

Clinical and immunological outcome after paediatric stem cell transplantation in inborn errors of immunity Lum, S.H.

Citation

Lum, S. H. (2021, January 20). Clinical and immunological outcome after paediatric stem cell transplantation in inborn errors of immunity. Retrieved from https://hdl.handle.net/1887/139163

Version: Publisher's Version

License: License agreement concerning inclusion of doctoral thesis in the

Institutional Repository of the University of Leiden

Downloaded from: https://hdl.handle.net/1887/139163

Note: To cite this publication please use the final published version (if applicable).

Cover Page



Universiteit Leiden

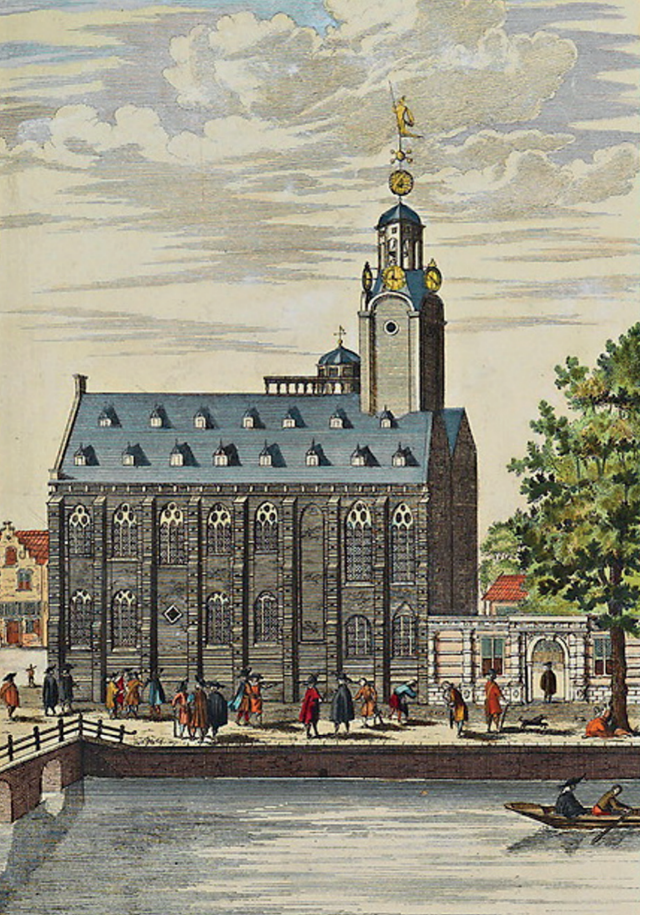


The handle http://hdl.handle.net/1887/139163 holds various files of this Leiden University dissertation.

Author: Lum, S.H.

Title: Clinical and immunological outcome after paediatric stem cell transplantation in

inborn errors of immunity **Issue date**: 2021-01-20



Chapter 9

Malignancy post-haematopoietic stem cell transplant in patients with primary immunodeficiency

Su Han Lum Mary Slatter

Abstract

Introduction: Haematopoietic cell transplantation (HCT) is a curative treatment for an expanding number of primary immunodeficiencies (PIDs). Malignancies are more common in patients with PID than in the general population and this review will discuss whether or not a successful HCT is expected to abolish or alter this risk. Second malignancy post HCT for a malignant disease is well known to occur, but generally less expected in patients transplanted for PID.

Areas covered: This article reviews recently published literature focusing on the pattern of malignancy in children with PID, incidence and risk factors for developing malignancy post-HCT for PID and possible strategies to reduce the risks.

Expert opinion: Survival post HCT for PID has improved dramatically in the last 20 years and the genomic revolution has led to an expanding number of indications. To improve long-term quality of life attention needs to focus on late effects, including the possibility of malignancy occurring more frequently than expected in the general population, understand the risks and improve the process of transplantation in order to minimise them. Further studies are needed.

Introduction

Primary immunodeficiency (PID) or inborn errors of immunity (IEI) comprise a large, heterogeneous group of disorders, often due to single-gene mutations that result in defects in the development and function of the immune system. After the description of Bruton's agammaglobulinaemia by Colonel Odgen Bruton in 1952, more than 350 gene mutations associated with immunodeficiency disorders have been identified. (1-3) Recent studies have shown that PID is increasingly recognised with a prevalence of up to one in 1200 people worldwide potentially living with a PID (4-8). These disorders have diverse phenotypes including infection, malignancy, allergy, auto-immunity and autoinflammation. Although patients with PID commonly present with infection and immune dysregulation, malignancy occurs with a higher incidence and manifests earlier in life, than in the general population, with an overall relative risk varying from 1.4 to 5-fold in registry-based studies. (9-11) It can be the first presentation of PID in some patients. Lymphoproliferative disorders (LPD) represent 50% of all reported cancer in patients with PID. After infections, malignancy is the second most common cause of death in patients with PID. (3, 12)

Advances in haematopoietic cell transplantation (HCT) and supportive care have resulted in significant improvements in survival for children with PID. Following the first successful HCTs for a child with severe combined immunodeficiency (SCID) and another for Wiskott-Aldrich syndrome (WAS) in 1968, transplant survival and graft outcomes have significantly improved in the modern era due to many factors: superior HLA matching technology, greater availability of alternative donors, graft engineering, additional cellular therapy, reduced toxicity conditioning regimens, pharmacokinetic-guided conditioning regimens, vigilant infection surveillance with more effective antimicrobial therapy and better supportive care and treatment for complications. These new developments enable precise personalised transplant care including patient-tailored conditioning regimens and precise prescription of graft components. (13-16) Over the last several decades, the number of HCTs performed for PID has increased and more than 2500 children have received HCTs for PID in Europe and North America. (17, 18) Transplant survival has improved to 90% for SCID with a matched family donor and 70% with a matched unrelated donor transplant. (19, 20) Similar transplant survival has been reported in non-SCID PID recently. (21, 22) Younger age at transplant has been consistently shown to be a significant positive predictor of transplant outcome. (21, 23). In addition, precise molecular diagnosis and understanding of the natural history of PID enable early identification of suitable candidates for HCT. In many patients with non-SCID PID, HCT has developed from being a last resort into the standard of care that corrects the defects in immunity.

Secondary malignancies are a well-known complication of conventional chemotherapy and radiation treatment for patients with various primary cancers and are also well-recognised as a complication among transplant survivors. Malignancy post-HCT (MPT) for haematological disorders has been reported since 1970 and several large studies have shown adult and paediatric transplant survivors are at high risk of MPT with reported incidences of up to 10-15% by 15 years post-HCT. More recent studies using competing risk analysis have shown a lower magnitude of risk of 3-7%. MPT has been reported in both autologous and allogeneic HCT recipients in children. (24-27). MPT has been described after HCT for aplastic anaemia, haemoglobinopathy and inborn errors of metabolism. For MPT for aplastic anaemia, the largest experience was reported by EBMT which included 748 patients who underwent HCT and 860 patients who were treated with immunosuppressive therapy (IST). Patients who developed haematological malignancies within 6 months or solid tumours within 12 months were excluded as these cancers might be due to the underlying diagnosis rather than therapy. The 10-year CIN of second cancer was 3.1% with HCT and 18.8% with IST. The risk of solid cancers was the same in the two groups but the pattern of tumours was different, primarily solid cancers after HCT and haematological malignancies after IST. The major risk factors for MPT were increasing age at HCT and the use of radiation therapy in the conditioning regimen. Deeg HJ et al. reported the risk of MPT for 700 patients who underwent HCT for aplastic anaemia or Fanconi anaemia. The risk of MPT at 20 years post-HCT was 14% for the entire cohort, rising to 42% in 79 patients with Fanconi anaemia. There were 18 solid MPT (all squamous cell carcinoma except one) and 5 haematological MPT. Radiation and use of azathioprine for chronic GvHD were the risk factors of MPT.(28) Although MPT has been described in haemoglobinopathies and inborn errors of metabolism, it is unclear if the risk is increased compared to the general population.

There is also an increased risk of a wide range of cancers associated with solid organ transplantation. The most extensive data come from a cohort study that analyzed the frequency of malignancy in over 175,000 solid organ transplant recipients between 1987 and 2008. (29)The most common organs transplanted included kidney, liver, heart, and lung (in 58, 22, 10, and 4 percent of cases, respectively). Malignancy was identified in over 10,656 cases, which correlated with a standardized incidence ratio (SIR) of 2.10 (95% CI 2.06-2.14) compared with the general population and an excess absolute risk (EAR) of 719 cases per 100,000 person-years. Malignancy occurred in more than 30 different primary sites, but those with a fivefold or greater increase, compared with the general population, included Kaposi sarcoma, skin, Non-Hodgkin lymphoma, liver, anus, vulva and lip.

Whilst allogeneic HCT corrects the underlying immune defect preventing recurrent lifethreatening infection, achieving immune competence post-HCT has been hypothesized to reduce the risk of malignancy in patients with PID. However, HCT may not eradicate the risk of malignancy completely. As transplant survival has improved, the number and length of follow-up of survivors after successful HCT is increasing. With these increases, it is now known that malignancy post-HCT may occur, causing significant morbidity including late and premature death in transplant survivors. This review will focus on the pattern of malignancy in children with PID, incidence and spectrum of malignancy in post-HCT for PID, proposed mechanism and risk factors for carcinogenesis in post-HCT for PID, proposed strategy to reduce the risk of MPT, how to identify patients at risk and how to manage these patients.

Pattern of Malignancy in Primary Immunodeficiency

It has been estimated that the overall risk of developing cancer in patients with PID ranges from 4 to 25% (table 1). (12, 30) In 2018, the United States Immune Deficiency Network (USIDNET) published the largest study which included 3658 PID patients who enrolled in the registry between 2003 and 2015. This registry study observed a 1.42-fold excess relative risk of cancer in patients with PID compared to the age-adjusted population. The greatest increase in cancer incidence was observed in lymphomas in both males (10-fold excess relative risk) and females (8-fold excess related risk). This excessive lymphoma risk was seen in patients with common variable immunodeficiency (CVID) which was the most common PID in the registry. Overall, males with PID had a 1.91-fold excess relative risk of cancer while females with PID had similar cancer rates compared to their age-adjusted population. There was no significant difference in the incidence of common cancers (lung, colon, breast and prostate) in patients with PID compared to the general population. (3) The Australian Society of Clinical Immunology and Allergy registry data which involved 1132 subjects showed that the SIR was significantly elevated for all cancer (SIR 1.6), cancer of the thymus gland (SIR 67.3), non-Hodgkin lymphoma (SIR 8.82), stomach cancer (SIR 6.10) and leukaemia (SIR 5.36) (31) The Netherlands reported 10% of cancer in 745 PID patients in the Dutch national registry between 2009 and 2012. Compared to the general Dutch population, PID patients had a 2.3-fold excess risk of developing cancer, and more than 10-fold increased risk for some solid tumours (thymus, endocrine organs) and haematological malignancies (leukaemia and lymphoma). (10)

Disorders which are associated with a higher incidence of cancer are CVID and ataxia telangiectasia which account for 30% and 24% of reported cancer among PID patients respectively. Approximately 30% of cases are reported in association with Wiskott Aldrich syndrome, severe combined immunodeficiency and selective IgG deficiency. (32, 33) The Immunodeficiency Cancer Registry database on immunodeficiency-associated cancer at the

University of Minnesota showed that nearly 60% of all reported malignancies in PID were lymphomas, 85% of which were non-Hodgkin lymphoma (34). An USIDNET study also showed that lymphomas were the most common cancer and accounted for 48% of all cancers in PID patients, followed by skin cancer (15%) and gastrointestinal (8%) and genitourinary cancers (8%). (3) Patients with PID develop lymphomas at younger ages compared to the general population. Cancers of all types in PID patients are likely to be disseminated at the time of diagnosis. Non-Hodgkin lymphomas in these patients are likely to be of B cell origin, be associated with EBV infection, of high histological grades and involve extranodal tissues, particularly the gut and central nervous system.

Although patients with PID are well-known to be at significant risk of developing malignancy, our understanding about the exact mechanisms of carcinogenesis in PID remains incomplete. One of the functions of the immune system is to recognise and destroy cancer cells without causing toxicity to normal tissues and to prevent cancer recurrence by long-term memory. Similar to the aetiology of cancer in childhood, carcinogenesis in children with PID is multifactorial (figure 1). The proposed mechanisms of increased risk of malignancy in PID encompass reduced immune surveillance, dysregulation of haematopoiesis and impaired DNA damage responses. (35) These can be classified into intrinsic factors and extrinsic factors. Intrinsic factors are mainly associated with haematological malignancies and encompasses impaired genetic stability (e.g. defective DNA repair such as ataxia telangiectasia, Nijmegen Breakage syndrome), genetic predisposition (e.g. leukocyte development defects such as severe congenital neutropenia and bone marrow failure syndrome, defective tumour suppression genes such as DOCK8 deficiency, CVID and autoimmune lymphoproliferative syndrome), and impaired immune function (e.g. X-linked lymphoproliferative disease, ITK deficiency, CD40 ligand deficiency). Extrinsic factors are commonly associated with carcinomas and include impaired clearance of oncogenic viruses (e.g. Epstein-Barr virus in XMEN disease, human papillomavirus in WHIM syndrome), chronic tissue inflammation and iatrogenic (e.g. radiation). Interplay between these factors and impaired immunosurveillance render PID patients at higher risk of developing cancers. (36, 37)

Incidence and spectrum of malignancy in transplant survivors of primary immunodeficiency

Whilst there are a number of single centre and multi-centre studies which reported MPT in children with haematological disorders (24, 38-46), there have been limited studies on MPT in children with PID (Table 2 and 3). The reported incidences of MPT in PID transplant survivors are 2.3% by Kamani *et al.*, 1.5% by Nelson *et al.* (non-malignant HCT, including PID) and 1.3%

of non-PTLD MPT by Unni *et al.* In Nelson's report, there were 6 malignancies in 318 patients who underwent allogeneic HCT for non-malignant disorders. The cumulative incidence of malignancy was $0.3 \pm 0.3\%$ and $2.3 \pm 1.2\%$ at 5 years and 10 years post-HCT respectively. This study showed that 15 times more malignancies occurred after HCT in the study population than expected in the age- and gender-matched Australian general population. In this cohort, one hundred and thirty patients had PID: a patient with SCID and another with CGD developed solid tumours but there was no further information documented. (47) Unni *et al.* carried out a retrospective study of 944 patients with PID who underwent HSCT in 2 specialised centres in the UK. Twelve patients (1.27%) developed non-PTLD malignancy at a median of 3.75 years (range 3 months to 11.2 years) post-HCT. (48) No patients received radiotherapy but all received chemotherapy with at least 1 alkylating agent for conditioning prior to HCT.

Spectrum of malignancy post haematopoietic cell transplantation

MPT is conventionally classified into three distinct groups: 1) lymphoma, including post-transplant lymphoproliferative disease (PTLD) 2) myelodysplasia (MDS) and acute myeloid leukaemia (AML) 3) solid tumours. The time course for developing MPT is variable. Leukaemia and lymphomas develop relatively early post-HCT whereas solid tumours have a longer latency period and are being increasingly reported because of improved transplant survival and longer follow-up. In 2003, Baker *et al.* reported multiple MPT in 8 (5.6%) of 137 patients who developed MPT after allogeneic HCT for various indications in Minnesota, including PID. All were solid MPT expect two who had AML/MDS. The first MPT was diagnosed at a median of 3.9 years post-HCT and the second MPT was diagnosed at a median of 6.3 years post-HCT. Three of these patients had recurrence of skin cancer and the remaining 5 patients had developed two distinct new cancers. (49) In 2019, Baker *et al.* published another large series from Seattle which showed that cumulative incidence of MPS by 30 years after HCT was 22%(50)

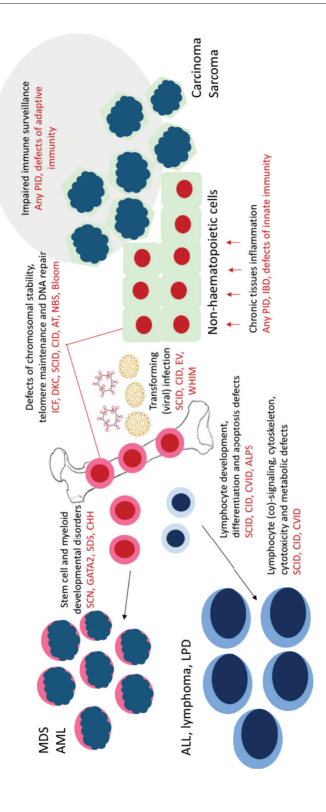


Figure 1: Proposed mechanism of carcinogenesis in patients with primary immunodeficiency

Post-transplant lymphoproliferative disease

PTLD is the most common MPT in the first year after HCT and solid organ transplantation. Most of these cases are related to impaired immune function during early post-HCT and in most cases, proliferation of EBV infection. CIBMTR and the Fred Hutchinson Cancer Research Centre reviewed the largest experience of PTLD in adult and paediatric cohorts which included 26901 allogeneic HCT survivors. This analysis excluded PID (n=532, 20 PTLD, 8.6%) and Fanconi aneamia (n=328, 1 PTLD, 0.3%). In this study, 127 PTLD were identified and the observed to expected ratio was 29.7% (95% CI, 24.7 to 35.2) compared to age-, sexand country-adjusted population. The risk factors for PTLD were T-cell depletion (RR 3.1-9.4), use of ATG (RR 3.8), HLA mismatched in the presence of T-cell depletion/ATG (RR 3.8), acute (RR 1.7) and chronic (RR 2.0) GvHD. Lower risks were found for T-cell depletion methods that remove both T and B cells. This analysis demonstrated a multiplicative effect of multiple risk factors on incidence; the incidence was low (0.2%) in 21686 patients with no major risk factors, but increased to 1.1%, 3.6% and 8.1% with 1, 2 or ≥3 major risk factors. (45)Kamani et al. reported 52 post HSCT malignancies in a large study of 2266 patients with PID giving an overall incidence of 2.3%. The most frequent malignancy was early-onset PTLD in 45 cases, which was associated with T cell depleted grafts. (51, 52)

Therapy-related acute myeloid leukaemia and myelodysplasia

In Nelson's report, three patients developed myelodysplastic syndrome (MDS) all of whom had received TBI and TCD grafts, 1 patient developed acute myeloid leukaemia (AML) and 3 developed solid tumours: a patient with Omenn's syndrome developed a desmoplastic squamous cell carcinoma 170 months after unrelated donor HSCT, a patient with combined immunodeficiency (CID) developed hepatocellular carcinoma 84 months after unrelated donor HCT and another patient developed a brain tumour after HCT for SCID. The incidence of non-PTLD malignancy was 0.3%.(47)

Solid malignancy post-transplant

A wide spectrum of solid MPT has been described and the incidence of solid MPT has been reported to rise over time and successive studies with follow-up to 20 years not showing a plateau in their occurrence (27, 52, 53). Overall, the reported cumulative incidence of solid MPT ranges from 1.2 to 1.6% at 5 years, from 2.2 to 6.1% at 10 years and from 3.8 to 14.9% at 15 years post-HCT. In the CIBMTR report, HCT survivors developed new solid malignancies at twice the expected rate compared to the general population (observed-to-expected ratio 2.1; 95% CI 1.8-2.5). The cumulative incidence of new solid MPT was 1% at 10 years, 2.2% at 15 years and 3.3% at 20 years post-HCT. (27) Similar results were reported by Kolb *et*

al. which included 1036 patients who were transplanted for malignancy, severe aplastic anaemia, PID and inborn errors of metabolism. Socie *et al* reported that solid MPT risk after HCT for childhood cancer increased over time to an estimated 11% at 15 years post-HCT and that age at HCT less than 10 years of age and high dose TBI are associated with higher risk of solid MPT. (24) Unni *et al.* carried out a retrospective study of 944 patients with PID who underwent HSCT in 2 specialised centres in the UK. Twelve patients (1.27%) developed non-PTLD malignancy at a median of 3.75 years (range 3 months to 11.2 years) post-HCT. (48) Overall, the reported cumulative incidence of solid MPT ranges from 1.2 to 1.6% at 5 years, from 2.2 to 6.1% at 10 years and from 3.8 to 14.9% at 15 years post-HCT.

All types of solid cancer have been described in HCT survivors and the types of cancer reported more frequently in HCT survivors include, melanoma, cancers of the oral cavity and salivary glands, brain, thyroid, uterine cervix, breast, bone and connective tissue. Squamous cell carcinoma (SCC) of the oral cavity is more common in HCT survivors, particularly among those with chronic GvHD, Fanconi anaemia, and prior chronic lichenoid lesions of the oral mucosa.

Proposed mechanism and risk factors of carcinogenesis in PID transplant survivors

The risk of developing MPT varies greatly among studies and is dependent on patient-, disease- and transplant-specific factors, including primary disease, age at HCT, chemotherapy agents, radiation dose and field, length and severity of immunodeficiency, graft-versus-host disease, infection and lifestyle (figure 2). PID transplant survivors represent a unique cohort compared to transplant survivors of malignancy. PID patients have an inherent risk of malignancy. Most PID patients do not receive pre-transplant chemotherapy and radiation is rarely used as conditioning regimen. Increased susceptibility to infection and extent of donor chimerism may play a role in patients with PIDs developing MPT.

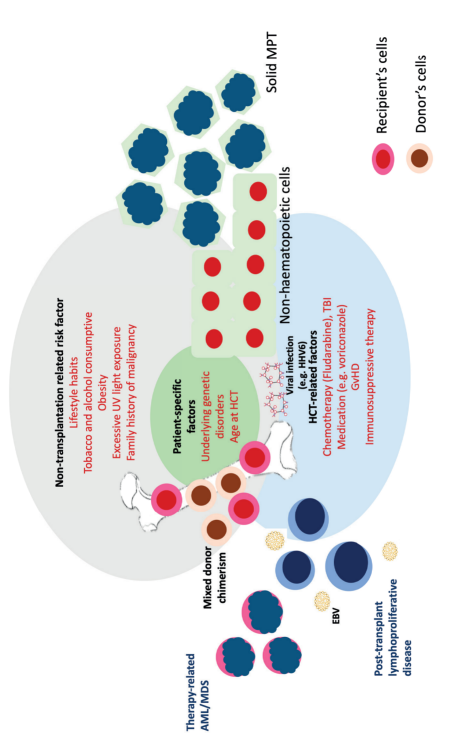


Figure 2: Proposed mechanism of carcinogenesis after HCT in patients with primary immunodeficiency

Patient-specific risk factors: Primary disease and age at transplant

Laffort et al. first described 9 patients with IL-2Ry- and JAK3-deficient SCID with extensive cutaneous human papillomavirus-associated warts. (54) The median onset was 8 years post-HCT (range 3-15 years). Subsequently Kamili et al. reported a further 6 affected patients. (55) The warts are resistant to conventional treatments and may pose a risk of malignant transformation. (56) The occurrence of these warts is not clearly associated with degree of donor chimerism or the level of T-lymphocyte reconstitution post-HCT, but may be associated with low numbers of NK cells or possibly poor NK cell function. Abd Hamid et al. reported 7 of 31 long-term survivors with IL-2Ry- and JAK3-deficient SCID to have extensive warts and in this study there was no difference in the mean values of NK cells at last follow up between those with or without warts or between conditioned and unconditioned recipients.(57) Although the pathophysiologic mechanism is not completely understood, evidence has been provided that lack of the common y chain function in cells such as keratinocytes may impair the secretion of chemokines that may guide the influx of protective immune cells. Therefore changing the cells in the haematopoietic system by HSCT or gene therapy which does not alter cells out-with the haematopoietic system leaves these patients at risk of developing HPV- associated warts and vigilance is required to monitor for any malignant transformation. Most centres would recommend HPV vaccination for these patients and further studies are needed to see if this will reduce the incidence

Kesserwan $\it{et\,al.}$ reported an association between Dermatofibrosarcoma protuberans (DFSP) and ADA deficient SCID patients. DFSP is normally a particularly rare malignant skin tumour in childhood with a low risk of metastasis.(58) The characteristic histologic finding is a spindle cell tumour with a storiform pattern which is CD34+. At the cytogenetic level it is associated with a characteristic chromosomal translocation (t[17;22] [q22;q13]) resulting in the COL1A1-platelet-derived growth factor β (PDGFB) fusion gene. Kesserwan $\it{et\,al.}$ described 8 patients with DFSP lesions. Mostly these presented as multiple small brown atrophic plaques, less than 1cm in diameter which had been present in some cases since birth. Cases have been reported post treatment with PEG-ADA and HSCT suggesting that despite immunological correction of the underlying disorder this association remains a risk. (58-60) Toxic metabolites in the skin and an increased propensity for DNA strand breaks in patients with ADA deficiency may be involved as mechanisms for this association. Careful histological and cytogenetic testing of lesions is required to make the diagnosis and treatment requires wide excision with margin control due to the infiltrative nature of the lesions.

The impact of age of risk factor of MPT has been studied in a number of large registry studies in HCT for malignancy. CIBMTR assembled a cohort of 1487 paediatric HCT and showed that

younger age (<10 years) at allogeneic HCT was associated with greater risk of MPT, especially brain and thyroid tumours (RR 3.7) but had no effect on autologous HCT. (24, 25) There is no study to analyse age at HCT in patients with PID.

Transplantation-related risk factors

Chemotherapy

Many studies have shown that patients who have been treated with chemotherapy and/ or radiotherapy followed by HCT have an increased risk of developing secondary cancer. However, MPT has also been reported in patients who only received chemotherapy alone for HCT. In animal models, a higher incidence of secondary cancer was also found in dogs that were given DLA-identical marrow from a litter mate after TBI than in controls dogs or dogs given only chemotherapy for conditioning.

In PID patients with increased susceptibility to cancer, use of chemotherapy particularly alkylating agents induce DNA breakage which may predispose to malignant transformation. In recent years the use of reduced intensity and reduced toxicity conditioning has increased which may reduce the risk of malignancy post-HSCT.

In Unni *et al.*'s study no patients received radiotherapy but all received chemotherapy with at least 1 alkylating agent for conditioning prior to HCT (25).

Fludarabine-based conditioning, moderate-severe chronic GVHD and chronic myeloproliferative or non-malignant disease were shown to be risk factors for second malignancy in adult patients.(61, 62). Shimoni *et al.* found no significant difference in the incidence of secondary malignancies in 931 adults receiving myeloablative, reduced intensity or reduced toxicity conditioning and postulated that there may be synergistic effects of DNA damage from an alkylator added to fludarabine related inhibition of DNA repair used in reduced intensity or toxicity regimens. (62)

Medication

Voriconazole is commonly used in patients with PID pre- and post-HSCT. A report documented an association between voriconazole and the development of squamous cell carcinoma post-allogenic HSCT.(63) A patient in Unni *et al.*'s study with XL-CGD had fungal granuloma pre-transplant, received voriconazole throughout transplant and developed actinic keratosis, followed by squamous cell carcinoma of the lower leg and an auricular basal cell carcinoma.

Graft versus host disease and viral infection

In Unni's study 2 patients who both had oral cGVHD and prolonged HHV6 viraemia post-HCT developed a parotid muco-epidermoid carcinoma (MEC) at 6 and 3 years post-HSCT. There are data which link parotid MEC to prolonged CMV infection, which can remain dormant in the salivary glands and so the presence of HHV6 in a patient with an immature immune system may play a role. Two additional patients had GVHD which together with prolonged immunosuppressive treatment may alter the microenvironment and depress normal tumour immune surveillance. (64)

Donor chimerism

In Unni et al.'s study, two patients lost donor engraftment in whole blood or B- cell and myeloid cell lineages before the onset of Philadelphia positive acute lymphoblastic leukemia in one and juvenile myelomonocytic leukemia in another. Both malignancies were confirmed to be recipient in origin. The first patient had RAG 2 deficiency and the second Griscelli syndrome, both of which predispose to malignancy. Alternatively, recipient stem cells surviving chemotherapy may have acquired genotoxic insults. Therefore the question arises that if they had had full donor chimerism this may not have happened.

Non-transplantation related risk factors

As for the general population lifestyle habits are important risk factors for the development of malignancies. Tobacco and alcohol consumption, obesity, excessive UV light exposure and a family history of malignancy may increase the risks.

Strategy to minimise risks of malignancy post haematopoietic cell transplantation

Optimising transplant related factors

Outcomes of transplant for PIDs have improved dramatically over the last 20 years due to multiple factors. Moving towards low intensity rather than highly toxic traditionally used myeloablative regimens has led to less toxicity both in the short-and long-term.

Whilst moving away from toxic alkylating agents may decrease the risk of malignancy, an increase in mixed chimerism may leave the patient with recipient cells with a predisposition to malignancy depending on the underlying disorder and therefore choosing the best conditioning regimens remains a balancing act. A personalized approach to each patient with the ability to use pharmacokinetic monitoring of conditioning agents and blood levels of

serotherapy drugs will enable physicians to maximise the chance of donor engraftment with good levels of donor chimerism and immune reconstitution whilst minimising short- and long-term toxicities. (65-68)

The ability of certain antibodies to open up the haematopoietic stem cell niche is an exciting prospect which may allow conditioning in the future without the need for toxic chemotherapy. CD45 is selectively expressed on all leucocytes and haemopoietic progenitors, but is absent on non-haemopoietic tissues. Straathoff and colleagues reported 16 patients with PID who were less than one year of age or had significant pre-existing co-morbidities. The conditioning regimen comprised alemtuzumab 0.2 mg/kg daily for 3 days for unrelated donors, or 0.1 mg/kg daily for matched sibling donors on day -8 to day -6, clinical grade rat anti-CD45 (YTH 24.5 and 54.12) 0.4 mg/kg on day -5 to day -2, fludarabine (30 mg/m² daily for 5 days on day -8 to day -4) and cyclophosphamide (300 mg/m² daily for 4 days on day -7 to day -4). Twelve patients were alive and well at the end of the study, one failed to engraft and was successfully re-transplanted and 3 died – none of conditioning toxicity. Donor chimerism was variable but high level and sufficient to cure disease in the survivors. (69)

A clinical trial is currently in progress using anti-CD117 antibody to treat patients with primary immunodeficiencies (AMG191 Conditioning/CD34+CD90 Stem Cell Transplant Study for SCID Patients, ClinicalTrials.gov Identifier: NCT02963064). This antibody to CD117 (otherwise known as c-kit receptor) selectively depletes haematopoietic stem cells. (70) Early results of this dose finding study reported that some donor stem cell chimerism, leading to donor T-and B-lymphocyte chimerism can be achieved.(71)

The level of donor chimerism is not just related to the conditioning regimen but also to factors such as stem cell dose in the transplant product. Good results have been obtained with well-matched donors using peripheral blood stem cells which lead to a higher stem cell dose than traditionally used bone marrow, without an increase in significant graft versus host disease when alemtuzumab is used.(72)

Other ways to improve chimerism post-transplant include the use of additional cells as an add-back either in the form of stem cells or T cell lymphocytes.

GVHD

Improved methods of HLA typing mean that better matched donors are now being used which decreases the risk of GVHD. In addition, timing and dosing of serotherapy used in the conditioning regimen are critical for the prevention of GVHD but need to be balanced against the impact on time to immune reconstitution particularly in the setting of PID with viral infections. Biomarkers to detect patients at risk of GVHD may become part of routine

practice enabling patient specific preventive strategies. If GVHD occurs prompt treatment is important. Methods such as Extracorporeal photopheresis (ECP) are now widely available which will improve previously poor outcomes of patients with steroid-resistant acute and chronic GvHD.(73) All these aspects will limit the impact of GVHD on carcinogenesis.

Optimizing non-transplant-related factors

Giving patients advice on lifestyle choices and avoiding high risk behaviours such as smoking and sun-bathing is important for minimising the risk of malignancy.

Surveillance for MPT in PID transplant survivors

All patients who undergo HCT for PID should have long-term follow up. Once they become adults, transition to adult services is very important. Patients should be counselled regarding the possible risk of malignancy and in addition to specific follow up according to their underlying diagnosis, type of HCT and any complications, should take part in routine screening for the general population together with encouraging self-examination.

Treatment of MPT in patients with underlying PID

There is no data about specific treatment of various malignancies post-transplant in patients with PID and so conventional treatment of the malignancy is recommended.

Conclusion

Second malignancy post HCT for a malignant disease is well known and well-established long-term follow up recommendations include surveillance for secondary malignancies. The occurrence of malignancy in patients with PID is known to be higher than in the general population, but the risk of malignancy post HCT in this group of patients is less well known. In theory if the underlying defect which may predispose to malignancy is only in the haematopoietic cells and the defect is cured, the risk of malignancy should be reduced. However, many PIDs, such as DNA repair defects, have features outside the haematopoietic cells which may therefore still pose a risk of developing disease-related malignancies. Incomplete donor chimerism post HCT may leave a patient at risk of malignancies in affected haematopoietic cell lines that are not completely corrected. In addition, the procedure of the transplant itself including use of alkylating agents and risk of complications such as GVHD and viral infection may increase risks of malignancy occurring. Therefore, it is important to look for malignancies as part of long-term follow up for patients post HCT for PID.

Expert Opinion and future perspectives

HCT is a curative treatment for an increasing number of PIDs. Survival after transplant has improved enormously in the last 20 years. Good quality of life with long-term healthy immune reconstitution and minimal late effects is therefore the goal. A systemic review or meta-analysis of existing data would be helpful to definite the risk factors of MPT.

Large scale, long-term follow-up of PID transplant survivors is required, including surveillance for malignancies, to improve our understanding of the spectrum of malignancies, better define the risks and therefore improve the way we perform HCT to minimise these risks. Multi-centre studies are needed.

A personalised approach to transplant will become more common. Biomarkers to identify patients at risk of developing malignancy may become available in order to allow tailoring of the approach to HCT and post HCT surveillance. PK monitoring with targeted dosing of chemotherapeutic agents and measuring levels of serotherapy used for conditioning, together with precise cell dosing in grafts, are becoming more common and in 5 years' time may become routine practise. Reduced toxicity conditioning will continue to evolve, for example using antibodies to open up the stem cell niche and thus avoid the use of DNA damaging alkylators which may have an effect on tissues out with haematopoietic cells.

Strategies to prevent and treat GVHD which has a role in carcinogenesis will continue to improve. For example, in our own centre the choice of donor for a patient with severe combined immunodeficiency if they do not have a matched sibling donor is a haploidentical parent using CD3+ alpha beta depletion together with CD19+ depletion to prevent GVHD and post-HCT LPD. (74, 75) Studies are ongoing using alternative techniques for haploidentical transplants including the use of post HCT cyclophosphamide, CD45RA depletion which retains the CD45RO-bearing T lymphocytes which are more likely to confer antiviral activity, giving additional caspase gene modified T cells which can be switched off if GVHD occurs, or CD45RA depleted cells post-HCT to improve immune reconstitution. (76-79) It may be that these techniques will supercede the use of unrelated donors, with their higher risk of causing GVHD, for all patients with PID.

Monitoring for infection and pre-emptive treatment will continue to improve. The use of T lymphocytes directed against specific viral epitopes will become more widely available. These can be either donor-derived or from third party banks. (80, 81) This will minimise the risks of carcinogenesis caused by viral infections such as CMV.

Finally, international standards for follow up of patients post HCT for PID should be developed which include surveillance for both disease-specific and other forms of malignancy which occur.

Table 1 Reported malignancies in PID patients from registry studies

Author Year	Ref	Year	Study cohort	Median I age at p dx of PID v (range) r	No. of patients with malignancy	No. of Median patients age at with diagnosis of malignancy malignancy (range)	Primary diagnosis of patients with malignancy	Type of cancer Remarks	Remarks
Mayor 2018	[3]	2003-	3658	₹ Z	171 (4.9%)	NA V	119 (70%) CVID 13 (9%)	82 (48%) Ivmphoma	1.42-fold excess
) -)) -) I	35% CVIDD				hypogammaglobulinemia		compared with
United			13% CGD				and agammaglobulinemia cancers	cancers	the age-adjusted
States			12% DiGeorge				8 (4.6%) WAS	14 (8%)	population
Immune			syndrome				4 SCID	genitourinary	Male PID: 1.91
Deficiency			7% SCID				3 ataxia telangiectasia	14 (8%)	excess relative risk
Network			7% WAS				3 hyperIgE STAT3	gastrointestinal	Female PID: no
(USIDNET)			4% HyperIgM				2 cartilage hair hypoplasia cancers	cancers	excess risk
			syndrome				1 DOCK8	10 (6%) breast	
			23% others				1 Activated PI3K delta	cancer	Male PID: 10-
							1 CID	(2%)	fold increase for
							16 others	endocrine	lymphoma
								cancers	Female PID: 8.34-
								6 (4%) head and	6 (4%) head and fold increase for
								neck cancers	lymphoma
								5 (3%) lung	
								cancers	
								2 (1%) bone	
								cancers	
								4 (2%)	
								unspecified	

Year	Ref Year	Study cohort	Median age at dx of PID (range)		No. of Median patients age at with diagnosis of malignancy (range)	Primary diagnosis of patients with malignancy	Type of cancer Remarks	
Jonk- man-Berk 2015	[10] 2009- 2012	745 687 (92.9%) alive	8.0 years (range, 0 - 80	60 (8.1%)	43.0 years (range, 3.0 -77.0)	44 (73.3%) 3 leukaemia hypogammaglobulinemia 17 lymphomas (78 CVID)	3 leukaemia PID had an overall 17 lymphomas 2.3-fold increase in (14 NHL. 3 HD) the life time chance	overall ease in
) -) 		and 58 (7.8%) died	years)					5 5
Nether- lands		before the study analysis					40 (66.7%) solid a malignancy cancers compared to	: > 0
		1118 had HCT (38					14 skin (13 SCC) general population 6 in Netherlands	ulation nds
		died)					gastrointestinal	
		61% predominantly					5 breast	
		antibody disorders					4 endocrine	
		11% phagocytic					3 lung	
		disorders					2 thymus	
		9% other well-de-					2 genital male	
		fined PID					2 bladder	
		8% predominantly					1 bone	
		T cell deficiencies					1 pancreas	
		4% Autoimmune						
		and immune dys-						
		regulation						
		4% unclassified im-						
		munodeficiencies						
		2% complement						
		deficiencies						
		1% defects in						
		innate immunity						

Author Ref Year Year	Ref	Year	Study cohort	Median age at dx of PID (range)	Median No. of age at patients dx of PID with (range) malignancy	No. of Median patients age at with diagnosis of malignancy (range)	Primary diagnosis of patients with malignancy	Type of cancer Remarks	. Remarks
Vajdic 2010	[31]	[31] 1990- 2008	1132	Median age at	58 (5.7%)	₹Z	53 predominantly antibody deficiencies	55 first cancer 3 second	Standardized incidence ratio
			78% predominately regis-	regis-	55 first			cancer	all cancers: 1.6
Australia		Mean	\sim	tration:	cancer				thymus: 67.3
		duration	7% complement	25 years	3 second			16 NHL	NHL: 8.82
		of FU:	deficiency	(range,	primary			1 HD	stomach: 6.10
		16 years	16 years 4.8% Combined T	0 – 95	cancer			4 leukaemia	leukaemia: 5.36
		(SD 5.4	and B cell immuno- years)	years)				3 stomach	
		years)	deficiency					2 thymus	
			5.8% other well-de-					1 adrenal	
			fined immunodefi-					1 thyroid	
			ciency syndromes					3 prostate	
			3.5% congenital de-					3 bladder	
			fects of phagocyte					1 ovary	
			number					12 breast	
			1.2% diseases of					4 melanoma	
			immune dysregu-					2 trachea,	
			lation					bronchus, lung	
								1 pancreas	
								4 colon	

CGD: chronic granulomatous disease; CID: combined immunodeficiency; CVID: combined variable immunodeficiency; FU: follow-up; HCT: haematopoietic cell transplantation; HD: Hodgkin's lymphoma; NA: not available; NHL: non-Hodgkin's lymphoma; PID: primary immunodeficiency; Ref: reference; SCID: severe combined immunodeficiency; SCC: squamous cell carcinoma; WAS: Wiskott-Aldrich syndrome

Table 2 Reported malignancy post allogeneic haematopoietic stem cell transplant in children with haematological disorders

Ref	f Year of	Study	Age at	No of MPT	Primary T	Type of	Interval be-	Donor for	Conditi-	Outco-	Remarks
	duration of FU		<u>.</u>		ν				7	patients with MPT	
18	[50] 1969-	4905	Adult	499	Z YZ	₹Z	Median 10.3	NA AN	NA AN		CI of MPT: 22%
	2014	(excluded	and	(11%)			years (range				at 30 years post-
		Fanconi	pae-	MPT			1.0 to 39.7				HCT
	Median	anaemia,	diatric	(ex-			years)				
	dura-	n = 20 and	cohorts	cluded							Highest incidence
	tion FU	non-hae-		SCC							of MPT was in
	(living):	matological	Median:	and							survivors exposed
	12.5	solid tumour,	34.5	BCC of							to unfractionated
	years	n=140)	years	skin)							(600-100cGy) or
	(range,		(range,								high dose frac-
	1.0 to 42	. 4500 hae-	0.3 –	81 had							tionated (14.4 to
	years)	matological	78.9	-Inw							17.5Gy) TBI.
		malignancy	years)	tiple							For patients who
		405 non-		MPT							received low dose
		malignant		69)							TBI, the incidence
		diseases (AA,		had 3							was comparable
		PID, HLH,		MPT, 9							to myeloablative
		haemoglob-		had 3							chemotherapy
		inopathy and		and 3							alone, but still
		others)		had 4)							2-fold higher than
											general popula-
											tion.

Omori		HCT/ cohort duration of FU	study cohort	Age at HCT	No of MPT	Primary diagnosis of patients who deve- loped MPT	Type of cancer	Interval be- tween HCT to MPT	Donor for Conditipations oning for who departients veloped who MPT developed MPT	٠ ه	Outco- me of patients with MPT	Remarks
2013	[46]	[46] 1995- 2010	370 113 AML	Adult and pae-	11 MPT in 10 pa-	3 ALL 2 AML 3 lympho-	1 thyroid papillary carcinoma	Median: 6.8 years (range, 1.4 to 15.2	Allogenic HCT Donor	All had TBI	3 died of MPT 1 died of	Incidence of invasive solid MPT 2.15±1.22 and
Single centre		Median duration	117 ALL 39 lympho-	diatric	tients	ma 2 MDS	1 submax- illary gland	years)	details: NA		aspiration pneumo-	6.46±2.82% at 5 and 10 years
Japan Japan		on Fu. 10.5 years (max 16.4 years)	34 MDS 41 CML 6 dyshema- topoiesis All allogenic	Medi- an: 36 years (range 1-72 years)			2 oesopha- geal cancer 1 oral cavity carcinoma 2 gastric cancer 1 ureteral cancer 1 head and neck cancer 1 border ma- lignant ovari- an tumour 1 extragonad- al germ cell				ت = ع	

													_
Author Year	Ref	Year of HCT/ duration of FU	Study cohort	Age at HCT	No of MPT	Primary diagnosis of patients who deve- loped MPT	Type of cancer	Interval be- tween HCT to MPT	Donor for Conditi- patients oning fo who de- patients veloped who MPT develop	Conditi- oning for patients who developed	Outco- me of patients with MPT	Remarks	
Rizzo	[27]	1964-	28874	Adult	EX-	AN	17 carcinoma	NA	Allogenic	NA	NA AN	CI of solid MPT:	
2009		1996	(excluded	and	cluded		in situ of skin		HCT			Competing risk	
			Fanconi	pae-	PTLD,		19 invasive		Donor			analysis: 1% at 10	
CIBMTR		Duration	Duration anaemia and	diatric	BCC		SCC of skin		details:			years, 2.2% at 15	
and		of FU:	PID)	cohorts	and				NA			years, 3.3% at 20	
Seattle		¥ N			leukae-		27 oral/pha-					years post-HCT	
			5916 (20.5%)		mia/		ryngeal					Kaplan-Meier	
			ALL	Medi-	MDS		18 melanoma					analysis: 2.5% at	
			7461 (25.8%)	an: 27			18 braub					10 years, 5.8% at	
			ANLL	years	189		16 thyroid					15 years, 8.8%	
			7594 (26.3)	(range	MPT		13 breast					at 20 years post-	
			CML	0.08			9 female					HCT	
			452 (1.6%)	-72.4			genital						
			other leukae-	years)			8 bronchus/					Latency: 65 MPT	
			mia				lung					occurred 1-4	
			1152 (4.0%)				7 liver					years and 100≥5	
			NHL				7 soft tissue					years post-HCT	
			242 (0.8%)				6 bone/joint						
			H				3 oesophagus					The rate of solid	
			507 (1.8%)				3 testes					MPT was at 2.1-	
			Myeloma				2 colon					fold higher than	
			158 (0.5%):				4 rectum /rec-					general popula-	
			other malig-				tosingmoid/					tion and reached	
			nancies				anus					3-fold among	
			2842 (9.8%)				12 other solid					patients followed	
							cancers					for ≥ 15 years	
												post-HCT	

Author Year	Ref Year of HCT/ duration of FU		Age at HCT	No of MPT	Primary diagnosis of patients who deve- loped MPT	Type of cancer	Interval be- tween HCT to MPT	Donor for Conditipations oning for who departments on the conditions on the condition of th	 Outco- me of patients with MPT	Remarks
		SAA								Risk factors:
		1298 (4.5%)								Conditioning
		MDS/MPD								radiotherapy (TBI
		906 (3.1%)								or limited field):
		Haemoglob-								non-SCC tumour
		inopathies								- RR 2.3 for TBI
		346 (1.2%)								Age at HCT: RR
		other								55.3 for < 10
										years; RR 6.2 if
		All allogeneic								10-19 years; 4.8
		HCT								if 20-29 years; no
		631 (2.2%)								excess risk if > 30
		Twin								years
		22030 (76.								cGvHD: SCC –
		3%) MSD								skin RR 11.0, oral
		2939 (10.2%)								RR 5.3
		MFD								Male: SCC – skin
		3066 (10.6%)								RR 11.9, oral RR
		QN								2.8
		208 (0.7%)								Most cases of
		other/uncer-								CNS, thyroid,
		tain								bone, soft tissue
										MPT occurred in
										patients who un-
										derwent HCT at <
										17 years of age

	t)	0				.u.		~		nin	-cell	([3R		neu	L								
Remarks	The observed to	expected ratio	was 29.7%		Risk factors:	T cell depletion:	RR 3.1 - 94	Use of ATG: RR	3.8	HLA mismatch in	presence of T-cell	depletion/ATG	use: RR 3.8	Acute (RR 1.7)	and chronic (RR	2) GvHD	Less effect when	T-cell depletion	used agents	which also re-	move B cells.					
Outco- me of patients with MPT	ΑΝ																									
Conditi- oning for patients who developed MPT	NA																									
Donor for Conditipations on the patients on the patients veloped who MPT MPT	ΝΑ																									
Interval be- tween HCT to MPT	105 (83%)	occurred	within first	year post-	HCT		22 (17%)	late onset,	1->10 years	post-HCT																
Type of cancer	PTLD																									
Primary diagnosis of patients who deve- loped MPT	ΑN																									
No of MPT	127	PTLD		Other	MPT	not	includ-	eq																		
Age at HCT	Adult	and	pae-	diatric	cohorts		Median:	56.6	years	(0.1-68	years)															
Study	26901	(exclud-	ed PID,	inherited	cancer pre-	disposition	syndromes,	initial diagno-	sis of NHL)		11582 ALL	14736 ANLL	14916 CML	5576 AA	6992 Other		All allogeneic	HCT	20873	(77.6%) MSD	2828 (10.5%)	MMFD	3005 (11.2%)	OD	195 (0.7) oth-	er/uncertain
Ref Year of HCT/ duration of FU	1964-	1996		Median	duration	of Fu NA																				
Ref																										
Author Year	Landgren [45]	2009		CIBMTR	and	Seattle		(update	of Curtis,	1999)																

Age at No of Primary Type of HCT MPT diagnosis cancer of patients	s	Type of cancer	Interval be- tween HCT	Donor for patients who de-	Conditi- oning for	Outco- me of patients	Remarks
who deve- loped MPT	hodeve-			veloped MPT	who developed MPT	with MPT	
Adult 32 10 ALL 32 thyroid	, ,	32 thyroid	Median: 8.5	23 allo-	7 chemo-	25 total	CI of thyroid MPT
and thyroid 7 CML carcinomas		carcinomas	years (range,	genic (21	therapy	thyroidec-	~0.05% at 20
57999 (82%) pae- carci- 4 AML	AML		0.6 to 22.2)	MSD; 2	alone	tomy	years post-HCT
haematologi- diatric nomas 1 MDS Median age		Median age		unrelated)	23 TBI +	5 subtotal	(0.2% in patients
cal cancer cohorts (23 1 NHL at diagnosis		at diagnosis		9 autolo-	chemo-	thyroidec-	0-10 years of age
8416 (12%) papil- 1 unclas- of MPT: 23.5		of MPT: 23.5		gons	therapy	tomy	at HCT.
solid tumour 15% lary, 9 sified years (range,		years (range,			1 tho-	1 Partial	
4391 (6%) AA <10 follicu- leukaemia 8.8 to 52.2		8.8 to 52.2		31 unma-	racoab-	1 lobecto-	18 (56%) present-
10% lar) 3 AA years)		years)		nipulated	dominal	my	ed with palpable
10-20 4 neuro-	neuro-			marrow	irradiation	23 ra-	nodules
blastoma	lastoma			1 PBSC	+ chemo-	dioactive	9 (28%) was as-
75% 1 breast	breast				therapy	iodine	ymptomatics and
(45%) allo- >20 cancer	ancer					ablation	diagnosed only
							on ultrasound
38988 (55%)						29 alive	surveillance.
s Median	ledian					and	
100 (0.1%) age at HCT:	ge at HCT:					disease	Risk factors:
11.2 years	1.2 years					free after	Age at HCT: RR
(range, 1.7	ange, 1.7					a median	24.6 for < 10
to 51.3	51.3					period of	years vs > 20
years)	ears)					5.5 years	years
						from MPT	TBI or thora-
						(range, 0.3	co-abdominal
						to 11.5)	radiotherapy (RR
							3.4)
						الم المرام د	Females RR 2 8 vs

		9% ost- ng in
Remarks	males cGvHD RR 2.9	CI of MPT: 5.9% at 20 years post-HCT, increasing by about 2% in each successive 5 years follow-up period. Most solid MPT occurs after 5 years, 3.8% at 20 years. Risk factors for PTLD: MMFD (RR 9.0) PID (RR 2.7) CMI (RR 2.5)
Outco- me of patients with MPT	relapse of primary disease 1 died of progres- sion of MPT	PTLD: 79% (34/43) died AML/MDS 79.4% (27/34) died Solid MPT: 42% died of solid MPT MPT
Donor for Conditi- patients oning for who de- patients veloped who MPT developed		₹ Z
		∢ Z
Interval be- tween HCT to MPT		PTLD Median: 0.3 years (range, 0.1 to 7.3 years) NA for MDS/ AML and solid MPT
Type of cancer		44 PTLD 36 AML/MDS 5 other leukaemia/ lymphoma 62 solid MPT Type of solid MPT 4 adenocarcinoma 11 BCC 4 breast 1 carcinoid (rectal) 5 carcinoma in situ
Primary diagnosis of patients who deve- loped MPT		₹
No of MPT		MPT in 137 pa-tients
Age at HCT		Adult and pae-diatric cohorts Median: 24 years (range, 0.1 to 67)
Study		3372 211 AA 117 PID 215 IEM 582 ALL 602 CML 119 MDS 342 NHL 124 HD 102 neuro- blastoma 138 Breast cancer 226 other
Year of HCT/ duration of FU		(49] 1974-2001 Median duration of follow-up: 5 years (range, 0.5 – 2.5 years)
Ref		[49]
Author Year		Baker 2003 (update of Bhatia, 1996) Single centre study Minne- sota

Remarks	in vitro TCE (RR	4.0)	Grade 3-4 GvhD	(RR 2.4)		Risk factors for	solid MPT:	Age ≥ 20 years at	HCT (RR 2.0)	TBI not significant		Risk factors for	AML/MDS:	230 of 34 had	autologous HCT	Risk highest for	PBSC (RR 3.1)								
Outco- me of patients with MPT																									
Conditi- oning for patients who developed																									
Donor for Conditipations on ing for who departments or who who MPT MPT																									
Interval be- tween HCT to MPT																									
Type of cancer	2 carcinoma	(primary	unknown)	1 CML	1 CNS, astro-	cytoma	1 CNS,	ependymoma	1 CNS, glioma	1 CNS, neu-	roectodermal	tumour	4 HD	3 parotid MEC	1 neuroblas-	toma	1 papillary	thyroid carci-	noma	1 renal cell	carcinoma	1 sarcoma,	bone	1 angiosar-	coma
Primary diagnosis of patients who deve-																									
No of MPT																									
Age at HCT																									
Study	1287 MFD	120 MMFD	373 MUD	399 MMUD		1193 autolo-	Sons																		
Ref Year of Study HCT/ cohort duration of FU																									
Author R Year																									

				_	_					S	90	St		%:	St-		%9	st-			had				
rks									CI of solid MPT:	1.6% at 5 years	post HCT; 6.1%	at 10 years post		Allogeneic: 6.4%	at 10 years post-		Autologous: 1.6%	at 10 years post-			All 6 skin SCC had	0			
Remarks									CI of so	1.6% a	post H	at 10 y	HCT	Alloger	at 10 y	H	Autolo	at 10 y	HCT		All 6 sk	CGVHD			
Outco- me of patients with MPT									NA A																
Conditi- oning for patients who developed									NA A																
Donor for Conditipations on the patients on the who developed who MPT MPT									×N ∀N																
Interval be- tween HCT to MPT									Median,	years (range)	Cervix, 3.3	(1.6-9.7)	Thyroid, 12.7	(7.5-18)	Oral cavity,	7.6 (4.7-11.7)	Liver, 10.8	(6.7-14.9)	Breast, 9.9	(2.6-17.1)					
Type of cancer	1 fibrosarco-	ma	1 liposarcoma	1 rhabdomyo-	sarcoma	1 lung SCC	1 oral cavity	SCC 1 skin SC	3 SCC	6 BCC	4 cervix uteri	2 salivary	gland	3 SCC of oral	cavity	2 breast	cancer	2 liver cancer	(had hepatitis	0	2 thyroid	1 astrocytoma	1 malignant	fibrous histio-	cytoma
Primary diagnosis of patients who deve- loped MPT									2 ALL	5 AML	1 AA	1 NHL	5 CML												
No of MPT									29	solid	MPT		Ë	cluded	PTLD	and	hae-	mato-	logical	malig-	nan-	cies			
Age at HCT									Adult	and	pae-	diatric	cohorts		Median:	33.9	years	(range,	1.5-71.5	years)					
Study									2129		327 ALL	648 AML	447 NHL	241 HD	392 CML	73 AA		1155 related	donor	213 UD	759 autolo-	gous			
Ref Year of Study HCT/ cohort duration of FU									[43] 1976-	1998		Median		tion of		years	(range,	0.1 to	21.1	years)					
									[43]																
Author Year									Bhatia	2001		Single	centre	City of	Hope										

		%, . at	s as- age ars r TBI (RR (RR
Remarks		CI of invasive solid MPT 0.9%, 4.3% and 11% at	5, 10, 15 years post-HCT Solid MPT was associated with age at HCT <10 years of age (RR 3.7) and high dose TBI (RR 3.1) Chronic GvHD lowered risk of solid tumours (RR 0.2) PTLD was associated with moderate/severe cGvHD (RR 6.5),
Outco- me of patients with MPT		All died	12 died (all pa- tients with brain MPT died)
Conditi- oning for patients who developed		¥Z	All had radiation
Donor for patients who developed		Allogenic HCT Donor	
Interval be- tween HCT to MPT		Median: 1.5 years 15 - first	year post- HCT 4 - between 12 and 18 months 1 - 4.9 years post-HCT Median: 6 years (range, 0.3 to 14.3 years)
Type of cancer	1 SCC oe- sophagus 1 synovial sarcoma	20 PTLD	year p HCT HCT 12 and month 12 invasive Media solid cancers years 9 brain cancer years) 5 thyroid papillary carcinoma 3 tongue SCC 3 melanoma 2 salivary MEC
Primary diagnosis of patients who deve- loped MPT		11 ALL 9 ANLL	Age at HCT Median: 5.8 years (range 0.4 to 16.1 years) 12 ALL 11 ANLL 2 un- specified leukaemia Age at HCT Median: 8.2 years (range,
No of MPT		45 (1.4%)	anoma in situ
Age at HCT		Pae- diatric cohort	
Study cohort		3182 2022 (63.5%)	
Year of HCT/ duration of FU		[24] 1964- 1992	Median duration of FU for patient survived > 1 year: 3.6 years (range, 1-20.7 years
Ref		[24]	
Author Year		Socie 2000	and Seattle

Autnor Year	Ref	Year of HCT/ duration of FU	Study cohort	Age at HCT	No of MPT	Primary diagnosis of patients who deve- loped MPT	Type of cancer	Interval be- tween HCT to MPT	Donor for patients who de- veloped MPT	Conditi- oning for patients who developed	Outco- me of patients with MPT	Remarks
						w.	2 osteosar-					MUD and MMFD
						years)	coma 1 MFH					(RK 7.5), ICD (\$\$ 4.8), and ATG (RR
Kolb	[42]	Before	1036	Adult	53	Ž	14 skin	Υ _Z	52 allo-	8 chemo-	10 died	3.1) Cumulative
1999	,	1986		and	(5.1%)		7 oral		genic	therapy		incidence of MPT
			302 AML	pae-			5 gut		1 autolo-	only		3.5±0.6% at 10
EBMT		Median	212 ALL	diatric			5 thyroid		Sons			years post-HCT
		duration	208 CML	cohorts			6 uterine/			45 radia-		and 12.8±2.6%
		of FU:	185 AA				cervix			tion plus		at 15 years post-
		10.7	35 lympho-	Medi-			4 breast			chemo-		HCT.
		years	ma	an: 21			3 brain			therapy		
		(range,	9 IEM	years			1 leukaemia					The rate of MPT
		5-22.1	32 solid	(range,								was 3.8-fold high-
		years	tumours	1 to								er that in an age-
				51.9								matched control
			All allogeneic	years)								population,
												More common
												in older patients
												and patients with
												CGVHD

Author Year	Ref	Year of HCT/ duration of FU	Study	Age at HCT	No of MPT	Primary diagnosis of patients who deve- loped MPT	Type of cancer	Interval be- tween HCT to MPT	Donor for Conditipations on the patients on the patients veloped who MPT develope	r be	Outco- me of patients with MPT	Remarks
Curtis	[40]	1964-	19229 (ex-	Adult	80	16 ALL	17 Buccal cav-	NA AN	NA A	NA A	36 died	CI of solid MPT:
1997		1992	cluded NHL,	and	solid	28 ANLL	ity/pharynx (1					2.2% at 10 years
			Fanconi and	pae-	MPT	14 CML	neuroblasto-					post-HCT and
CIBMTR		Median	PID)	diatric		8 AA	ma of nasal					6.7% at 15 years
and		dura-		cohorts		4 lympho-	cavities; 4					post-HCT
Seattle		tion of	4245 (22%)			ma/other	MEC)					
		FU: 3.5	ALL	Median:			1 colon					Ratio to observed
		years	5208 (27.1%)	25.5			2 rectum					to expected cas-
		(range,	ANLL	years			3 liver (2 ma-					es: 2.7 (p<0.001)
		>1- 25	4885 (25.4%)	(range,			lignant fibrous					
		years)	CML	0.2			histiocytomas)					Risk factors:
			729 (3.8)	to 67			2 lung					Younger age at
			NHL	years)			7 breast					HCT
			561 (2.9%)				1 cervix					Higher dose TBI
			other cancer				1 uterine					Chronic GvHD
			643 (3.3%)				2 testis					Male
			MDS				11 brain (4 as-					
			2159 (11.2%)				trocytoma, 6					
			AA				glioblastoma,					
			799 (4.2%)				1 PNEC)					
			Other				8 thyroid					
							9 bone and					
			533 (2.8%)				connective					
			syngeneic				tissue (3					
			15217 (79%)				chondrosar-					
							coma,					

									%6:	post-		1PT:	/ears		pani		MDS	ars			-BI (RR	was	ctor	L-)BSC	>35
Remarks									CI of MPT: 9.9%	at 13 years post-	HCT	CI of solid MPT:	5.6% at 13 years	post-HCT	PTLD plateaued	at 1.6% at 4	years, AML/MDS	at 2.1% 9 years		Risk factors:	Solid MPT: TBI (RR	6.0); cGVHD was	not a risk factor	for skin MPT	AML/MDS: PBSC	(RR 5.8), age >35
Outco- me of patients with MPT									36 died																	
Conditi- oning for patients who developed MPT									AN																	
Donor for patients who de- veloped MPT									38 related NA	donor	e ud		18 autolo-	Sons												
Interval be- tween HCT to MPT									PTLD	Median: 0.2	years (range,	0.1 0 3.0	years)		AML/MDS	Median: 3.0	(range 0.3 to	9 years)		Solid tu-	mours:	Median: 4.0	(0.2 - 13)	years)		
Type of cancer	1 osteo-	sarcoma, 2	rhabdomyo-	sarcoma, 2	fibrosarcoma,	1 unspecified)	11 melanoma	5 other	22 EBV-relat-	ed PTLD	11 AML/MDS	2 NHL	1 HD		17 solid MPT	3 melanoma	2 brain tu-	mour	3 BCC	1 SCC	1 osteosar-	coma	1 papillary	carcinoma of	thyroid	1 malignant
Primary diagnosis of patients who deve- loped MPT									26 leukae-	mia	25 lympho-	ma	8 PID	1 AA	1 IEM											
No of MPT									51	(2.4%)	devel-	oped	53	MPT												
Age at HCT									Adult	and	pae-	diatric	cohorts		Medi-	an: 20	years	(range,	0.2	to 67	years)					
Study	MSD	2265 (11.8%)	MMFD	1079 (5.6%)	ΔN	135 (0.7%)	other/uncer-	taın	2150		451 ALL	403 AML	419 CML	72 MDS	201 NHL	86 HD	75 NHL	75 NBL	30 Breast	cancer	150 AA	92 PID	91 IEM	80 others		
Year of Study HCT/ cohori duration of FU									1974-	1995		Median	dura-	tion of	FU: 3.1	years	(range,	0.6-18.8	years)							
Ref									[38]																	
Author Year									Bhatia	1996		Single	centre	Minne-	sota											

Author Ref Year of Study Year HCT/ cohort duration of FU	of FU	Study cohort n	Age at HCT	No of MPT	Primary diagnosis of patients who deve- loped MPT	Type of cancer	Interval be- Donor for Conditi- tween HCT patients oning for to MPT who de- patients veloped who MPT developed	Donor for patients who de- veloped MPT	Outco- me of patients with MPT	Remarks
		1063 MFD/				fibrous histio-				at HCT (RR 3.5)
		MMFD				cytoma				PTLD: in vitro TCD
		337 UD				1 breast				(RR 11.9), ATG (RR
		750 autolo-				cancer				5.9), MMD (RR
		Sons				1 prostate				8.9), PID (RR 2.5)
						cancer				
						1 carcinoma				
						in situ of cer-				
						vix and vulva				
						1 adeno-				
						carcinoma				
						of unknown				
						primary				

Author	Rof	Vear of	Study	Age at	No of	Drimary	Tyne of	Interval he. Donor for Conditi-	Donor for	Conditi	Outro-	Remarks
Year		HCT/ cohort duration of FU	cohort	HCT	MPP	ı, t²⊢	cancer	tween HCT	patients who de- veloped MPT	, g	me of patients with MPT	2
Wither-	[38]	[38] 1970-	2145	Adult	35	15 ALL	16 NHL	Median: 1.0	¥N Y	32 had TBI	30 died of	30 died of Age-adjusted
spoon		1987		and	(1.6%)	10 ANLL	6 leukaemia	year (range,			secondary	secondary incidence of
1989			1926 leukae-	pae-		4 CML	3 glioblasto-	1.5 months			cancer	secondary cancer
		Duration mia	mia	diatric		4 AA	ma	to 13.9				was 6.69 times
Single		of FU:	320 AA	cohorts		1 Hod-	3 melanoma	years)			5 survived	higher than that
centre		₹ Z				gkin's	3 SCC				after	of primary cancer
Seattle			152 synge-	Age at		1 myelofi-	1 hepatic ade-				successful	in the general
			neic	HCT		brosis	nocarcinoma				treatment	population
			1980 MFD/	Medi-			1 adenocar-				of second-	
			MMFD	an: 19			cinoma of				ary cancer	Risk factors:
			13 UD	years			rectum				(2 melano-	acute GvHD, ATG,
			101 autolo-	(range,			1 lung adeno-				ma, 1 BCC,	ma, 1 BCC, anti-CD3 mono-
			Sons	2-58)			carcinoma				1 vulvar	clonal body, TBI,
							1 invasive vul-				carcinoma,	T cell depletion,
							var carcinoma				1 B-NHL)	HLA mismatch

Bu: Busulfan; CB: cord blood; CGD: chronic granulomatous disease; CI: cumulative incidence; CIBMTR: Centre for International Blood and Marrow Transplant Research; CML chronic myeloid leukaemia; CNS: central nervous system, Cy: cyclophosphamide; EBV: Epstein-Barr virus; Flu: Fludarabine; FU: follow-up GvHD: graft-AA: aplastic anaemia; ALL: acute lymphoblastic leukaemia; ANLL: acute nonlymphoblastic leukaemia; AML: acute myeloid leukaemia; BCC: basal cell carcinoma; versus-host disease; HCC: hepatocellular carcinoma; HD: Hodgkin's lymphoma; HLH: haemophagocytic lymphohistiocytosis; JAK3: Janus Kinase 3; IEM: inborn error of metabolism; IMML: juvenile myelomonocytic leukaemia; LAD1: leucocyte adhesion deficiency type 1; MDS: myelodysplastic syndrome; MEC: mucoepidermoid matched unrelated donor, MMUD: mismatched unrelated donor; MPT: malignancy post-transplant; NA: not available; NBL: neuroblastoma; NHL: non-Hodgkin's carcinoma; Melp: Melphalan; MFD: mismatched family donor; MFH: mismatched family donor; MSD: matched sibling donor; MMD: mismatched donor; MUD: lymphoma; NFKB2: nuclear factor kappa B 2; PBSC: peripheral blood stem cell; PID: primary immunodeficiency; PNEC: primitive neuroectodermal cancer; PTLD: oos-transplant lymphoproliferative disease; RCC: renal cell carcinoma; RMS; rhabdomyosarcoma; RR: relative risk; SCID: severe combined immunodeficiency; SCCC: squamous cell carcinoma; TBI: total body irradiation; TCD: T cell depletion; Thio: thiotepa; Treo: treosulfan; UD: unrelated donor; WAS: Wiskott-Aldrich syndrome

Table 3 Reported malignancy post haematopoietic stem cell transplant in patients with primary immunodeficiency

Author Year	Year of HCT/ Duration of FU	Ref	Study cohort Age at HCT		No of Primary post-HCT diagnosis cancer	Type of cancer	Interval between HCT to malignancy	Donor	Conditioning	Outcome	Remarks
Unni	N A	[48]	944 NA	12	4 SCID	1 ALL, 1	Median: 3.75	4 MSD	7 Flu-Melp	2 died	RAG2
2018			Chil-	(1.27%)	(RAG2, JAK3,	AML, 1	years (range,	marrow	3 Flu-Treo		SCID
			dren	-uou	2 unknown)	right occip-	3 months to	4 MUD	1 Bu-Flu	10 alive	with
				PTLD		ital Ewing	11.2 years)	marrow	1 Bu-Cy	with	ALL had
				malig-		sarcoma, 1		1 MMUD		median	100%
				nancy		embryonal		marrow	10 alemtuzum-	follow-up of	recipient
						RMS of		2 MUD	ab	13.2 years	myeloid
				Age at		right cheek		PBSC	2 no serother-	(range 2	chimer-
				HCT:	1 CGD	BCC (ear)		1 matched	apy	years 1	ism;
				85		and left		CB		month to	leukae-
				months		lower leg			None had TBI	18 years)	mia was
				(range,		SCC in situ					recipient
				4-204	1 CD40	RCC					in origin.
				months)	ligand						
					1 Griscelli/	JMML					Griscelli
					HLH sys-						with
					temic						JMML
					1 GATA2	Toe mela-					had
						noma					100%
					1 LAD1	RCC					recipient
					1 MHC	Parotid					myeloid
					class I	MEC					chimer-
					1 NEFB2	Parotid					ism;
					systemic	MEC					leukae-
					1 WAS	SCC (gas-					mia was
						trostomy					recipient
						site)					in origin.

Author Year	Year of HCT/ Duration of FU	Ref	Ref Study cohort Age at HCT	Age at HCT	No of post-HCT cancer	No of Primary post-HCT diagnosis cancer	Type of cancer	Interval between HCT to malignancy	Donor	Conditioning	Outcome	Remarks
Nelson 2017	1982- 2007 Median duration of FU: 4.5 years (range, 0 to 25.4 years)	[47]	318 71 acquired AA 51 inherited AA 69 SCID 24 WAS 37 other PID 14 HLH 14 thalassae- mia 38 IEM 149 MFD 52 MMFD 78 MUD 38 MMUD 1 unknown	Median: 3 years (range, 0-14 years)	(%6:1)9	1 AA 2 FA 1 CGD 1 SCID 1 Thalassae- mia	2 leukae- mia 4 solid tumours	Median: 9.2 years (range, 0.4-14.5 years)	4 M FD 2 M M FD	2 had radiation	3 died	Cumula- tive inci- dence of MPT was 0.3±0.3% at 5 years post- HCT and 2.3±1.2% at 10 years post- HCT
Kamani 2011 CIBMTR	1968- 2003 Duration of FU: NA	[51]	2266 1075 SCID 360 WAS 831 Others	Median: 1 year (range, 1.2 months to 47 years)	52 (2.3%) all malig- nancy Median age at)	25 (2.35%) SCID 12 (3.3%) WAS 15 (1.8%) other	45 PTLD (17 con- firmed positive EBV) 3 MDS 1 AML	Median: 3.7 months (range, 0.9 to 169.6 months)	9 family donors 11 MUD 19 hap- loidentifical donors 10 MUD/	22 BuCy 1 BuCyEtopo- side 8 TBICy 5 Cy 9 none 1 CyCytarabine	40 (77%) died Cause of death: 29 post-trans- plant malig-	PID patients were at a relatively low risk of devel-oping

Author Year	Year of HCT/ Duration of FU	Ref	Author Year of Ref Study cohort Age at Year HCT/ HCT OFFU	No of Primary post-HCT diagnosis cancer	Type of cancer	Interval between HCT to malignancy	Donor	Conditioning Outcome Remarks	Outcome	Remarks
			630 MSD	HCT:	1 SCC		3 fetal	1 FluCy	4infection	malig-
			961 other	15.6	1 HCC			1 FluMelpThio	3 GVHD	nancy
			family donors	months	1 brain		40 marrow	40 marrow 1 Methyl+Cam- 4 unknown	4 unknown	post-HCT
			528 UD	(range	tumour		5 CB	path		com-
				1.2			2 PBSC	2 unknown		pared
				months			3 fetal			to their
				to 265.2			2 unknown			historical
				months						risk of
							29 had			cancer.
							ex-vivo T			
							depletion			T cell
										deple-
										tion ap-
										peared
										to
										correlate
										with
										PTLD
										develop-
										ment.

ymphohistiocytosis; IEM: inborn error of metabolism; JAK3: Janus Kinase 3; JMML. juvenile myelomonocytic leukaemia; LAD1: leucocyte adhesion deficiency Type 1; ALL: acute lymphoblastic leukaemia; AML: acute myeloid leukaemia; BCC: basal cell carcinoma; Bu: Busulfan; CB: cord blood; CGD: chronic granulomatous disease; Cy: cyclophosphamide; EBV: Epstein-Barr virus; Flu: Fludarabine; GvHD: graft-versus-host disease; HCC: hepatocellular carcinoma; HLH: haemophagocytic MDS: myelodysplastic syndrome; MEC: mucoepidermoid carcinoma; Melp: Melphalan; Methyl: methylprednisolone; MFD: mismatched family donor; MSD: matched donor; MUD: matched unrelated donor; MMUD: mismatched unrelated donor; NA: not available; NFKB2: nuclear factor kappa B 2; PBSC: peripheral blood stem cell; PID: primary immunodeficiency; PTLD: post-transplant lymphoproliferative disease; RCC: renal cell carcinoma; SCID: severe combined immunodeficiency; SCC: squamous cell carcinoma; TBI: total body irradiation; Thio: thiotepa; Treo: treosulfan; WAS: Wiskott-Aldrich syndrome

References:

- 1. Bruton OC. Agammaglobulinemia. Pediatrics. 1952;9(6):722-8.
- Picard C, Bobby Gaspar H, Al-Herz W, Bousfiha A, Casanova JL, Chatila T, et al. International Union of Immunological Societies: 2017 Primary Immunodeficiency Diseases Committee Report on Inborn Errors of Immunity. I Clin Immunol. 2018;38(1):96-128.
- 3. Mayor PC, Eng KH, Singel KL, Abrams SI, Odunsi K, Moysich KB, et al. Cancer in primary immunodeficiency diseases: Cancer incidence in the United States Immune Deficiency Network Registry. J Allergy Clin Immunol. 2018;141(3):1028-35.
- 4. Boyle JM, Buckley RH. Population prevalence of diagnosed primary immunodeficiency diseases in the United States. J Clin Immunol. 2007;27(5):497-502.
- 5. Kobrynski L, Powell RW, Bowen S. Prevalence and morbidity of primary immunodeficiency diseases, United States 2001-2007. | Clin Immunol. 2014;34(8):954-61.
- Shillitoe B, Bangs C, Guzman D, Gennery AR, Longhurst HJ, Slatter M, et al. The United Kingdom Primary Immune Deficiency (UKPID) registry 2012 to 2017. Clin Exp Immunol. 2018;192(3):284-91.
- 7. Kirkpatrick P, Riminton S. Primary immunodeficiency diseases in Australia and New Zealand. J Clin Immunol. 2007;27(5):517-24.
- 8. Bousfiha AA, Jeddane L, Ailal F, Benhsaien I, Mahlaoui N, Casanova JL, et al. Primary immunodeficiency diseases worldwide: more common than generally thought. J Clin Immunol. 2013;33(1):1-7.
- 9. Kinlen LJ, Webster AD, Bird AG, Haile R, Peto J, Soothill JF, et al. Prospective study of cancer in patients with hypogammaglobulinaemia. Lancet. 1985;1(8423):263-6.
- 10. Jonkman-Berk BM, van den Berg JM, Ten Berge IJ, Bredius RG, Driessen GJ, Dalm VA, et al. Primary immunodeficiencies in the Netherlands: national patient data demonstrate the increased risk of malignancy. Clin Immunol. 2015;156(2):154-62.
- 11. Resnick ES, Moshier EL, Godbold JH, Cunningham-Rundles C. Morbidity and mortality in common variable immune deficiency over 4 decades. Blood. 2012;119(7):1650-7.
- 12. Mueller BU, Pizzo PA. Cancer in children with primary or secondary immunodeficiencies. J Pediatr. 1995;126(1):1-10.
- 13. Slatter MA, Gennery AR. Hematopoietic cell transplantation in primary immunodeficiency conventional and emerging indications. Expert Rev Clin Immunol. 2018;14(2):103-14.
- 14. Slatter MA, Gennery AR. Advances in hematopoietic stem cell transplantation for primary immunodeficiency. Expert Rev Clin Immunol. 2013;9(10):991-9.
- 15. Shah RM, Elfeky R, Nademi Z, Qasim W, Amrolia P, Chiesa R, et al. T-cell receptor alphabeta(+) and CD19(+) cell-depleted haploidentical and mismatched hematopoietic stem cell transplantation in primary immune deficiency. J Allergy Clin Immunol. 2018;141(4):1417-26 e1.
- 16. Lum SH, Hoenig M, Gennery AR, Slatter MA. Conditioning Regimens for Hematopoietic Cell Transplantation in Primary Immunodeficiency. Curr Allergy Asthma Rep. 2019;19(11):52.
- 17. Gennery AR, Slatter MA, Grandin L, Taupin P, Cant AJ, Veys P, et al. Transplantation of hematopoietic stem cells and long-term survival for primary immunodeficiencies in Europe: entering a new century, do we do better? J Allergy Clin Immunol. 2010;126(3):602-10 e1-11.
- 18. Griffith LM, Cowan MJ, Kohn DB, Notarangelo LD, Puck JM, Schultz KR, et al. Allogeneic hematopoietic cell transplantation for primary immune deficiency diseases: current status and critical needs. J Allergy Clin Immunol. 2008;122(6):1087-96.

- 19. Pai SY, Logan BR, Griffith LM, Buckley RH, Parrott RE, Dvorak CC, et al. Transplantation outcomes for severe combined immunodeficiency, 2000-2009. N Engl | Med. 2014;371(5):434-46.
- 20. Grunebaum E, Mazzolari E, Porta F, Dallera D, Atkinson A, Reid B, et al. Bone marrow transplantation for severe combined immune deficiency. JAMA. 2006;295(5):508-18.
- 21. Lum SH, Flood T, Hambleton S, McNaughton P, Watson H, Abinun M, et al. Two decades of excellent transplant survival for chronic granulomatous disease: a supraregional immunology transplant center report. Blood. 2019;133(23):2546-9.
- 22. Lum SH, Anderson C, McNaughton P, Engelhardt KR, MacKenzie B, Watson H, et al. Improved transplant survival and long-term disease outcome in children with MHC class II deficiency. Blood. 2020.
- 23. Ferrua F, Galimberti S, Courteille V, Slatter MA, Booth C, Moshous D, et al. Hematopoietic stem cell transplantation for CD40 ligand deficiency: Results from an EBMT/ESID-IEWP-SCETIDE-PIDTC study. J Allergy Clin Immunol. 2019;143(6):2238-53.
- 24. Socie G, Curtis RE, Deeg HJ, Sobocinski KA, Filipovich AH, Travis LB, et al. New malignant diseases after allogeneic marrow transplantation for childhood acute leukemia. J Clin Oncol. 2000;18(2):348-57.
- 25. Danner-Koptik KE, Majhail NS, Brazauskas R, Wang Z, Buchbinder D, Cahn JY, et al. Second malignancies after autologous hematopoietic cell transplantation in children. Bone Marrow Transplant. 2013;48(3):363-8.
- 26. Martin A, Schneiderman J, Helenowski IB, Morgan E, Dilley K, Danner-Koptik K, et al. Secondary malignant neoplasms after high-dose chemotherapy and autologous stem cell rescue for high-risk neuroblastoma. Pediatr Blood Cancer. 2014;61(8):1350-6.
- 27. Rizzo JD, Curtis RE, Socie G, Sobocinski KA, Gilbert E, Landgren O, et al. Solid cancers after allogeneic hematopoietic cell transplantation. Blood. 2009;113(5):1175-83.
- 28. Deeg HJ, Socie G, Schoch G, Henry-Amar M, Witherspoon RP, Devergie A, et al. Malignancies after marrow transplantation for aplastic anemia and fanconi anemia: a joint Seattle and Paris analysis of results in 700 patients. Blood. 1996;87(1):386-92.
- 29. Engels EA, Pfeiffer RM, Fraumeni JF, Jr., Kasiske BL, Israni AK, Snyder JJ, et al. Spectrum of cancer risk among US solid organ transplant recipients. JAMA. 2011;306(17):1891-901.
- 30. Filipovich AH, Mathur A, Kamat D, Shapiro RS. Primary immunodeficiencies: genetic risk factors for lymphoma. Cancer Res. 1992;52(19 Suppl):5465s-7s.
- 31. Vajdic CM, Mao L, van Leeuwen MT, Kirkpatrick P, Grulich AE, Riminton S. Are antibody deficiency disorders associated with a narrower range of cancers than other forms of immunodeficiency? Blood. 2010:116(8):1228-34.
- 32. TSB G. Lymphoproliferative disorders and malignancies related to immunodeficiencies. In: Princples and Practice of Pediatric Oncoloyg, 6th e., Pizza PA, Paplack DG (eds), Lippincott Willaims & Wilkins, Phiadelphia, PA. 2006:748.
- 33. Levine AM. Lymphoma complicating immunodeficiency disorders. Ann Oncol. 1994;5 Suppl 2:29-35.
- 34. Filipovich AH, Heinitz KJ, Robison LL, Frizzera G. The Immunodeficiency Cancer Registry. A research resource. Am J Pediatr Hematol Oncol. 1987;9(2):183-4.
- 35. Finn OJ. Immuno-oncology: understanding the function and dysfunction of the immune system in cancer. Ann Oncol. 2012;23 Suppl 8:viii6-9.
- 36. Hauck F, Voss R, Urban C, Seidel MG. Intrinsic and extrinsic causes of malignancies in patients with primary immunodeficiency disorders. J Allergy Clin Immunol. 2018;141(1):59-68 e4.

*Important review documenting causes of malignancies in patients with PID.

- 37. Verhoeven D, Stoppelenburg AJ, Meyer-Wentrup F, Boes M. Increased risk of hematologic malignancies in primary immunodeficiency disorders: opportunities for immunotherapy. Clin Immunol. 2018;190:22-31.
- 38. Witherspoon RP, Fisher LD, Schoch G, Martin P, Sullivan KM, Sanders J, et al. Secondary cancers after bone marrow transplantation for leukemia or aplastic anemia. N Engl J Med. 1989;321(12):784-9.
- 39. Bhatia S, Ramsay NK, Steinbuch M, Dusenbery KE, Shapiro RS, Weisdorf DJ, et al. Malignant neoplasms following bone marrow transplantation. Blood. 1996;87(9):3633-9.
- 40. Curtis RE, Rowlings PA, Deeg HJ, Shriner DA, Socie G, Travis LB, et al. Solid cancers after bone marrow transplantation. N Engl | Med. 1997;336(13):897-904.
- 41. Curtis RE, Travis LB, Rowlings PA, Socie G, Kingma DW, Banks PM, et al. Risk of lymphoproliferative disorders after bone marrow transplantation: a multi-institutional study. Blood. 1999;94(7):2208-16.
- 42. Kolb HJ, Socie G, Duell T, Van Lint MT, Tichelli A, Apperley JF, et al. Malignant neoplasms in long-term survivors of bone marrow transplantation. Late Effects Working Party of the European Cooperative Group for Blood and Marrow Transplantation and the European Late Effect Project Group. Ann Intern Med. 1999;131(10):738-44.
- 43. Bhatia S, Louie AD, Bhatia R, O'Donnell MR, Fung H, Kashyap A, et al. Solid cancers after bone marrow transplantation. | Clin Oncol. 2001;19(2):464-71.
- 44. Cohen A, Rovelli A, Merlo DF, van Lint MT, Lanino E, Bresters D, et al. Risk for secondary thyroid carcinoma after hematopoietic stem-cell transplantation: an EBMT Late Effects Working Party Study. J Clin Oncol. 2007;25(17):2449-54.
- 45. Landgren O, Gilbert ES, Rizzo JD, Socie G, Banks PM, Sobocinski KA, et al. Risk factors for lymphoproliferative disorders after allogeneic hematopoietic cell transplantation. Blood. 2009;113(20):4992-5001.
- 46. Omori M, Yamashita H, Shinohara A, Kurokawa M, Takita J, Hiwatari M, et al. Eleven secondary cancers after hematopoietic stem cell transplantation using a total body irradiation-based regimen in 370 consecutive pediatric and adult patients. Springerplus. 2013;2:424.
- 47. Nelson AS, Vajdic CM, Ashton LJ, Le Marsney RE, Nivison-Smith I, Wilcox L, et al. Incident cancers and late mortality in Australian children treated by allogeneic stem cell transplantation for non-malignant diseases. Pediatr Blood Cancer. 2017;64(1):197-202.
- 48. Unni MNM, Elfeky R, Rao K, Nademi Z, Chiesa R, Amrolia P, et al. Non-posttransplant lymphoproliferative disorder malignancy after hematopoietic stem cell transplantation in patients with primary immunodeficiency: UK experience. J Allergy Clin Immunol. 2018;141(6):2319-21 e1.

*Important study from the UK documenting the occurrence of malignancy post trasnplant for PID excluding lymphoproliferative disorders.

- 49. Baker KS, DeFor TE, Burns LJ, Ramsay NK, Neglia JP, Robison LL. New malignancies after blood or marrow stem-cell transplantation in children and adults: incidence and risk factors. J Clin Oncol. 2003;21(7):1352-8.
- 50. Baker KS, Leisenring WM, Goodman PJ, Ermoian RP, Flowers ME, Schoch G, et al. Total body irradiation dose and risk of subsequent neoplasms following allogeneic hematopoietic cell transplantation. Blood. 2019;133(26):2790-9.

51. Kamani NR, Kumar S, Hassebroek A, Eapen M, LeRademacher J, Casper J, et al. Malignancies after hematopoietic cell transplantation for primary immune deficiencies: a report from the Center for International Blood and Marrow Transplant Research. Biol Blood Marrow Transplant. 2011;17(12):1783-9.

** Landmark study of a large number of patients transplanted for PID who developed malignancy post transplant which were mainly lymphoproliferative disorders

- 52. Majhail NS, Brazauskas R, Rizzo JD, Sobecks RM, Wang Z, Horowitz MM, et al. Secondary solid cancers after allogeneic hematopoietic cell transplantation using busulfan-cyclophosphamide conditioning. Blood. 2011;117(1):316-22.
- 53. Curtis RE, Metayer C, Rizzo JD, Socie G, Sobocinski KA, Flowers ME, et al. Impact of chronic GVHD therapy on the development of squamous-cell cancers after hematopoietic stem-cell transplantation: an international case-control study. Blood. 2005;105(10):3802-11.
- 54. Laffort C, Le Deist F, Favre M, Caillat-Zucman S, Radford-Weiss I, Debre M, et al. Severe cutaneous papillomavirus disease after haemopoietic stem-cell transplantation in patients with severe combined immune deficiency caused by common gammac cytokine receptor subunit or JAK-3 deficiency. Lancet. 2004;363(9426):2051-4.

*Important report demonstrating that patients with common gamma chain and JAK-3 defiency are at risk of HPV associated warts even after correction of the haematopoietic stem cell lineages post transplant.

- 55. Kamili QUA, Seeborg FO, Saxena K, Nicholas SK, Banerjee PP, Angelo LS, et al. Severe cutaneous human papillomavirus infection associated with natural killer cell deficiency following stem cell transplantation for severe combined immunodeficiency. J Allergy Clin Immunol. 2014;134(6):1451-3 e1.
- 56. Goldschmidt MH, Kennedy JS, Kennedy DR, Yuan H, Holt DE, Casal ML, et al. Severe papillomavirus infection progressing to metastatic squamous cell carcinoma in bone marrow-transplanted X-linked SCID dogs. J Virol. 2006;80(13):6621-8.
- 57. Abd Hamid IJ, Slatter MA, McKendrick F, Pearce MS, Gennery AR. Long-term outcome of hematopoietic stem cell transplantation for IL2RG/JAK3 SCID: a cohort report. Blood. 2017;129(15):2198-201.

*Important report demonstarting the association of a rare skin cancer with ADA deficiency.

- 58. Kesserwan C, Sokolic R, Cowen EW, Garabedian E, Heselmeyer-Haddad K, Lee CC, et al. Multicentric dermatofibrosarcoma protuberans in patients with adenosine deaminase-deficient severe combined immune deficiency. J Allergy Clin Immunol. 2012;129(3):762-9 e1.
- 59. Carroll D, Ramani P, Lander AD. Giant-cell fibroblastoma in a patient with a bone-marrow transplant. Pediatr Surg Int. 2003;19(6):495-6.
- 60. Rubocki RJ, Parsa JR, Hershfield MS, Sanger WG, Pirruccello SJ, Santisteban I, et al. Full hematopoietic engraftment after allogeneic bone marrow transplantation without cytoreduction in a child with severe combined immunodeficiency. Blood. 2001;97(3):809-11.
- 61. Eapen M, Ahn KW, Orchard PJ, Cowan MJ, Davies SM, Fasth A, et al. Long-term survival and late deaths after hematopoietic cell transplantation for primary immunodeficiency diseases and inborn errors of metabolism. Biol Blood Marrow Transplant. 2012;18(9):1438-45.
- 62. Shimoni A, Shem-Tov N, Chetrit A, Volchek Y, Tallis E, Avigdor A, et al. Secondary malignancies after allogeneic stem-cell transplantation in the era of reduced-intensity conditioning; the incidence is not reduced. Leukemia. 2013;27(4):829-35.
- 63. Wojenski DJ, Bartoo GT, Merten JA, Dierkhising RA, Barajas MR, El-Azhary RA, et al. Voriconazole exposure and the risk of cutaneous squamous cell carcinoma in allogeneic hematopoietic stem cell transplant patients. Transpl Infect Dis. 2015;17(2):250-8.

- 64. Melnick M, Sedghizadeh PP, Allen CM, Jaskoll T. Human cytomegalovirus and mucoepidermoid carcinoma of salivary glands: cell-specific localization of active viral and oncogenic signaling proteins is confirmatory of a causal relationship. Exp Mol Pathol. 2012;92(1):118-25.
- 65. Marsh RA, Lane A, Mehta PA, Neumeier L, Jodele S, Davies SM, et al. Alemtuzumab levels impact acute GVHD, mixed chimerism, and lymphocyte recovery following alemtuzumab, fludarabine, and melphalan RIC HCT. Blood. 2016;127(4):503-12.
- Chiesa R, Standing JF, Winter R, Nademi Z, Chu J, Pinner D, et al. Proposed Therapeutic Range of Treosulfan in Reduced Toxicity Pediatric Allogeneic Hematopoietic Stem Cell Transplant Conditioning: Results From a Prospective Trial. Clin Pharmacol Ther. 2019.
- 67. Ivaturi V, Dvorak CC, Chan D, Liu T, Cowan MJ, Wahlstrom J, et al. Pharmacokinetics and Model-Based Dosing to Optimize Fludarabine Therapy in Pediatric Hematopoietic Cell Transplant Recipients. Biol Blood Marrow Transplant. 2017;23(10):1701-13.
- 68. Oostenbrink LVE, Jol-van der Zijde CM, Kielsen K, Jansen-Hoogendijk AM, Ifversen M, Muller KG, et al. Differential Elimination of Anti-Thymocyte Globulin of Fresenius and Genzyme Impacts T-Cell Reconstitution After Hematopoietic Stem Cell Transplantation. Front Immunol. 2019;10:315.
- 69. Straathof KC, Rao K, Eyrich M, Hale G, Bird P, Berrie E, et al. Haemopoietic stem-cell transplantation with antibody-based minimal-intensity conditioning: a phase 1/2 study. Lancet. 2009;374(9693):912-20.
- 70. Derderian SC, Jeanty C, Walters MC, Vichinsky E, MacKenzie TC. In utero hematopoietic cell transplantation for hemoglobinopathies. Front Pharmacol. 2014;5:278.
- 71. Agarwal R DC, Proshaska S, et al. . Toxicity-Free Hematopoietic Stem Cell Engraftment Achieved with Anti-CD117 Monoclonal Antibody Conditioning. Biol Blood Marrow Transplant 2019;25:S92.
- 72. Slatter MA, Rao K, Abd Hamid IJ, Nademi Z, Chiesa R, Elfeky R, et al. Treosulfan and Fludarabine Conditioning for Hematopoietic Stem Cell Transplantation in Children with Primary Immunodeficiency: UK Experience. Biol Blood Marrow Transplant. 2018;24(3):529-36.
- 73. Alfred A, Taylor PC, Dignan F, El-Ghariani K, Griffin J, Gennery AR, et al. The role of extracorporeal photopheresis in the management of cutaneous T-cell lymphoma, graft-versus-host disease and organ transplant rejection: a consensus statement update from the UK Photopheresis Society. Br J Haematol. 2017;177(2):287-310.
- 74. Rastogi N, Katewa S, Thakkar D, Kohli S, Nivargi S, Yadav SP. Reduced-toxicity alternate-donor stem cell transplantation with posttransplant cyclophosphamide for primary immunodeficiency disorders. Pediatr Blood Cancer. 2018;65(1).
- 75. Balashov D, Shcherbina A, Maschan M, Trakhtman P, Skvortsova Y, Shelikhova L, et al. Single-Center Experience of Unrelated and Haploidentical Stem Cell Transplantation with TCRalphabeta and CD19 Depletion in Children with Primary Immunodeficiency Syndromes. Biol Blood Marrow Transplant. 2015;21(11):1955-62.
- Neven B, Diana JS, Castelle M, Magnani A, Rosain J, Touzot F, et al. Haploidentical Hematopoietic Stem Cell Transplantation with Post-Transplant Cyclophosphamide for Primary Immunodeficiencies and Inherited Disorders in Children. Biol Blood Marrow Transplant. 2019;25(7):1363-73.
- 77. Kurzay M, Hauck F, Schmid I, Wiebking V, Eichinger A, Jung E, et al. T-cell replete haploidentical bone marrow transplantation and post-transplant cyclophosphamide for patients with inborn errors. Haematologica. 2019;104(10):e478-e82.
- 78. Touzot F, Moshous D, Creidy R, Neven B, Frange P, Cros G, et al. Faster T-cell development following gene therapy compared with haploidentical HSCT in the treatment of SCID-X1. Blood. 2015;125(23):3563-9.

- 79. Maschan M, Blagov S, Shelikhova L, Shekhovtsova Z, Balashov D, Starichkova J, et al. Low-dose donor memory T-cell infusion after TCR alpha/beta depleted unrelated and haploidentical transplantation: results of a pilot trial. Bone Marrow Transplant. 2018;53(3):264-73.
- 80. Naik S, Nicholas SK, Martinez CA, Leen AM, Hanley PJ, Gottschalk SM, et al. Adoptive immunotherapy for primary immunodeficiency disorders with virus-specific T lymphocytes. J Allergy Clin Immunol. 2016;137(5):1498-505 e1.
- 81. Ip W, Silva JMF, Gaspar H, Mitra A, Patel S, Rao K, et al. Multicenter phase 1/2 application of adenovirus-specific T cells in high-risk pediatric patients after allogeneic stem cell transplantation. Cytotherapy. 2018;20(6):830-8.

 \mathbb{C}