Late-Onset, Noninfectious Pulmonary Complications following Allogeneic Hematopoietic Stem Cell Transplantation: A Nationwide Cohort Study of Long-Term Survivors

Ole Henrik Myrdal[^a,b] Trond Mogens Aaløkken[^b,c] Phoi Phoi Diep[^b,d,e] Ellen Ruud[^b,d] Lorentz Brinch[^f] Kristian Foss[^b,c] Henrik Mangseth[^a] Johny Kongerud[^a,b] Liv Ingunn Sikkeland[^a,b] May B. Lund[^a,b]

[^a]: Department of Respiratory Medicine, Oslo University Hospital, Rikshospitalet, Oslo, Norway; [^b]: Institute of Clinical Medicine, University of Oslo, Oslo, Norway; [^c]: Department of Radiology, Oslo University Hospital, Oslo, Norway; [^d]: Department of Pediatric Oncology and Haematology, Oslo University Hospital, Rikshospitalet, Oslo, Norway; [^e]: Division of Pediatric and Adolescent Medicine, Department of Pediatric Research, Oslo University Hospital, Oslo, Norway; [^f]: Department of Haematology, Oslo University Hospital, Rikshospitalet, Oslo, Norway

**Keywords**
Allogeneic hematopoietic stem cell transplantation · Bronchiolitis obliterans syndrome · High-resolution CT · Pulmonary function · Long-term follow-up

**Abstract**
**Background:** Survivors of allogeneic hematopoietic stem cell transplantation (allo-HSCT) are at risk for pulmonary adverse events. Data on late-onset noninfectious pulmonary complications in long-term adult survivors of allo-HSCT are limited and incomplete. **Objectives:** This study aimed (1) to determine occurrence and degree of pulmonary sequelae in adult survivors of allo-HSCT and (2) to identify associations between pulmonary function, high-resolution CT (HRCT), and clinical characteristics. **Method:** In a nationwide, single-center cross-sectional study, 103 survivors (aged median [range] 35 [17–58] years, 53% females) were examined 17 (6–32) years after allo-HSCT and compared with healthy controls (n = 105). Methods included pulmonary function tests and HRCT. **Results:** Chronic graft-versus-host disease was diagnosed in 33% of survivors, including 12% with bronchiolitis obliterans syndrome (BOS). Mean lung volumes (TLC, FVC, and FEV1) and gas diffusing capacity were >80% of predicted for the survivors as a group, but significantly lower than in healthy controls. Pathological HRCT findings were detected in 48% of the survivors (71% airways disease, 35% interstitial lung disease, and 24% apical subpleural interstitial thickening). Air trapping (%) on HRCT correlated with % predicted FEV1, p < 0.001. In a multiple logistic regression model, both BOS and pathological findings on HRCT were associated with chemotherapy prior to allo-HSCT, p < 0.05. **Conclusions:** Long-term allo-HSCT survivors had significantly lower pulmonary function than age- and gender-matched healthy controls and nearly half had pathological findings on HRCT. Longitudinal data will determine if pulmonary sequelae will remain stable or progress. We recommend lifelong monitoring of pulmonary function in allo-HSCT survivors. HRCT provides additional information, but is not suited for surveillance.

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Correspondence to:
Ole Henrik Myrdal, omyrda@ous-hf.no
Introduction

Allogeneic hematopoietic stem cell transplantation (allo-HSCT) is an established, life-saving treatment for a diversity of malignant and nonmalignant disorders [1]. It is, however, associated with numerous adverse effects that may complicate long-term outcomes [2].

Pulmonary complications have been reported in one-third of allo-HSCT recipients, affecting morbidity and mortality [3]. Factors associated with pulmonary complications are graft-versus-host disease (GVHD), conditioning regimen, underlying lung disease, and factors related to treatment prior to allo-HSCT [3]. In brief, pulmonary complications may develop due to either infectious or noninfectious causes. Late-onset noninfectious pulmonary complications (LONIPCs) may be caused by chemotheraphy used in the myelosuppressive conditioning regimen [4], total body irradiation [5, 6], or as an immunological response induced by the graft [7]. The latter is typically late-occurring in the form of chronic GVHD (cGVHD) [8]. The most frequent LONIPC is bronchiolitis obliterans, which is linked to cGVHD [9]. Bronchiolitis obliterans is diagnosed by its histological pattern of fibrogenic deposition in small airways. A lung biopsy is needed, which requires an invasive and potentially harmful procedure. Therefore, the clinically based diagnosis bronchiolitis obliterans syndrome (BOS) is usually preferred. Criteria for the diagnosis of cGVHD and BOS have been published by the National Institutes of Health (NIH) [10]. The diagnosis of BOS is primarily based on pulmonary function tests with supporting criteria based on high-resolution computed tomography (HRCT) and differs from the diagnosis of BOS following lung transplantation [11]. Also, interstitial lung diseases (ILDs) may occur in survivors of allo-HSCT and are often labeled LONIPCs as well [9]. The ILDs have different appearances and histological findings [12]. One is pleuroparenchymal fibroelastosis which is characterized by elastic fibrosis, involving lung tissue in the upper lobes [13]. Hence, the development of LONIPCs is complex, with a range of rare conditions from ILDs to BOS, and there may also be an overlap between these disorders in the same patient [3]. To our knowledge, HRCT and pulmonary function tests have been applied jointly in only a few clinical studies with focus on late complications in adult allo-HSCT survivors [14–16]. One was a large (n = 198), prospective study, median 6 years after allo-HSCT, reporting 44 survivors with LONIPCs, mainly in terms of BOS and ILD [14]. Other studies applying HRCT have been restricted to patients with previously diagnosed cGVHD/ BOS [15–17]. Hence, little is known of LONIPCs in adult allo-HSCT survivors with observation periods exceeding 1 decade. The present study was carried out within the context of a large and comprehensive project investigating health conditions in young, very long-term survivors of allo-HSCT [18–21]. All survivors were examined with both HRCT and pulmonary function tests. We aimed to assess the occurrence and degree of late pulmonary sequelae in a nationwide cohort of survivors who had been treated with allo-HSCT in childhood, adolescence, or early adulthood median 17 years previously. We also aimed to identify associations between clinical characteristics, HRCT findings, and pulmonary function.

Materials and Methods

Design and Study Population

The study was conducted at Oslo University Hospital from August 2014 to February 2016, as a part of a single-center, nationwide cross-sectional study covering a broad range of late treatment-related effects after allo-HSCT [18–21]. The survivors were eligible for inclusion if they were younger than 30 years at transplantation, older than 16 years at examination, and had minimum 5 years of observation (n = 157) (Fig. 1). All participating survivors (n = 103) gave their written informed consent. The study was approved by the South-East Regional Committee for Medical and Health Research Ethics (2014/370).
Healthy Controls

For pulmonary function testing, 105 healthy, never-smoking subjects (age [median, range] 35 [20–59] years, 56% females) with no history of cancer or pulmonary disease were recruited among university and hospital employees through local advertisements. The controls did not undergo HRCT examination since it was considered unethical to expose healthy subjects to irradiation.

Clinical Assessment

Clinical data, including gender, age, BMI, smoking habits, medical history, physician-diagnosed lung disease, and current medication, were recorded. Routine blood tests included hemoglobin (Hb) levels.

Pulmonary Function

Measures of pulmonary function were performed according to the American Thoracic Society/European Respiratory Society (ATS/ERS) guidelines [22, 23] and included total lung capacity (TLC), vital capacity (VC), residual volume (RV), forced vital capacity (FVC), forced expiratory volume in 1 s (FEV1), and diffusing capacity of the lung for carbon monoxide (DLCO). DLCO values were also corrected for Hb levels, obtained on the same day as pulmonary function testing. Since Hb correction of DLCO did not significantly influence the results of the analyses, the variable DLCO Hb was not reported in the Results section. The predicted values for spirometry and DLCO were taken from the Global Lung Initiative [24, 25] and static volumes from the European Community for Steel and Coal [26]. All tests were performed with the Jaeger Master Screen Body (Erich Jaeger, Wurzburg, Germany). BOS was diagnosed according to NIH criteria [10]: (1) new-onset airway obstruction with FEV1/FVC <0.70, and (2) FEV1 <75% of predicted, with 10% decline over <2 years, and (3) absence of infection, and (4) one of 2 supportive features of air trapping, either on expiratory HRCT or by residual volume (RV) >120% of predicted. Due to the cross-sectional study design, we cannot tell if airway obstruction was new-onset or not, or if FEV1 had declined >10% in 2 years. Restrictive impairment and impaired gas diffusing capacity were defined as, respectively, TLC and DLCO <80% of predicted. We calculated the lower limit of normal in order to confirm that the cutoff 80% predicted fitted the lower 5th percentile of the reference values [27]. We used the term pulmonary sequelae for describing the combination of impairments in pulmonary function and pathological findings on HRCT.

Acquisition and Review of HRCT Images

CT examinations were performed with a LightSpeed 16 scanner (GE Healthcare, Milwaukee, WI, USA) and obtained in the supine position, during deep inspiration and breath-holding. Tube current settings were adjusted to each patient’s weight, but with low dose references (noise index: 40). No intravenous contrast material was used. In all subjects, supplementary expiratory scans were obtained with 1.25-mm section thickness at 10-mm intervals. Supplementary prone position scans were obtained when considered necessary to differentiate between fine reticular fibrosis and dependent atelectasis. The images were reviewed on a PACS (Picture Archiving and Communication System) screen in random order and in consensus by 2 experienced chest radiologists (T.M.A. and K.F.) and 1 pulmonologist (O.H.M.) with special training in interpretation of HRCTs. The reviewers were blinded to lung function and clinical data. The presence, extent, and distribution of interstitial findings were evaluated according to the HRCT criteria of ILD recommended by the Nomenclature Committee of the Fleischner Society [28]. HRCT-detected ILD was defined as reticular pattern and/or ground glass opacities. Airways disease was defined as bronchiectasis, and/or air trapping, and/or mosaic pattern, and/or centrilobular micronodules. In the setting of long-term survivors of allo-HSCT with the risk of LONIPCs, the HRCT findings were categorized into 3 major groups: airways disease, apical irregular subpleural interstitial thickening, and other signs of ILD. The distribution of pathological findings was evaluated in 4 zones: (a) above the aortic arch, (b) between the aortic arch and the level of the carina, (c) between the level of the carina and the level of the inferior pulmonary veins, and (d) below the inferior pulmonary veins. The extent of involvement of pathological findings was evaluated for each lung zone. The severity of bronchiectasis was scored either as bronchial wall thickening without distinct ectasis or bronchiectasis in localizations (a–d). The extent of air trapping in each zone was assigned a score based on the percentage of lung parenchyma involved, with an overall score...
Late-Onset Pulmonary Complications after Allo-HSCT

of involvement for each patient derived by summing the scores of the 4 CT levels. Subsegmental air trapping comprising <5% of the lung parenchyma was considered normal [27].

Statistical Analysis
Student’s t test or Mann Whitney U test was used, as appropriate, to compare continuous data between groups. χ² test or Fisher’s exact test was used to compare categorical variables. Univariate and multiple logistic regression analyses were used to detect associations between dependent variables and relevant covariates, controlling for extraneous effects. The independent variables which entered the regression models were those hypothesized a priori for biological or clinical reasons or found to be significant at a 20% level by univariate analysis. Pulmonary function in patients with BOS was analyzed separately since the diagnostic criteria for BOS are based on such tests [10]. A 2-sided p value <0.05 was considered significant. Standard statistical analyses were performed with SPSS software (IBM SPSS statistics, version 26).

Results
A total of 157 survivors fulfilled the inclusion criteria and 103 (66%) participated in the study (Fig. 1). The 54 nonparticipating survivors included more males than females (69% vs. 47%, p = 0.01), were (median [range] years) younger at the time of allo-HSCT 14 (0.8–30) versus 20 (0.3–30), p = 0.02, and had shorter observation time 12 (5–24) versus 17 (6–32), p < 0.001 than the included survivors. The nonresponders were comparable to the included survivors with respect to diagnosis prior to transplantation.

Patient Characteristics
Clinical characteristics are outlined in Table 1. Hematological malignancies comprised the underlying diagno-

sis in 77 (75%) survivors of whom 46 (60%) had received chemotherapy prior to allo-HSCT. Patients with chronic myeloid leukemia had not received chemotherapy routinely, in contrast to those with other malignancies. Myeloablative regimens with cyclophosphamide/busulfan or cyclophosphamide had been applied in 101 (98%) survivors. Only 7 subjects had been treated with total body irradiation. Chronic GVHD was diagnosed in 34 (33%) survivors. Survivors with cGVHD were comparable to those without cGVHD with respect to gender, age, observation time, BMI, and smoking habits. Among subjects with benign underlying disorders (n = 26), 4 (15%) had developed cGVHD compared to 30 (39%) among those with malignant disorders (n = 77) (p = 0.03). Chronic GVHD was associated with chemotherapy prior to allo-HSCT (p = 0.01) and history of acute GVHD (p = 0.001), but not with donor match. Mean (SD) Hb was 15.3 (1.0) for males and 13.6 (0.9) for females.

Pulmonary Function
Pulmonary function for the allo-HSCT survivors and the healthy controls is shown in Figure 2. Twelve survivors were diagnosed with BOS. They had increased RV and reduced FVC and FEV₁, hence confirming that they met the NIH criteria. The survivors without BOS had significantly lower TLC, FVC, FEV₁, and DLCO than the healthy controls, whereas RV was comparable. Among the allo-HSCT survivors, 4% had restrictive impairment and 17% had impaired gas diffusing capacity. No case of impairment was observed among the healthy controls. Twelve of the 13 survivors with asthma had been diagnosed after HSCT, and 4 of them had BOS. Ever-smokers

Fig. 2. Pulmonary function in 103 long-term survivors of allo-HSCT and 105 healthy controls. Comparison between controls and allo-HSCT w/o BOS. Data presented as % predicted (mean [95% CI]). Allo-HSCT, allogeneic hematopoietic stem cell transplantation; BOS, bronchiolitis obliterans syndrome.
patients who had received chemotherapy prior to allo-HSCT had lower DLCO % predicted than those who had not received such treatment (mean [95% CI] % predicted 78 [73–82] versus 87 [82–91], \( p = 0.003 \)). Of the survivors with impaired DLCO, 2 had BOS.

**HRCT Findings**

HRCT findings are presented in Tables 2 and 3, Figure 3, and online supplementary Figure 1 (for all online suppl. material, see www.karger.com/doi/10.1159/000520824).

Comparison is between the different findings and normal. One subject may have several findings. Data presented as % of predicted value (95% CI). Allo-HSCT, ALLOGENEIC hematopoietic stem cell transplantation; TLC, total lung capacity in % of predicted value; RV, residual volume; FVC, forced vital capacity; FEV1, forced expiratory volume in 1 s; DLCO, diffusing capacity for carbon monoxide.

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**Table 2. Pulmonary function in 103 long-term survivors of allo-HSCT according to findings on HRCT**

| Types of pathological findings (n = 64) | Survivors w/ normal findings | Survivors w/ pathological findings | \( p \) value |
|----------------------------------------|-----------------------------|-----------------------------------|-------------|
| airways disease                         | (n = 54)                    | (n = 49)                          | 0.13        |
| interstitial lung disease               | (n = 35)                    | (n = 17)                          | 0.02        |
| apical subpleural thickening            | (n = 12)                    |                                   | 0.05        |

| TLC   | RV      | FVC    | FEV1   | FEV1/FVC | DLCO  |
|-------|---------|--------|--------|----------|-------|
| 105 (102–109) | 112 (106–118) | 96 (93–100) | 94 (90–98) | 0.81 (0.78–0.82) | 93 (88–98) |
| 102 (98–106) | 121 (113–130) | 90 (86–94) | 82 (76–88) | 0.75 (0.71–0.78) | 95 (91–100) |

Comparison is between the different findings and normal. One subject may have several findings. Data presented as % of predicted value (95% CI). Allo-HSCT, ALLOGENEIC hematopoietic stem cell transplantation; TLC, total lung capacity in % of predicted value; RV, residual volume; FVC, forced vital capacity; FEV1, forced expiratory volume in 1 s; DLCO, diffusing capacity for carbon monoxide.
Late-Onset Pulmonary Complications after Allo-HSCT

(except one) of the survivors who fulfilled the pulmonary function criteria for BOS had findings consistent with airways disease. The most frequent single finding on HRCT was air trapping, which was detected in 24 survivors, with mean 29% of the lung parenchyma involved, ranging from 5 to 77%. The degree of air trapping in % of lung volume correlated with RV% predicted (r = 0.610, p = 0.002) and inversely with FEV1% predicted (r = −0.771, p < 0.001) (online suppl. Fig. 1c). There was no association between pathological findings on HRCT and cGVHD without BOS (p = 0.926). Age at allo-HSCT was not a risk factor for pneumological abnormalities in our cohort (p = 0.728). In a multiple logistic regression model, controlling for gender, age at allo-HSCT, and observation time, pathological findings on HRCT and BOS were both associated with chemotherapy prior to allo-HSCT (p < 0.041 and p < 0.040) (Table 3).

Discussion

The main findings of this study were that long-term allo-HSCT survivors had significantly lower pulmonary function than age- and gender-matched healthy controls and nearly half of them had pathological findings on HRCT. Although mean lung volumes (TLC, FVC, and FEV1) and gas diffusing capacity were above 80% predicted for the entire study group, one-third of the survivors had some kind of impairment (17% impaired gas diffusing capacity, 12% had developed BOS, and 4% had restrictive impairment). Survivors who had received chemotherapy prior to allo-HSCT had 2.4-fold and 4.5-fold increased risk for, respectively, pathological findings on HRCT and BOS. The predominant pathological findings on HRCT were signs of airways disease, but also various patterns of ILD, including apical irregular subpleural interstitial thickening suggestive of pleuroparenchymal fibroelastosis, were found. The latter is a rare radiological pattern seen, in particular, after allo-HSCT and lung transplantation [13]. Our findings were indicative of pleuroparenchymal fibroelastosis because upper lobe pleural thickening with associated subpleural fibrosis was present and without involvement of lower lobes. However, detailed radiological features of pleuroparenchymal fibroelastosis are still largely unknown, and definite diagnosis requires histological confirmation [29].

The survivors included in the present study were young adults (median 35 years old), and approximately three-quarters were never-smokers. The former/current smokers had few pack-years (median 3 pack-years) which probably explains why their pulmonary function was comparable to the never-smokers. Four of the 12 survivors with physician-diagnosed asthma after HSCT also had BOS. It is possible that respiratory symptoms caused by BOS may have led to misclassification of asthma.

Pulmonary function was well preserved for the survivors who had not developed BOS. However, compared to the healthy controls, also the survivors without BOS had significantly reduced TLC, FVC, FEV1, and gas diffusing capacity. The latter was reduced, in particular, in subjects who had received intravenous high-dose courses of chemotherapy for malignant blood disorders prior to allo-HSCT. This finding is in line with reports from various other studies indicating that DLCO is the most sensitive test for detecting chemotherapy-induced lung injury [30–35]. In previous studies, we have found impaired gas diffusing capacity in long-term lymphoma survivors after high-dose therapy with autologous stem cell transplantation [33] and in very long-term adult survivors of childhood acute lymphoblastic leukemia [34]. Late adverse ef-

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Table 3. Odds ratio for selected risk factors related to, respectively, pathological findings on HRCT and BOS in 103 long-term survivors of allo-HSCT

| Independent variables          | Pathological findings on HRCT | BOS |
|-------------------------------|-------------------------------|-----|
|                               | odds ratio | 95% CI | p value | odds ratio | 95% CI | p value |
| Female gender                 | 1.7         | 0.7–3.9 | 0.205   | 2.3         | 0.6–9.2 | 0.222   |
| Age at allo-HSCT, years       | 1.0         | 1.0–1.1 | 0.502   | 1.0         | 0.9–1.1 | 0.970   |
| Observation time, years       | 1.0         | 0.9–1.1 | 0.565   | 0.9         | 0.8–1.0 | 0.149   |
| Chemotherapy prior to allo-HSCT | 2.4         | 1.0–5.4 | 0.041   | 4.5         | 1.1–18.6 | 0.040   |

Findings on HRCT include signs of airways disease, interstitial lung disease, and apical irregular subpleural interstitial thickening. Allo-HSCT, allogeneic hematopoietic stem cell transplantation; BOS, bronchiolitis obliterans syndrome.
fects of chemotherapy on gas diffusing capacity have also been shown in patients treated for lung cancer [32] and breast cancer [35]. Although the underlying mechanisms of cytotoxic lung injury are multifactorial and remain unclear, it is thought that microvascular damage may be a common denominator and an important feature [30, 31].

Our findings suggest that in combination with pulmonary function tests, HRCT is a useful tool for diagnosing BOS. In the present study, all but one subject with BOS also had pathological findings on HRCT consistent with airways disease. However, signs of airways disease (air trapping, small airway wall thickening, bronchiectasis, and mosaic attenuation) on HRCT are too common and too unspecific to be used alone to diagnose BOS. Air trapping was the single most frequent finding on HRCT in our study and occurred in nearly one-fourth of the survivors of whom only half fulfilled the pulmonary function criteria for BOS. To our knowledge, only one other large-sample study has used both HRCT and pulmonary function tests in order to identify long-term noninfectious lung complications in young adult survivors of allo-HSCT [14]. A few smaller studies have reported HRCT findings in survivors who had already been diagnosed with airway obstruction by previous pulmonary function testing [15–17]. It is, however, difficult to compare HRCT findings from different studies due to various study designs, definitions, and methods of reporting. Air trapping is defined as retention of air distal to airway obstruction, best visualized on expiratory HRCT scans. It is the most common finding on HRCT suggestive of BOS. We chose to report air trapping as a percentage of lung parenchyma, which is a well-established method [16, 28, 36, 37]. We found a strong correlation between % air trapping on HRCT and FEV1% predicted and RV% of predicted. This is in agreement with previous studies that have reported similar associations between airflow obstruction and air trapping in comparable allo-HSCT study populations [15–17]. The prevalence of BOS was 12% in our study, which is comparable to 11% reported by Bergeron et al. [14]. This is interesting, given the difference in follow-up between the 2 studies (median 17 vs. 6 years). Due to the cross-sectional design of our study, we cannot make assumptions regarding longitudinal changes. On the other hand, since our data are comparable with that of Bergeron et al. [14] with a median follow-up time of 6 years, we may assume that most of the cases with BOS do occur within the first few years after allo-HSCT. This is in line with reports from earlier studies [38, 39]. Although we do not know if the occurrence of pulmonary sequelae found in our study will increase or remain unchanged – or even regress – in the future, we think it is unlikely that the findings on HRCT and the impairments in pulmonary function will change in a clinically significant degree since they were – in general – of a mild nature. However, this assumption may be challenged by new biological research focusing the cellular processes of aging and their association with the premature development of age-related diseases seen in cancer survivors [40]. Since long-term comorbidities observed in cancer survivors seem to mimic the phenotypes of aging, they may be caused by some kind of interaction between therapeutic exposures and the underlying biology of aging [40].

One strength of the present study is the single-center, national patient cohort, uniformly treated according to standardized national protocols. Also, at follow-up, all medical tests were undertaken at the same site and carried out by a limited number of highly selected and experienced staff dedicated to the study, and the same type of equipment was used for, respectively, HRCT and pulmonary function testing. Another strength is an age- and gender-matched group of healthy controls. HRCT was applied in all survivors, not only those with diagnosed or suspected pulmonary complications, which has been the choice in other studies [16, 37]. From a research point of view, we obviously would have liked to have HRCT results also for the healthy controls, but since it was considered unethical to expose healthy subjects to irradiation for study purposes, the controls were used for comparing pulmonary function data only. Since we aimed to investigate late pulmonary sequelae in very long-term survivors, the long observation time (median 17 years) may be seen as a strength. However, very long observation time also implies survival bias. In our study, 45% of the allo-HSCT patients were deceased at the time of survey. Since national legislation prevents access to information on individual causes of death, we do not know to what extent pulmonary complications may have affected mortality. In a recent long-term study of mortality including almost 4,500 allo-HSCT survivors, Wong et al. [41] reported that if a patient survives the first 2 years after allo-HSCT, the 5-year survival rates the next 15 years exceed 85%. Furthermore, they found that cGVHD accounted for increasingly fewer deaths among long-term survivors and also a relatively stable mortality rate for pulmonary cause of death throughout the years [41]. Another weakness is the lack of pretreatment data that prevents analysis of longitudinal changes. The patients’ age at diagnosis ranged from 0.3 to 30 years. Although pulmonary function testing had not been routinely carried out prior to allo-HSCT and about one-third of the patients would have been too...
young to obtain reliable tests, it might have been possible to track down baseline pulmonary function data from a subset of the patients. Pretreatment HRCT, on the other hand, was not accessible for any patient since it had not been part of the initial routine before allo-HSCT workup for any of the survivors. Furthermore, the cross-sectional design of the study does not allow us to study causal relationships, just describe associations. In a very long-term follow-up study, we think an attendance rate of 66% is satisfactory. However, we cannot rule out that nonresponse bias may have affected the external validity and generalizability of the results since the nonresponders comprised more males, were younger at the time of treatment, and had shorter observation time. In conclusion, after a median 17 years of observation, allo-HSCT survivors had significantly lower pulmonary function than age- and gender-matched healthy controls and nearly half of them had pathological findings on HRCT. The survivors were young adults (median age 35 years), and longitudinal data are needed to determine if their pulmonary sequelae will remain stable or progress throughout their adulthood. We therefore recommend lifelong monitoring of pulmonary function in allo-HSCT survivors. Pulmonary function testing is cost-effective, readily available, safe, and easy to perform. HRCT may provide additional information and may be indicated for clinical reasons in selected patients. However, HRCT is still a cumbersome and expensive method, and it is also poorly suited for surveillance due to radiation exposure.

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Statement of Ethics

The study was approved by the Regional Committee for Medical and Health Research Ethics (2014/370) and the Data Protection Officer at Oslo University Hospital. Written informed consent was obtained from all participants.

Conflict of Interest Statement

The authors have no conflicts of interest to declare.

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Author Contributions

M.B.L., T.M.A., L.B., and E.R. designed the study. O.H.M., P.P.D., K.F., and H.M. collected the data. O.H.M., P.P.D., and L.I.S. participated in data analysis. O.H.M., L.I.S., T.M.A., J.K., and M.B.L. interpreted the results and drafted the manuscript. The final version was approved by all authors.

Data Availability Statement

All data generated or analyzed during this study are included in this article and its supplementary material files. Further enquiries can be directed to the corresponding author.
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