Review

Long-term complications and side effects after allogeneic hematopoietic stem cell transplantation: an update

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Allogeneic hematopoietic stem cell transplantation (allo-HSCT) is an effective therapy for various malignant and non-malignant diseases. Many patients have now been followed for two or three decades post-transplant and are presumed to be cured. With the tremendous advances achieved in terms of supportive care, it is reasonable to expect outcomes to improve steadily and consequently increasing numbers of transplant survivors will be facing life after the initial transplant experience. Although long-term allo-HSCT survivors generally enjoy good health, for many others, cure or control of the underlying disease is not accompanied by full restoration of health. The burden of long-term morbidity borne by allo-HSCT survivors is substantial, and long-term follow-up of patients who received allo-HSCT is now widely recommended. Immediate survival is no longer the sole concern after allo-HSCT. The goals should also include complete recovery of the overall health status with normal physical and psychological functioning. Long-term side effects after allo-HSCT include non-malignant organ or tissue dysfunction, changes in quality of life, infections related to abnormal immune reconstitution and secondary cancers. Many of these can be attributed to the deleterious effects of chronic graft-versus-host disease. The aims of this review are to provide an update on the recent research evidence in the field. Blood Cancer Journal (2011) 1, e16; doi:10.1038/bcj.2011.14; published online 29 April 2011

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Introduction

Allogeneic hematopoietic stem cell transplantation (allo-HSCT) is an effective therapy for various malignant and non-malignant diseases. Many patients have now been followed for two or three decades post-transplant and are presumed to be cured. At present, the allo-HSCT transplant activity is steadily increasing worldwide. This increase is mainly driven by the use of the so-called reduced intensity conditioning (RIC) regimens, and because of access to alternative stem cell sources from human leukocyte antigen-matched or mismatched unrelated adult donors and from cord blood. A recent analysis, published from the Seattle Group, found a substantial reduction in the hazard of death over the past decade when comparing the period from 1993–1997 to 2003–2007. Improved outcomes appear to be related to reductions in organ damage, infection and severe acute graft-versus-host disease (GVHD). With the tremendous advances achieved in terms of supportive care, it is reasonable to expect outcomes to improve substantially, and consequently increasing numbers of transplant survivors will be facing life after the initial transplant experience. Although long-term allo-HSCT survivors generally enjoy good health, for many others cure or control of the underlying disease is not accompanied by full restoration of health. When looking to all-cause mortality in a cohort of almost 1500 patients who survived 2 or more years after allo-HSCT, mortality rates remain twice as high as that of the general population among 15-year survivors of allo-HSCT, and that the survivors face challenges affecting their health and well-being. Relapse of primary disease (29%) and chronic GVHD (cGVHD; 22%) were the leading causes of premature death. Moreover, in patients who have survived for at least 5 years after allo-HSCT without recurrence of the original disease, life expectancy is still not fully restored. Mortality rates remained four- to ninefold higher than those expected in the general population for at least 30 years after transplantation, providing an estimated 30% lower life expectancy, regardless of current age. In rank order, the leading causes of excess deaths were second malignancies and recurrent disease, followed by infections, cGVHD, respiratory diseases and cardiovascular diseases. The burden of long-term morbidity borne by allo-HSCT survivors is substantial, and long-term follow-up of patients who received allo-HSCT is now widely recommended. Allo-HSCT survivors were twice as likely as siblings to develop a chronic condition, and 3.5 times to develop severe/life-threatening conditions, whereas allo-HSCT survivors with cGVHD were 4.7 times as likely to develop severe/life-threatening conditions. In terms of health-care utilization by long-term survivors after allo-HSCT, more than 95% of the patients reported medical contact and the prevalence of general physical examination increased with time, highlighting the need for increased awareness of long-term side effects after allo-HSCT.

At present, immediate survival is no longer the sole concern after allo-HSCT, as many patients can survive the acute complications of the procedure and remain free of their original disease for several years. As a matter of fact, the primary aim of allo-HSCT is the cure from the primary disease. However, the goals should also include complete recovery of the health status with normal physical and psychological functioning but also normal family and social integration with good subjective well-being. Long-term side effects after allo-HSCT include non-malignant organ or tissue dysfunction, changes in quality of life, infections related to abnormal immune reconstitution and secondary cancers. Many of these can be attributed to the deleterious effects of cGVHD. Almost all organs and tissues can...
be the target for the long-term side effects after allo-HSCT. Nevertheless, one should also take into account the common diseases or other comorbidities that the patient might have had before transplantation. Also, one should consider the issues of physiological aging and the impact of lifestyle and environment. Among the transplant community, it is common to define all events occurring beyond 3 months after allo-HSCT as late complications, and separate them into delayed (from 3 months to 2 years), late (2–10 years) and very late events (>10 years). Figure 1 summarizes the different steps that may be considered as risk factors for long-term side effects after allo-HSCT. These risk factors can be related to the patient and disease features or to the transplant technique itself, including the choice of conditioning regimen, the choice of the graft source and the occurrence of long-term cGVHD. Despite an ever-growing knowledge of the natural history of the side effects of allogeneic bone marrow transplantation, long-term complications and the features of cGVHD symptoms are heterogeneous in nature, time of onset, duration and severity. The latter parameters may depend on the intensity of the conditioning regimen and immunosuppressive therapies used. The aims of this review are not to provide a comprehensive list of the different late complications occurring after allo-HSCT, but rather an update on the recent research evidence in the field. Recommended screening and preventive practices were published elsewhere and will not be discussed here. Also, specific complications encountered in the pediatric allo-HSCT population, will not be addressed.

Role of the conditioning regimen

Intensity of the conditioning regimen, especially the use of full-dose myeloablative total body irradiation (TBI) is likely to have a major role in the onset of diverse late complications after allo-HSCT. Several studies have demonstrated an impact of TBI on cataract formation with an incidence of <20% in the no-TBI group versus more than 80% in the single-dose TBI group. Spermatogenesis provides another example of the deleterious impact of high-dose TBI, with 85% of the patients receiving TBI developing azoospermia versus 51% in the no-TBI group. The incidence of diabetes and hypertension is also significantly increased after allo-HSCT as compared with patient receiving autologous transplantation or to their siblings. Besides, the use of radiation therapy appears to be deleterious especially when it comes to diabetes. As non-myeloablative or RIC regimens are being increasingly used in the allogeneic setting, one might question whether the use of RIC will reduce long-term toxicity after allo-HSCT? At present, it is well established that RIC can reduce short-term acute toxicity. However, it has been proven that the probability of cGVHD is not reduced after RIC because RIC regimens are mainly used in older patients, in more advanced diseases with peripheral blood stem cells being the main graft source. In one study assessing chronic kidney disease in patients that had received RIC versus myeloablative regimens, there were no significant differences in term of renal complications.

Role of the stem cell source and chronic GVHD

The graft source has a major role in long-term side effects, especially cGVHD. A large meta-analysis that included more than 1100 patients from nine international randomized trials, demonstrated a significantly higher incidence of extensive cGVHD when using granulocyte colony-stimulating factor-mobilized peripheral blood stem cells versus bone marrow. On the other hand, the incidence and severity of cGVHD is likely lower after unrelated cord blood transplantation. Chronic GVHD, and its different target organs (Table 1), is one of the most serious consequence of allo-HSCT, involving many physical, human, emotional, societal, financial and intellectual resources for the patients and caregivers. Indeed, active cGVHD has a significant impact on many aspects of the overall health status. In multivariable analyses, subjects with active cGVHD were more likely to report adverse general health, mental health, functional impairments, activity limitation and pain than were those with no history of cGVHD. However, health status did not differ between those with resolved cGVHD and those who never had cGVHD. Also, the percentage of chronic GVHD patients with poor general health status will remain very high at any time after transplantation. Avascular necrosis is one example demonstrating that cGVHD and its corollary of long-term immunosuppressive therapy are significant risk factors for this complication. The lung is another important target for cGVHD with different complications being described thus far. The non-infectious complications can be divided into the bronchiolitis obliterans syndrome, the bronchiolitis obliterans-organizing pneumonia and the idiopathic pneumonia syndrome. The incidence of pulmonary non-infectious complications in patients surviving more than 3 months after allo-HSCT transplant is around 10% at 2 years and the outcome of patients with cGVHD who develop pulmonary complications is significantly worse than cGVHD patients without pulmonary complications. Bronchiolitis obliterans syndrome is usually considered as a severe pulmonary manifestation affecting the small airways. Its incidence can range from 2 to 14% and the overall survival is around 13% at 5 years. The clinical presentation is often insidious, although some patients may have dry cough, progressive dyspnea or wheezing, without...
The diagnosis of bronchiolitis obliterans syndrome is based on an expiratory flow <75% of predicted in addition to specific features on high-resolution computed tomography scan and absence of infection in the respiratory tract. An increasing body of data could show a strong association between bronchiolitis obliterans syndrome and pulmonary cGVHD.

### Table 1: Signs and symptoms of chronic GVHD (reproduced from Filipovich et al.\textsuperscript{24})

| Organ or site | Diagnostic (sufficient to establish the diagnosis of chronic GVHD) | Distinctive (seen in chronic GVHD, but insufficient alone to establish a diagnosis of chronic GVHD) | Other features\textsuperscript{a} | Common (seen with both acute and chronic GVHD) |
|--------------|-----------------------------------------------------------------|-----------------------------------------------------------------|-------------------------------|-----------------------------------------------|
| Skin         | Poikiloderma                                                   | Depigmentation                                                   | Sweat impairment              | Erythema                                      |
|              | Lichen planus-like features                                     |                                                                 | Icthyosis                     | Maculopapular rash                            |
|              | Sclerotic features                                             |                                                                 | Keratosis pilaris             | Periungual erythema                           |
|              | Morphea-like features                                          |                                                                 | Hypopigmentation              | Pruritus                                      |
|              | Lichen scleros-like features                                   |                                                                 | Hyperpigmentation             |                                               |
| Nails        | Dystrophy                                                      | Longitudinal ridging, splitting, or brittle features             |                               |                                               |
|              | Onycholysis                                                    | Pterygium unguius                                                |                               |                                               |
|              | Nail loss (usually symmetric; affects most nails)\textsuperscript{b} | Nail loss (usually symmetric; affects most nails)\textsuperscript{b} |                               |                                               |
| Scalp and body hair | New onset of scarring or nonscarring scalp alopecia (after recovery from chemoradiotherapy) | Scaling, papulosquamous lesions Xerostomia | Thinning scalp hair, typically patchy, coarse, or dull (not explained by endocrine or other causes) Premature gray hair | Gingivitis Mucositis Pain |
| Mouth        | Lichen-type features                                           | Mucocoele                                                        |                               |                                               |
|              | Hyperkeratotic plaques                                          | Mucosal atrophy                                                  |                               |                                               |
|              | Restriction of mouth opening from sclerosis                     | Pseudolamellibranches\textsuperscript{a}                        |                               |                                               |
|              |                                                               | Ulcers\textsuperscript{a}                                        |                               |                                               |
| Eyes          | New-onset dry, gritty, or painful eyes\textsuperscript{c}       | Cricatrical conjunctivitis                                       | Photophobia                   |                                               |
|              |                                                               | Keratoconjunctivitis sicca\textsuperscript{c}                    | Hyperpigmentation             |                                               |
|              |                                                               | Confluent areas of punctuate keratopathy                         | Blepharitis (erythema of the eyelids with edema) |                                               |
| Genitalia    | Lichen planus-like features                                     | Erosions\textsuperscript{b}                                      |                               |                                               |
|              | Vaginal scarring or stenosis                                    | Fissures\textsuperscript{b}                                      |                               |                                               |
|              |                                                               | Ulcers\textsuperscript{b}                                        |                               |                                               |
| GI tract      | Esophageal web                                                  | Exocrine pancreatic insufficiency                                 |                               |                                               |
|              | Strictures or stenosis in the upper to mid third of the esophagus\textsuperscript{b} |                               |                               |                                               |
| Liver         |                                                                |                                                                |                               |                                               |
| Lung          | Bronchiolitis obliterans diagnosed with lung biopsy             | Bronchiolitis obliterans diagnosed with PFTs and radiology\textsuperscript{a} |                               |                                               |
| Muscles, fascia, joints | Joint stiffness or contractures secondary to sclerosis |                                                                | Edema                         |                                               |
| Hematopoietic and immune |                                                                |                                                                | Muscle cramps                 |                                               |
|              |                                                                |                                                                | Arthralgia or arthritis       |                                               |
|              |                                                                |                                                                | Thrombocytopenia              |                                               |
|              |                                                                |                                                                | Eosinophilia                  |                                               |
|              |                                                                |                                                                | Lymphopenia                   |                                               |
|              |                                                                |                                                                | Hypo or                       |                                               |
|              |                                                                |                                                                | hypergammaglobulinemia        |                                               |
|              |                                                                |                                                                | Autoantibodies (AIHA and ITP) |                                               |
|              |                                                                |                                                                | Pericardial or pleural effusions |                                               |
|              |                                                                |                                                                | Ascakes                        |                                               |
|              |                                                                |                                                                | Peripheral neuropathy         |                                               |
|              |                                                                |                                                                | Nephrotic syndrome            |                                               |
|              |                                                                |                                                                | Myasthenia gravis             |                                               |
|              |                                                                |                                                                | Cardiac conduction            |                                               |
|              |                                                                |                                                                | abnormality or cardiomyopathy |                                               |

Abbreviations: AIHA, autoimmune hemolytic anemia; ALT, alanine aminotransferase; AST, aspartate aminotransferase; BOOP, bronchiolitis obliterans-organizing pneumonia; GVHD, graft-versus-host disease; ITP, idiopathic thrombocytopenic purpura; PFTs, pulmonary function tests.\textsuperscript{a}Can be acknowledged as part of the chronic GVHD symptomatology if the diagnosis is confirmed.\textsuperscript{b}In all cases, infection, drug effects, malignancy, or other causes must be excluded.\textsuperscript{c}Diagnosis of chronic GVHD requires biopsy or radiology confirmation (or Schirmer test for eyes).
In contrast to pulmonary complications, cardiac and vascular complications usually occur ‘very late’ after allo-HSCT with a relatively low magnitude of risk during the first decade. Indeed, after 20 years of follow-up, the incidence of arterial events might be as high as 22%25,26. In addition to the allo-HSCT treatment itself, the usual cardiovascular risk factors such as smoking, hypertension, diabetes or obesity were shown to be significant factors for an arterial event after allo-HSCT.25 From the pathophysiological standpoint, data published in 2002 suggested that host endothelial cells are a target of alloreactive donor cytotoxic T cells, and there is an extensive loss of microvessels in the skin of patients with cGVHD, which may lead to tissue fibrosis, the hallmark lesion of cGVHD.27 Renal complications are also frequent after allo-HSCT with an incidence around 23% after a median time of 2.5 years after transplantation. Older age, hypertension, poor kidney function before transplantation, but also acute and cGVHD with the use of calcineurin inhibitors were shown to be risk factors for chronic kidney disease. The effect of TBI is still controversial.28-30

**Late secondary malignancies after allo-HSCT**

Different categories of secondary malignancies can occur after allo-HSCT, including post-transplant lymphoproliferative disorders, late relapse of the primary disease, donor-type secondary leukemia/other malignancy and de novo solid tumors. As part of this Review, we will focus on secondary solid tumors. TBI and the chemotherapeutic drugs used before allo-HSCT as part of the conditioning regimen can induce new secondary malignancies after allo-HSCT. This is attributed to the mutagenic risk of irradiation and chemotherapy, the genetic predisposition of the patient to develop cancer, prolonged immunosuppression, and in elderly patients, to age-related risk.

One of the first largest reports, including more than 1000 patients transplanted before December 1985, could show that the incidence of secondary malignancies is around 3.5% at 10 years and 12.8% at 15 years, which is 3.8-fold higher than that of an age-matched control population.31 This study was recently updated. This update showed that the incidence of secondary malignancies continued to increase with longer follow-up time compared with age-matched control population.32 A more recent very large analysis including 28,874 patients who received allo-HSCT, showed that solid cancers occurred twice more frequently than expected in the general population and this risk reached threefold among patients followed for 15 years or more. Of note, allo-HSCT survivors irradiated at young ages had increased risks of solid cancers.33 Indeed, the risk of secondary solid cancer may vary depending on the type of conditioning regimen. In a study including acute and chronic myeloid leukemia patients receiving high-dose Busulfan and cyclophosphamide conditioning, the cumulative incidence of solid cancers at 10 years was 1.2% for acute myeloid leukemia and 2.4% for chronic myeloid leukemia, which is 1.4 times higher in comparison with general population incidence rates. Significantly elevated risks were observed for tumors of the oral cavity, esophagus, lung, soft tissue and brain. Chronic GVHD was an independent risk factor for all solid cancers, and especially for cancers of the oral cavity.34 The picture may also be different when it comes to some specific cancers such as thyroid cancer in which younger age was found to be the strongest risk factor of secondary thyroid cancer.35 A joint EBMT/Seattle study showed that the cumulative incidence of breast cancer is around 5% at 20 years and the median time to breast cancer diagnosis is around 12.5 years. Cumulative incidence was higher among survivors who received TBI than those who did not receive TBI. In multivariable analysis, increased risk was associated with longer time since transplantation, use of TBI and younger age at transplantation.36

**Conclusion**

Allo-HSCT has the potential to cure a significant proportion of patients with otherwise fatal diseases. However, allo-HSCT recipients are exposed to long-term complications and side effects. All of these late complications present a great diversity in respect to frequencies, time of onset, risk factors, prevention strategies, treatment approaches and outcomes. In many patients, long-term survivorship is not accompanied by full restoration of health. The long-term side effects are usually multifactorial and complex process, and the natural history of such late effects will likely change in the future because the transplant techniques have been changing significantly over the last three decades. Chronic GVHD remains the most challenging risk factor. A preventable death with a functioning hematopoietic allograft is not a marker of success. Patients can have a major role through engagement in health-maintenance behaviors.

The long-term goal is maintenance of health and to ensure the best possible quality of life. Establishment of a close partnership between the transplant centre, the organ-specific specialties and local primary care providers to provide lifelong counseling and prevention therapy is a key factor for the long-term success of any transplant program.

**Conflict of interest**

The authors declare no conflict of interest.

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