%. Mean age at transplantation was 10 (±4.7) years. HSCT survivors showed reduced a vertical growth of the mandibular ramus (p=0.003). This persisted among individuals below 12 years of age (p=0.017). The HSCT group showed delayed dental eruption, though there was no statistically significant difference (p=0.3668). The HSCT group showed stature deficit, increased weight, and body mass index (Z-score stature: -0.28; Z-score weight: 0.38, respectively).

CONCLUSIONS

Pediatric HSCT has decreased vertical craniofacial growth compared to their matched controls. There might be an association between reduced craniofacial vertical growth and reduced estature growth. Further studies to quantitatively investigate the impact of different myeloablative regimens in craniofacial skeletal growth and development.

KEYWORDS: Hematopoietic Stem-Cell Transplantation; Pediatrics; Bone Marrow Transplantation; Skull/Growth and Development; Facial Bones/Growth and Development.

OBJECTIVES: To assess the craniofacial skeletal growth in pediatric hematopoietic stem-cell transplantation (HSCT) survivors in comparison with age-sex matched-paired controls.

MATERIALS AND METHODS

Between 2015 and 2018, long-term pediatric HSCT survivors were selected from Instituto do Tratamento do Câncer Infantil, Instituto da Criança, Hospital das Clínicas da Faculdade de Medicina da Universidade de São Paulo.

The case group (HSCT) inclusion criteria were as follows: 1) HSCT for the treatment of hematological malignancies; 2) age at the time of HSCT ≤18 years; and 3) panoramic radiograph taken at least 6 months after HSCT. The exclusion criteria were...
as follows: 1) diagnosis of skeletal disorders; 2) previous orthodontic treatment; 3) the presence of orthodontic appliances, and 4) history of craniofacial trauma. The control group (CONTROL) individuals were selected matched by age-sex from Instituto de Ciências e Tecnologia, Faculdade de Odontologia, UNESP – São José dos Campos – São Paulo – Brazil.

Information regarding date of birth, sex, height, weight, medical history, and myeloablative conditioning regimen was retrieved from the medical records. Oral and maxillofacial health information was retrieved from the dental records.

Panoramic radiographs were obtained at Instituto de Radiologia, InRad-HC-FMUSP, using the Orthophos CD (Siemens, Bensheim, Germany) with imaging settings of 60–90 kVp, 9–12 mAs and 12 s of exposure. All radiographic images were downloaded with 256 gray levels, 3188 x 1709 pixels, and 300 dpi resolution in digital format (JPEG) compatible with Image J (1.50c4 for Mac OS Sierra 10.12.6).

Craniofacial growth was assessed radiographically using linear and angular distances between predetermined topographic anatomical points in the jaws (Figure 1) (17). Two blinded observers were previously trained for conducting cephalometry studies. The dental age was estimated by the Nolla tooth development stage (18).

The sample size was estimated before data collection based on a review of previous studies (10,13,14). A primary error probability of 5% and statistical power of 80% was assumed. Statistical comparisons were undertaken through the Student t-test using Excel for Mac (Microsoft version 15.37).

This study was approved by the Research Ethics Committee of the Faculdade de Medicina, Universidade de São Paulo (CEP/CONEP number 05139018.9.0000.0068). The parents or legal guardians of the participants provided written consent for their children to participate in the study.

RESULTS

Patients Characteristics and Oncologic Treatment

Fifty panoramic radiographic images were analyzed (25 HSCT in the group and 25 CONTROL in the group). Radiographic images were obtained on an average of 2.43 (± 2.0) years after HSCT.

Age and sex of the control group were matched to the study group. Sixteen male and nine female patients composed each group. The mean age at transplantation was 10 (± 4.7) years. The mean age during radiography was 12.2 (± 3.8) years.

Fifty-six percent of the primary cancer diagnosis was acute lymphoblastic leukemia, 36% was acute myeloid leukemia, and 8% was myelodysplastic syndromes. Individualized primary malignant diagnosis has been described in Supplementary file 1.

Pre-HSCT myeloablative conditioning included different combinations of cyclophosphamide (76%), fludarabine (40%), busulfan (20%), melphalan (20%), and etoposide (8%). Eighty percent of the included patients (n=20) received myeloablative pre-HSCT total body irradiation divided into six sessions of 200 cGy, totaling 1200 cGy for each patient. All patients received methotrexate and corticosteroids.

In 21% of the cases, autogenous HSCT was performed and the remaining were allogeneic. Of these, 36.3% received stem cells from related donors, 36.3% from unrelated donors, and 27.2% from the umbilical cord. The mean age during HSCT was 10 (± 4.7) years.

Height and Weight Assessment

Younger (age < 12 years) male and female HSCT patients were below the stature average and had above-average weight for their age (Z-score height: -0.13; Z-score weight: +1.8). In older patients (age > 12 years), this was seen in the male-only comparison (Z-score height: -1.5; Z-score weight: +0.68). Female patients showed decreased average stature and weight (Z-score height: -1.4; Z-score weight: -0.95). Table 1 shows the absolute height and weight mean values, body mass index (BMI), and SD (Z-score) for each age group.

Cephalometric Assessment

HSCT group showed a smaller vertical growth in the mandibular ramus with significant statistical differences (p=0.003). The comparison of individuals younger than 12 years did not show significant difference (p=0.13). The comparison of individuals above 12 years of age showed a significant reduction in the mandibular ramus vertical growth (p=0.017). Table 2 presents vertical extensions and the mean values of mandibular ramus for both groups.

There were no statistically significant differences in the comparison of the other linear or angular cephalometric assessments. Table 3 presents the mean values for both groups.

Figure 1 - Maxillomandibular cephalometric assessment. A. posterior side of the right jaw; B. anterior nasal spine; C. posterior side of the left maxilla; D. head of the left jaw; E. left mandibular angle; F. mentonian fossa; G. right mandibular angle; H. right jaw head; I. apex of the inferior interincisal alveolar ridge.
Dental age estimation

HSCT patients showed delayed dental formation and eruptive chronology, though there was no statistically significant difference between the groups \( (p=0.3686) \). Figure 2 presents the correlation between chronological and dental age according to the dental maturation stage (18).

**DISCUSSION**

HSCT plays an important role in the management of hematologic high-risk pediatric malignancies with elevated long-term disease-free survival rates (1-3). However, associated late complications include craniofacial impairment and low stature (10,19).

In this study, we found reduced vertical growth in the mandibular ramus of HSCT patients in comparison to healthy controls \( (p=0.003) \). This difference persisted in patients older than 12 years \( (p=0.017) \), but was not seen in the younger group \( (<12\text{ years, } p=0.13) \), suggesting that HSCT plays an active role lowering stature and impairing the craniofacial skeleton. Previous reports indicate a positive correlation between somatic and craniofacial growth (20). In addition,
the observed statistically significant differences that persisted in the > 12 year-old group suggest that the puberal spurt does not lead to the catch-up growth of craniofacial bones (21). HSCT patients showed late dental eruption compared to their matched controls. Dental eruption was previously assumed to be a causative effect of reduced vertical mandibular growth in early age HSCT survivors (16). However, further studies are required to establish a positive correlation between delayed dental eruption and reduced craniofacial vertical growth.

Craniofacial changes are associated with increased risk of sleep apnea, respiratory disorders, and impaired neurodevelopment (22). The interpretation of our results allows us to infer that the vertical deficit of the face observed in our results may be a contributing factor to the development of respiratory diseases during adulthood, and further studies with this population are needed.

The early detection of craniofacial deformities is important to provide childhood HSCT survivors proper care and mitigate the quality of life impairment associated with oncologic treatments. Panoramic radiographs present as a trustworthy and cost-effective tool for craniofacial growth screening early malocclusion diagnosis (23,24).

The post-transplantation period of evaluation (mean 2.43 years, min.: 6 months; max.: 7 years) did not considerably affect the comparative analyses of craniofacial growth for the following reasons. First, patients aged below 12 years undergoing HSCT have a significantly higher risk for dental and jaw aberrations compared with healthy individuals. Second, the age at transplant has a higher impact on dental development and craniofacial growth compared with conditioning regimens.

The main limitation of this study was the impossibility of separately studying the different myeloablative regimens due to the high individualization of chemotherapeutic agents to minimize the side effects. However, this limitation does not invalidate the present results since no differences were observed between the prevalence of dental anomalies and different myeloablative chemo-radiotherapy protocols (14).

The comparative analysis of craniofacial growth and development showed statistically significant differences in the group of patients over 12 years of age. These results suggest that unwanted HSCT effects include late craniofacial skeletal changes. However, more long-term studies are needed to study these variables.

■ CONCLUSIONS

Compared to the control group, pediatric HSCT patients had delayed skeletal changes due to impaired craniofacial growth and development.

Statistically significant differences were observed in the growth of the Madibular Ramus, which was maintained in individuals over 12 years of age.

■ AUTHOR CONTRIBUTIONS

Frascino AV contributed in writing of the article. Collassanti MD contributed in HSCT and oncological data. Fava M contributed in Radiographic and dental supervision. Odone-Filho V contributed in Main supervision, text review.

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## APPENDIX

### Supplementary file 1 - Individual malignant diagnosis.

| ID | SEX  | AGE AT HSCT(Y) | MALIGNANT DIAGNOSIS |
|----|------|----------------|---------------------|
| 1  | female | 2.3611         | AML                 |
| 2  | male  | 3.4473         | AML                 |
| 3  | female | 4.6785         | ALL                 |
| 4  | male  | 6.5116         | ALL                 |
| 5  | male  | 6.7852         | AML                 |
| 6  | female | 7.4145         | MDS                 |
| 7  | male  | 7.6936         | ALL                 |
| 8  | female | 7.6936         | ALL                 |
| 9  | male  | 8.4323         | ALL                 |
| 10 | female | 8.4432         | AML                 |
| 11 | male  | 8.4706         | ALL                 |
| 12 | female | 9.554          | MDS                 |
| 13 | female | 9.554          | ALL                 |
| 14 | male  | 10.202         | ALL                 |
| 15 | male  | 10.492         | ALL                 |
| 16 | male  | 11.083         | ALL                 |
| 17 | male  | 11.644         | ALL                 |
| 18 | male  | 12.252         | AML                 |
| 19 | male  | 13.576         | AML                 |
| 20 | female | 13.953         | AML                 |
| 21 | female | 14.367         | AML                 |
| 22 | male  | 14.547         | ALL                 |
| 23 | male  | 14.843         | AML                 |
| 24 | male  | 15.141         | ALL                 |
| 25 | male  | 15.157         | ALL                 |

ALL, acute lymphoblastic leukemia; AML, acute myeloid leukemia; and MDS, myelodysplastic syndrome.