



2021 STATUS REPORT REGIMEN-RELATED TOXICITY AND SUPPORTIVE CARE WORKING COMMITTEE

Working Committee Leadership

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INTRODUCTION

- a. Minutes and overview plan from 2020 TCT meeting ([Attachment 1](#))

PROPOSALS MOVING FORWARD FOR SCORING (None)

Given the unique circumstances of this year, as well as a backlog of unfinished existing studies, the RRT working committee was tasked with selecting 0-2 proposals from the received batch of proposals to be considered for future discussion and consideration at the digital TCT 2021. After due consideration of all the proposals, none of the proposals could be selected to be forwarded to the next step. The proposals were deemed not to have high enough individual scientific merit to be competitive with other proposals from all the CIBMTR working committees, nor competitive with the existing portfolio of the ongoing RRTWC studies. In addition, with several proposals, there were concerns of overlap with ongoing CIBMTR or BMT CTN studies or recently published working committee publications.

PROPOSALS DROPPED BECAUSE THEY OVERLAP WITH EXISTING STUDIES OR ARE NOT FEASIBLE DUE TO LIMITATIONS OF AVAILABLE PATIENTS OR DATA

- a. PROP 2007-03 Evaluating the impact of high-dose post-transplantation cyclophosphamide (PTCy) on the risk of sinusoidal obstruction syndrome (SOS) and early mortality in B-cell acute lymphoblastic leukemia treated with inotuzumab prior to allogeneic blood or marrow transplant (Jonathan A. Webster/ Ephraim Fuchs/ George Yaghmour).
- b. PROP 2009-07 Incidence and risk factors of engraftment syndrome in autologous hematopoietic cell transplant recipients and its impact on outcomes (Muhammad Bilal Abid).
- c. PROP 2009-11 Allogeneic hematopoietic cell transplant activity and outcomes over last three decades- a CIBMTR analysis (Bhagirathbhai Dholaria/ Bipin Savani/ Mehdi Hamadani).
- d. PROP 2010-17 Predictors of increased melphalan exposure and transplant-related mortality in patients undergoing FM-based allogeneic transplant (Edmund Waller/ Karen Swiss/ Pritesh Patel).
- e. PROP 2010-62 Comparison of commonly used reduced-intensity conditioning regimens in haploidentical hematopoietic cell transplant with post-transplant cyclophosphamide (Bhagirathbhai Dholaria/ Bipin Savani).
- f. PROP 2010-68 Determining the contributing factors for and estimating the probability of developing non-infectious pulmonary toxicity in allogeneic hematopoietic cell transplant recipients (Tamila Kindwall-Keller/ Benjamin Lobo).

- g. PROP 2010-82 Organ toxicities with high dose post-transplant cyclophosphamide in haploidentical donor stem cell transplantation: Incidence, risk factors and impact on survival (Dipenkumar Modi/ Joseph Uberti/ Bipin Savani).
- h. PROP 2010-142 Trend in survival in patients undergoing allogeneic hematopoietic stem cell transplantation (Nidhi Sharma/ Yvonne Efebera).
- i. PROP 2010-162 Predicting 100 day non-relapse mortality after allogeneic hematopoietic cell transplantation using a machine learning algorithm (Akshay Sharma/ Neel S. Bhatt/ Li Tang/ Robert Davis/ Oguz Akbilgic).
- j. PROP 2010-163 Does palifermin impact the length of transplant admission? (Hemalatha Rangarajan/ Prakash Satwani).
- k. PROP 2010-175 Trends of major organ injuries amongst children and young adults following allogeneic hematopoietic cell transplantation for hematologic malignancies (Hemalatha Rangarajan/ Prakash Satwani).
- l. PROP 2010-181 Renal dysfunction prior to CD19 Chimeric antigen receptor T (CAR T) cell therapy in large B cell lymphoma (LBCL) (Christina A. Bachmeier/ Michael D. Jain/ Frederick L. Locke).
- m. PROP 2010-200 Predicting kidney outcomes post-hematopoietic cell transplant (HCT) using artificial intelligence (AI) based approaches (Sethu Madhavan/ Xia Ning, Hannah Choe).
- n. PROP 2010-269 Retrospective analysis of the association between geriatric assessment measures and post-transplant outcomes in older autologous stem cell transplant recipients (Pashna N. Munshi/ Rebecca Olin/ Andrew Artz).
- o. PROP 2010-270 A multicenter engraftment prediction model for patients undergoing allogeneic hematopoietic cell transplantation (Amany Keruakous/ Carrie Yuen).
- p. PROP 2010-307 Acute cardiovascular morbidities and mortality within 100 days after hematopoietic stem cell transplant (HSCT) (Zubair Shah).

PROPOSALS NOT ACCEPTED FOR CONSIDERATION AT THIS TIME DUE TO RELATIVE SCIENTIFIC IMPACT COMPARED TO ONGOING STUDIES AND/OR OTHER PROPOSALS

- a. PROP 2010-09 Impact of kinetic based busulfan dosing in patients receiving allogeneic hematopoietic stem cell transplant for acute myeloid leukemia (AML)/myelodysplastic syndrome (MDS) (Taylor J. Fitch/ Christopher E. Dandoy).
- b. PROP 2010-46 Incidence of and risk factors associated with venous thromboembolism after allogeneic transplant (Saurabh Chhabra/ Mehdi Hamadani).
- c. PROP 2010-51 Determining the optimal anti-thymocyte globulin dosing in patients with hematologic malignancies (Leland Metheny/ Michael Byrne/ Marcos de Lima).
- d. PROP 2010-54 Update on hematopoietic cell transplantation (HCT)-comorbidity index (CI) (Sanghee Hong/ Leland Metheny).
- e. PROP 2010-63 Clinical characteristics of engraftment syndrome and its impact on allo-HCT outcomes (Bhagirathbhai Dholaria/ Bipin Savani).

- f. PROP 2010-115 Risk factors for engraftment syndrome and its impact on clinical outcomes in pediatric allogeneic stem cell transplant recipients: A contemporary analysis (Hemalatha Rangarajan/ Prakash Satwani).
- g. PROP 2010-127 Microbial contamination of hematopoietic stem cell products and its impact on early transplant outcomes: A CIBMTR analysis (Hemalatha Rangarajan/ Prakash Satwani).
- h. PROP 2010-145 A risk-score for bronchiolitis obliterans syndrome after allogeneic hematopoietic cell transplantation (Sagar S. Patel/ Betty Hamilton/ Navneet Majhail/ Celattin Ustun).
- i. PROP 2010-225 Evaluating the impact of post-transplant cachexia on allogeneic hematopoietic stem cell transplantation outcomes (Asmita Mishra/ Joseph Pidala).
- j. PROP 2010-252 Impact of non-infectious encephalopathy (PRES) on outcomes post allogeneic hematopoietic stem cell transplant in children (Hemalatha Rangarajan/ Prakash Satwani).
- k. PROP 2010-321 Value of comorbidities assessment with HCTCI to predict post transplantation toxicities and survival in modern era (Lohith Gowda/ Mohammed Sorrow).
- l. PROP 2010-337 Is the hematopoietic cell transplantation specific comorbidity index (HCT-CI) predictive of NRM and OS in Second allogeneic hematopoietic stem cell transplants? (Ana A. Tomas/ Roni Tamari/ Miguel-Angel Perales)

STUDIES IN PROGRESS

- a. **RT17-01** Allogeneic hematopoietic stem cell transplant outcome for patients with end stage renal disease on dialysis. Status: Manuscript Preparation. The primary objectives of this study are to: 1) evaluate the impact of renal function measured by estimated glomerular filtration rate (eGFR) on allo-HCT transplant outcomes; 2) describe the characteristics and outcomes of patients on renal replacement therapy at the time of allo-HCT; and 3) explore the utilization of degrees of renal dysfunction based on eGFR to optimize the HCT-Comorbidity Index (HCT-CI). The study is in the manuscript submission phase with the goal to publication by Jan 2021.
- c. **RT18-01b** Developing a modified hematopoietic stem cell transplantation-comorbidity index for adolescents and young adults with non-malignant diseases. A modified hematopoietic cell transplantation-comorbidity index for pediatric recipients of allogeneic transplantation for non-malignant diseases. Status: Manuscript Preparation. The study aims to: 1) describe the frequency of the HCT-CI defining comorbidities and other health related biomarkers in pediatric and young adult patients and analyze the effect of each on overall survival (for non-malignant diseases) in pediatric and young adult patients; 2) create a broader risk score for pediatric and young adult patients using only weighted pre-HCT comorbidities and biomarkers that are shown to affect outcomes; 3) compare the new risk score to the standard HCT-CI in each population. The study results were presented (orally) at the 62nd Annual ASH Meeting in Dec 2020 (virtually). The study is in manuscript preparation stage with the goal to move to submit manuscript by June 2021.
- d. **RT18-02** The effect of obesity on outcomes after alternative donor stem cell transplants. Status: Analysis. The study aims to: 1) investigate the difference in non-relapse mortality risk related and overall survival to obesity (as measured by BMI, weight, and body surface area) between patients who underwent MUD or MRD transplants, haploidentical transplants, and umbilical cord transplants; 2) determine impact of BMI, weight, and body surface area on NRM, OS, RFS, engraftment rates, aGVHD and cGVHD rates after alternative stem cell transplant. The study is currently manuscript preparation with goal to move to submit manuscript by July 2021.

- e. **RT18-03** An analysis of non-infectious pulmonary toxicities in myeloablative total body irradiation vs. chemotherapy-based conditioning regimens after allogeneic hematopoietic cell transplantation for hematologic malignancies/diffuse alveolar hemorrhage is a result of complex interaction between conditioning regimen, graft source, and engraftment. Status: Manuscript Preparation. The study aims to 1) assess the incidence and risk factors of non-infectious pulmonary toxicities over time, specifically investigating conditioning regimen and intensity, and graft source; 2) evaluate the impact non-infectious pulmonary toxicity on non-relapse mortality (NRM) and overall survival (OS); 3) assess the presence of concurrent infection(s) with BOS, COP, DAH, or IPS and how this influences NRM or OS. The study is in manuscript preparation with the goal to move to submit manuscript by May 2021.
- f. **RT19-01** Analysis of comorbidity-associated toxicity at the regimen level. Status: Data File Preparation. The study aims to 1) evaluate the comorbidity-specific risk of non-relapse mortality and overall mortality within patients receiving pre-defined conditioning regimens, and 2) within patients stratified by conditioning intensity groups (myeloablative, reduced-intensity, and non-myeloablative, and 3) explore toxicities associated with specific conditioning regimen stratified by preexisting comorbidities. The study is currently in data file preparation with goal to move to manuscript preparation by July 2021.
- g. **RT19-02** Hemorrhagic cystitis as a complication of hematopoietic cell transplantation in the posttransplant cyclophosphamide graft-versus-host disease prophylaxis era compared to other allogeneic hematopoietic cell transplantation. Status: Protocol Development. The study aims to determine the incidence and severity of HC in patients who received PTCy as part of GVHD prophylaxis, 2) to describe disease characteristics and pre-transplant regimens in patients that developed HC after receiving PTCy-based GVHD prophylaxis and 3) to evaluate survival outcomes in PTCy patients with HC. The study is in protocol development with the goal to move to data file preparation by July 2021.
- h. **RT20-01** Toxicities of older adults receiving allogeneic hematopoietic cell transplant compared to younger patients. Status: Protocol Development. The study aims to 1) examine toxicities by decade in adults receiving allo transplants for hematologic malignancies and 2) examine impact of comorbidities on toxicities. The study is currently in protocol development, we hope to begin data file preparation by July 2021.

PUBLICATIONS, SUBMITTED PAPERS, PRESENTATIONS

- a. **RT14-02** Epperla N, Li A, Logan B, Fretham C, Chhabra S, Aljurf M, Chee L, Copelan E, Freytes CO, Hematti P, Lazarus HM, Litzow M, Nishihori T, Olsson RF, Prestidge T, Saber W, Wirk B, Yared JA, Loren A, Pasquini M. Incidence, risk factors for and outcomes of transplant-associated thrombotic microangiopathy. *British Journal of Haematology*. 2020 Jun 1; 189(6):1171-1181. doi:10.1111/bjh.16457. Epub 2020 Mar 2.
- b. **RT18-01a** Expanded comorbidity definitions improve applicability of the hematopoietic stem cell transplantation-comorbidity index for children, adolescents, and young adults with hematologic malignancies undergoing allogeneic stem cell transplantation. *Poster presentation at the ASH 2020 Annual Meeting*.
- c. **RT18-01b** Expanded comorbidity definitions improve application of the hematopoietic cell transplantation comorbidity index (HCT-CI) for children and young adults with non-malignant diseases receiving allogeneic hematopoietic cell transplantation. *Oral presentation at the ASH 2020 Annual Meeting*.
- d. **RT18-02** The effect of obesity on outcomes after alternative donor stem cell transplants. *Poster presentation at the TCT 2021 Annual Meeting*.

- e. **RT18-03** An analysis of non-infectious pulmonary toxicities in myeloablative total body irradiation vs. chemotherapy-based conditioning regimens after allogeneic hematopoietic cell transplantation for hematologic malignancies/diffuse alveolar hemorrhage is a result of complex interaction between conditioning regimen, graft source, and engraftment. *Poster presentation at the ASH 2020 Annual Meeting.*



A G E N D A

CIBMTR WORKING COMMITTEE FOR REGIMEN-RELATED TOXICITY AND SUPPORTIVE CARE

Orlando, FL

Saturday February 22, 2020, 12:15 – 2:15 pm

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1. Introduction

Dr. Stadtmauer opened the meeting at 12:15 pm by welcoming the working committee members for attending the Regimen-Related Toxicity and Supportive Care Working Committee (RRTWC) meeting. He introduced the RRTWC leadership, welcomed the incoming chair Dr. Mohamed Sorror and thanked Dr. Shin Mineishi for his contributions to the committee over the year. Dr. Stadtmauer then stated the goals and limitations of the RRTWC and introduced Dr. Chhabra to the podium to continue.

2. Accrual summary

The accrual summary was not presented in order to provide more time for the discussion of RT studies that are ongoing, published or presented in the last year, and the proposed studies to be presented at the meeting.

3. Presentations, published or submitted papers

Dr. Chhabra gave an overview of the studies published and submitted in the past year. He also presented new areas of data collection on both the TED and CRF forms, rules of authorship, goals, expectations and limitations. Dr. Chhabra brought up the importance of contributing at each step of the study.

- a. **RT13-02** Sabloff M, Chhabra S, Wang T, Fretham C, Kekre N, Abraham A, Adekola K, Auletta JJ, Barker C, Beitinjaneh AM, Bredeson C, Cahn J-Y, Diaz MA, Freytes C, Gale RP, Ganguly S, Gergis U, Guinan E, Hamilton B, Hashmi S, Hematti P, Hildebrandt G, Holmberg L, Hong S, Lazarus HM, Martino R, Muffly L, Nishihori T, Perales M-A, Yared J, Mineishi S, Stadtmauer EA, Pasquini MC, Loren AW. *Comparison of high doses of total body irradiation in myeloablative conditioning*

prior to hematopoietic cell transplantation. **Biology of Blood and Marrow Transplantation: Journal of the American Society for Blood and Marrow Transplantation.**

doi:10.1016/j.bbmt.2019.08.012. Epub 2019 Aug 29.

- b. **RT14-01** Parikh SH, Satwani P, Ahn KW, Sahr NA, Fretham C, Abraham AA, Agrawal V, Auletta JJ, Abdel-Azim H, Copelan E, Diaz M-A, Dvorak CC, Frangoul HA, Freytes CO, Gadalla SM, Gale RP, George B, Gergis U, Hashmi S, Hematti P, Hildebrandt GC, Keating AK, Lazarus HM, Myers K, Olsson RF, Prestidge T, Rotz S, Savani BN, Shereck E, Williams KM, Wirk B, Pasquini MC, Loren AW. *Survival trends in infants undergoing allogeneic hematopoietic cell transplant.* **Journal of the American Medical Association Pediatrics.** doi:10.1001/jamapediatrics.2019.0081. Epub 2019 Mar 18. PMC6503511.
- c. **RT14-02** Epperla N, Li A, Logan B, Fretham C, Chhabra S, Aljurf M, Chee L, Copelan E, Freytes CO, Hematti P, Lazarus HM, Litzow M, Nishihori T, Olsson RF, Prestidge T, Saber W, Wirk B, Yared JA, Loren A, Pasquini M. *Incidence, Risk Factors for and Outcomes of Transplant-Associated Thrombotic Microangiopathy.* **British Journal of Haematology.** In press.
- d. **RT14-03** Zinter MS, Logan BR, Fretham C, Sapru A, Abraham A, Aljurf MD, Arnold SD, Artz A, Auletta JJ, Chhabra S, Copelan E, Duncan C, Gale RP, Guinan E, Hematti P, Keating AK, Marks DI, Savani BN, Olsson R, Ustun C, Williams KM, Pasquini MC, Dvorak CC. *Comprehensive prognostication in critically ill pediatric hematopoietic cell transplant patients: Results from merging the Center for International Blood and Marrow Transplant Research (CIBMTR) and Virtual Pediatric Systems (VPS) Registries.* **Biology of Blood and Marrow Transplantation.** doi:10.1016/j.bbmt.2019.09.027. Epub 2019 Sep 26.
- e. **RT17-01a** Dias A, Farhadfar N, Wang T, Fretham C, Murthy H, Logan B, Mineishi S, Savani B, Stadtmauer E, Chhabra S, Wingard JR, Ganguly S, Pasquini MC. *Impact of Renal Dysfunction Measured By Estimated Glomerular Filtration Rate (eGFR) on Outcomes after Allogeneic Hematopoietic Cell Transplantation (HCT).* **Presented at ASH, December 2019.**
- f. **RT17-01b** Farhadfar N, Murthy H, Wang T, Fretham C, Dias A, Logan B, Mineishi S, Savani B, Stadtmauer E, Chhabra S, Ganguly S, Wingard JR, Pasquini MC. *Development of the Renal Adjusted Hematopoietic Cell Transplant Comorbidity Index (RA-HCT-CI) Using Different Levels of Renal Dysfunction According to Estimated Glomerular Filtration Rate (eGFR).* **Presented at TCT, February 2020.**
- g. **RT18-04** Broglie L, Fretham C, Al-Seraihy, George B, Kurtzberg J, Loren A, MacMillan M, Martinez C, Davies SM, Pasquini M. *Pulmonary complications in pediatric and adolescent patients following allogeneic hematopoietic cell transplantation.* **Biology of Blood and Marrow Transplantation: Journal of the American Society for Blood and Marrow Transplantation.** doi:10.1016/j.bbmt.2019.06.004. Epub 2019 Jun 12.

4. Studies in progress

Dr. Chhabra presented the studies in progress. The older studies will be given priority to finish this year. The rest of the studies are on schedule to meet their current goals.

- a. **RT17-01** Allogeneic hematopoietic stem cell transplant outcome of patients with end stage renal disease on dialysis (N Farhadfar/A Dias/JR Wingard/H Murthy/S Ganguly) **Manuscript preparation**
- b. **RT18-01** A Modified Hematopoietic Cell Transplantation (HCT) Risk Assessment Tool for Pediatric and Young Adult Patients Undergoing Allogeneic Transplantation. (B Friend/L Broglie/G Schiller/M Thakar/M Sorrow) **Analysis**
- c. **RT18-02** The effect of obesity on outcomes after alternative donor stem cell transplants (M Abou-Ismaïl/G Ravi/L Metheny/M de Lima) **Datafile preparation**
- d. **RT18-03** An Analysis of Non-Infectious Pulmonary Toxicities in Total Body Irradiation versus Chemotherapy-Based Conditioning Regimens after Allogeneic Hematopoietic Cell

Transplantation for Hematologic Malignancies (S Patel/B Hamilton/N Majhail/C Ustun) **Datafile preparation**

- e. **RT19-01** Analysis of comorbidity-associated toxicity at a regimen-based level (R Shouval/ B Savani/ A Nagler) **Protocol development**
- f. **RT19-02** Hemorrhagic cystitis as a complication of hematopoietic stem cell transplantation in the post-transplant cyclophosphamide graft-versus-host disease prophylaxis era compared to other allogeneic stem cell transplants (K Adekola/ N Ali/ O Frankfurt/ L Metheny/ J Moreira/ M de Lima) **Protocol development**

5. Proposals

Future/proposed studies

Dr. Chhabra provided information on the proposals received this year. There were 23 proposals for the RRTWC. Seventeen were dropped due to overlap with current or recent studies or feasibility issues, which left six proposals to be presented at the meeting. The proposal themes were seropositivity (CMV, HTLV), toxicities, prior cancer diagnosis, post-transplant cyclophosphamide and prognostic tools for transplant. Dr. Chhabra invited Dr. Mineishi to the podium.

- a. **PROP 1911-167** CMV Serotype and Graft Failure (M Pamukcuoglu/ M Arora) (Attachment 4)
Dr. Mineishi introduced Dr. Pamukcuoglu. The hypothesis is that graft failure may occur from dominant recipient CMV positive NK/T cells as opposed to donor CMV positive NK/T cells. The aim is to determine the impact of pre-transplant recipient/donor CMV serotype on graft failure. The CIBMTR identified 9592 cases of first allo with peripheral blood grafts between 2008 and 2018 for all hematologic malignancies and conditioning intensities.
The primary aim of the study is to determine the impact of pre-transplant donor/recipient CMV status on graft failure.
There was discussion around the definition of graft failure, and the variability of data collected. There was discussion around Letemovir for prevention of CMV reactivation and loss of grafts. Exploring CMV as an infection post-transplant. Letemovir is not approved in pediatric patients, differs by center and peripheral blood. Since primary graft failure incidence is <5%, might be important to include bone marrow and non-malignant cases to increase numbers. Non-malignant cases have higher graft failure. Olsson et al. published study on graft failure with pediatric cases included and found CMV serostatus was not a significant predictor for graft failure based on patient data between 1995 and 2008.
- b. **PROP 1911-181** Significance of a prior cancer diagnosis as exclusion criteria in clinical trials among patients with Hematological malignancies undergoing hematopoietic stem cell transplant (HSCT) (A Kansagra/ S Hashmi/ B Savani/ M Hamadani/ S Devine) (Attachment 5)
Dr. Mineishi introduced Dr. Anderson. The hypothesis of this study is there is no difference in progression-free survival between patients on clinical trials where there are guidelines or acceptance of cases with prior cancers and standard of care patients.
The CIBMTR found 1000 cases of adults with AML or MDS transplanted between 6/2/2011 and 4/18/2014 with similar criteria to the BMT-CTN 0901 clinical trial cohort.
The primary aim of the study is to evaluate overall and progression-free survival for both cohorts. The secondary aim is to evaluate the risk of secondary malignancy after transplant. Dr. Pasquini commented that there does not appear to be a strict exclusion of prior cancers in the BMT CTN 0901 cohort and the type of prior malignancy may matter. Curative cancer treatment is the endpoint with competing outcomes. Dr. Anderson said within a population of transplants in-situ cancers and melanoma may make a difference. There was a comment that this might not be the right clinical trial to compare it to. The impact of cancer on outcomes is important to study, and the FDA is interested in reducing barriers to clinical trials. Dr. Stadtmauer questioned how the population was chosen. Dr. Pasquini commented that the

comparison does not have to solely be to CTN 0901 but to a combination of clinical trials potentially.

- c. **PROP 1911-198** Outcomes and safety of hematopoietic stem cell transplants from HTLV positive donors and HTLV positive recipients (M Janakiram/ G Okov) (Attachment 6)

Dr. Savani introduced Dr. Janakiram. The hypotheses of this proposal are as follows; 1) patients who receive transplants for HTLV-1 positive donors will have the same outcomes as transplants from HTLV-1 negative donors, 2) the rate of secondary malignancies will be similar between groups, 3) the rate of secondary malignancies will be similar in HTLV-1 positive recipients for both auto and all transplants, 4) the progression-free survival and non-relapse mortality will be comparable between cohorts.

The CIBMTR identified 31,919 cases who received their first allo transplant for any donor or graft source between 2008 and 2018 (26,839 HTLV-1 negative recipients, 217 HTLV-1 positive recipients, and 4863 unknown HTLV-1 seropositivity). Additionally, there were 13,577 cases of auto transplants with any graft source between 2008 and 2018.

The primary aims are to examine progression-free survival, non-relapse mortality, and secondary malignancies. The secondary aims are to examine overall survival and incidence of HTLV-1 related complications.

There was discussion around the impact of this study and partnership with other registries. There was a comment about feasibility of comparing 217 HTLV-1 positive recipients to 2500 HTLV-1 negative cases. Dr. Janakiram responded that a matching pair analysis may assist with this issue. Dr. Yared from University of Maryland raised the question that the HTLV-1 positive donors in the allo transplants may be related donors. There was a question as to how long HTLV associated cancers develop. Dr. Janakiram stated these cancers take around 30 years to develop. There was discussion that the endpoint might need to include the 30 year follow-up and second primary malignancies. There was discussion regarding reaching out to centers to verify HTLV status. There was a comment that donors may be excluded on HTLV status, if this is not the case, donors increase and there may be a need for a different end point.

- d. **PROP 1911-234** Patterns of Venous Occlusive Disease in Patients with AML and ALL in the era of Monoclonal Antibodies and Antibody Drug Conjugates (L Gowda/ M Byrne/ P Kebriaei/ D Porter) (Attachment 7)

Dr. Savani introduced Dr. Kebriaei. The hypothesis is that the risk of VOD has changed with new drugs for treatment of acute leukemias.

The CIBMTR identified 664 cases who underwent allo transplant and received monoclonal antibodies prior to transplant for AML or ALL between 2008-2019.

The primary aim is to evaluate the incidence and severity of VOD of patients treated pre-transplant with gemtuzamab or inotuzamab. The secondary aims are to examine the timing of VOD, mortality and morbidity for monoclonal antibodies, risk factors for VOD, predictors of survival for VOD, impact of antibodies on VOD risk, severity and death, relapse, non-relapse mortality and acute and chronic GVHD for those who developed VOD.

There was a comment that gemtuzamab does not lead to increased risk for VOD. There was a comment that the granularity of the CIBMTR data may be insufficient and that the dosing information and schedule would be important and those have changed over time but are not available in the registry. There was discussion around the changes in gemtuzamab use and the availability of date of use and resolution of VOD. There was discussion regarding the methodology of this study given not all the data are applicable to all cases. There was discussion around the availability of ICU stays and defibrotide information which are not captured during the entire study period. There was a comment that this study might have low statistical power. It was commented that this would be a better study than what has been published so far.

- f. **PROP 1911-46** The PTCY-CI and PTCY-CDRI: prognostic tools for the use of post-transplant cyclophosphamide based GVHD prophylaxis regimens in allogeneic stem cell transplants for malignant conditions (R Shapiro/ R Romee/ A Bashey) (Attachment 8)
Dr. Stadtmauer introduced Dr. Shapiro. The hypotheses of this study is PT-Cy impacts non-relapse mortality and should be included in HCT-CI and DRI scores to accurately predict risk scores.
The CIBMTR identified 8353 cases of adults who received PT-Cy for the first allo transplant for a hematologic malignancy.
The primary aim is to validate the HCT-CI and DRI risk score using PT-Cy prophylaxis regimens. The secondary aims are to develop novel HCT-CI and DRI risk scores which incorporate transplant related factors including PT-Cy use.
Dr. Stadtmauer commented that most of the cases in this proposal are haploidentical. There was discussion that PT-Cy has already been tested in HCT-CI and DRI. Dr. Shapiro commented that the importance of looking at PT-Cy in these risk scores is there will be lower non-relapse mortality with use which may impact scoring systems. Dr. Pasquini commented that HCT-CI has been looked at in PT-Cy setting and it has been validated as a predictor for NRM. There was discussion that the intensity of conditioning may be important to HCT-CI scores where lower intensity related to increased scores. Dr. Sorrow commented that reclassifying risk with PT-Cy compared to other immune suppression and different setting does not produce different scores or better predictive models. There was a comment that with a new score it would increase the number of scores physicians need to consider when treating patients.
- g. **PROP 1911-60** Toxicities of Older Adults Receiving Allogeneic Hematopoietic Cell Transplant Compared to Younger Patients (R Jayani/ H Muff) (Attachment 9)
Dr. Stadtmauer introduced Dr. Jayani. The hypothesis of the study is that older adults experience a higher rate of toxicities than younger patients.
The CIBMTR identified 24,581 cases of adults who underwent allo transplant between 2008 and 2018 (15,794 <60 years old, 8,787 60+ years old).
The primary aim of the study is to understand the toxicities that most impact older adults. Secondary aims include evaluating the frequency of organ toxicity early after transplant, late effects of transplant in both groups, impact of comorbidities on organ toxicity and late effects and acute and chronic GVHD rates and severities.
There was discussion of the cut point for age, 70+ years might be more standard with CIBMTR studies, age could also be evaluated continuously or by decade. There was a comment that continuous age would allow model flexibility. There was discussion around quantifying toxicities as individual toxicities and as a sum score. There was a comment that older people are predisposed to more toxicities and important to modify regimens impact potential toxicities. Dr. Pasquini commented that is a different endpoint looking at all the toxicities and a breakdown by regimen. There was a comment that ICU information, HCT-CI and pulmonary toxicities are important. Dr. Sorrow commented that groups with increased toxicity of management might be the same as younger. Dr. Stadtmauer mentioned this might be a complimentary study to the BMT-CTN study looking at non-relapse mortality and toxicity with a composite endpoint and examining regimen related tool.

The meeting was adjourned at 2:15 pm.

Working committee Overview Plan for 2020-2021							
Study number and title	Current status	Goal with date	Total hours to complete	Total hours to 2021 goal	Hours allocated to 6/30/20	Hours allocated 7/1/20-6/30/21	Total Hours allocated
RT17-01: AlloHCT outcome of patients with end stage renal disease on dialysis.	Manuscript prep	Submit by April 2020 Published by July 2021	60	60	50	10	60
RT18-01: A modified HCT risk assessment tool for pediatric and young adult patients undergoing alloHCT.	Analysis	Submit by May 2020 Published by July 2021	110	120	110	10	120
RT18-02: The effect of obesity on outcomes after alternative donor stem cell transplants.	Data file prep	Manuscript preparation by May 2020 Submitted by July 2021	200	200	130	70	200
RT18-03: An analysis of non-infectious pulmonary toxicities in regards to conditioning regimens, graft source and early vs. delayed engraftment.	Data file prep	Manuscript preparation by May 2020 Submitted by July 2021	160	160	90	70	160
RT19-01: Analysis of comorbidity-associated toxicity at a regimen based level.	Data file preparation	Analysis by July 2020 Submitted by July 2021	290	290	180	110	290
RT19-02: Hemorrhagic cystitis as a complication of HCT in the Pt-Cy GVHD prophylaxis era compared to other alloHCTs.	Protocol development	Data file preparation by July 2020 Manuscript preparation by July 2021	330	260	100	160	260

RT20-01: VOD in AML and ALL with pre-transplant monoclonal antibodies	Protocol pending	Protocol Development by July 2020 Manuscript preparation by July 2021	330	260	0	260	260
RT20-02: Toxicities of older adults receiving allogeneic hematopoietic cell transplant compared to younger patients	Protocol pending	Protocol Development by July 2020 Analysis by July 2021	330	200	0	200	200

Oversight Assignments for Working Committee Leadership (March 2020)	
Edward Stadtmauer	<p>RT18-02: The effect of obesity on outcomes after alternative donor stem cell transplants.</p> <p>RT20-01: VOD in AML and ALL with pre-transplant monoclonal antibodies</p>
Bipin Savani	<p>RT18-03: An analysis of non-infectious pulmonary toxicities in regards to conditioning regimens, graft source and early vs. delayed engraftment.</p> <p>RT19-01: Analysis of comorbidity-associated toxicity at a regimen based level.</p> <p>RT19-02: Hemorrhagic cystitis as a complication of HCT in the Pt-Cy GVHD prophylaxis era compared to other alloHCTs.</p>
Mohamed Sorrow	<p>RT18-01: A modified HCT risk assessment tool for pediatric and young adult patients undergoing alloHCT.</p> <p>RT20-02: Toxicities of older adults receiving allogeneic hematopoietic cell transplant compared to younger patients</p>