

Methodology Employed for Annual Report on Hematopoietic Cell Transplant Center-Specific Survival Rates Last Updated December 14, 2020

Table of Contents Rationale for a fixed effects censored data logistic regression model......4 Details of fixed effects censored data logistic regression and confidence limits......5 ١. Patient demographics......7 Risk factors included in final multivariate model7

CIBMTR* is a research collaboration between the National Marrow Donor Program* (NMDP)/ Be The Match* and the Medical College of Wisconsin Milwaukee Campus

Medical College of Wisconsin 9200 W Wisconsin Ave, Suite C5500 Milwaukee, WI 53226 USA (414) 805-0700 Minneapolis Campus

National Marrow Donor Program/ Be The Match 500 N 5th St Minneapolis, MN 55401 USA (763) 406-5800

Introduction

The purpose of the annual report on transplant center-specific survival rates is to provide potential hematopoietic cell transplant (HCT) recipients, their families, and the general public with a comparison of survival rates among the centers in the C.W. Bill Young Cell Transplantation Program (CWBYCTP) network. Transplant centers may use these reports for quality improvement initiatives. Reporting center-specific survival rates is a requirement of the Stem Cell Therapeutic and Research Act of 2005 (re-authorized in 2010 and 2015), and prior to that, the 1990 Transplant Amendments Act. Because centers vary considerably in the risk level of cases treated, a statistical model was developed to adjust for several risk factors known or suspected to influence outcome. The outcome reported is one-year overall survival, for recipients of allogeneic HCT in the United States only. No attempts are made to incorporate other outcomes, such as relapse or disease-free survival.

The first center-specific risk-adjusted comparisons were published in 1994 [1] and yearly since then. The current iteration of the report prepared by the Center for International Blood and Marrow Transplant Research (CIBMTR) includes recipients of both unrelated and related donor transplants facilitated by the CWBYCTP for a three-year time window. The methodology for this analysis has undergone various transformations over the years. The methodology in current use has been employed since 2005, thus allowing direct comparisons over the most recent time periods. This method adjusts for risk using a censored data logistic regression model [2] [3] [4] that allows inclusion of recipients with incomplete one-year follow-up. Note that although the method has remained the same, the types of patients studied changed with the inclusion of related-donor transplants in the 2010 report, which may affect comparisons over time. A riskadjusted one-year survival rate is calculated for each center, based on results of the censored data logistic regression.

Results are available via the CWBYCTP website

(<u>http://bloodcell.transplant.hrsa.gov/research/transplant_data/us_tx_data/index.html</u>), and a version of this report, as approved by HRSA, is distributed to HCT centers. This information is available online at <u>www.bethematch.org/tcdirectory/search</u>. Raw numbers of transplants and surviving recipients are published for each center, stratified by diagnosis and age. Each center included in the report performed at least one unrelated or related donor transplant over the three-year window of time for analysis.

Methods

Recipients and data

The current analysis includes first unrelated or related donor transplants performed in a threeyear time interval, with follow-up through one year after the last recipient was transplanted. The rolling three-year window of transplants for inclusion was adopted with the 2011 report, replacing a rolling five-year window used previously. This change was based on the recommendation of the 2010 Center-Specific Outcomes Analysis Forum [5], in order to represent more current transplant center outcomes. A minimum of one-year follow-up is required for all eligible cases. All U.S. transplant centers that performed at least one HCT in the time interval are considered for inclusion in the report, provided they had sufficient data with at least one year of follow-up available. Typically, about 180 U.S. transplant centers are included in the analysis, with about 25,000 first allogeneic transplants performed by domestic transplant centers in the CWBYCTP network during this time.

Demographics of the included cases are provided in tables for recipients of unrelated donor transplants and recipients of related donor transplants, broken down by donor type according to unrelated vs. baseline and follow-up data used for the analysis are provided to the CIBMTR by the transplant centers at the time of transplant (baseline), and at 100 days, six months and annually post-transplant, using standardized forms. Race was self-reported by recipients or by the staff at the center.

Risk factors considered

Based on the recommendation of the Center-Specific Outcomes Analysis Forum in 2010 and 2012, variables recognized as clinically important were forced into the model regardless of whether they were statistically significant. After careful discussion with clinical and statistical transplant experts, the following essential risk factors were included in the model:

- Recipient age
- Recipient race (self-reported)
- Recipient Karnofsky / Lansky Performance Status score at transplant
- Coexisting disease (HCT-specific comorbidity index (HCT-CI), Sorror [6])
- Low body mass index (BMI) of adults age 18 or older, or low BMI or weight-for-age percentile for age<18
- Recipient cytomegalovirus (CMV) serology
- History of mechanical ventilation
- History of invasive fungal infection
- Prior autologous transplant
- Diagnosis and disease status / stage
- AML ELN risk group (Döhner et al. [7])
- AML transformed from Myelodysplastic (MDS) / myeloproliferative (MPN) diseases
- Number of induction cycles to achieve latest CR before HCT for AML and ALL patients in CR
- Time from diagnosis to transplant for acute myelogenous leukemia (AML) and acute lymphoblastic leukemia (ALL) not in CR1 or PIF (used as surrogate for length of CR)
- ALL cytogenetic risk group (Moorman et al. [8])
- MDS IPSS-R prognostic risk category/score at HCT (Greenberg et al. [9])
- MDS with predisposing conditions
- Resistant disease in non-Hodgkin lymphoma (NHL) and Hodgkin lymphoma (HL) only
- Plasma cell disorder disease status at HCT
- Year of transplant
- Donor type: matched sibling donor vs. other related vs. unrelated donor

- HLA matching^{*}
- Recipient and donor gender
- Donor age (unrelated bone marrow or peripheral blood stem cells (PBSC) donors only)
- Socioeconomic status (median household income) based on zip code of residence of recipient

In addition, the following variables were believed to be of uncertain clinical relevance, and so they were included in the model only if statistically significant (p<0.05).

- Recipient ethnicity
- Therapy related AML or MDS
- ALL molecular marker BCR/ABL at any time between diagnosis and HCT
- T-cell lineage in ALL, Philadelphia chromosome in ALL
- Del 17p in CLL
- NHL subtype
- Multiple myeloma cytogenetics risk group (Palumbo et al. [10])
- Multiple myeloma International Staging System (ISS) stage at diagnosis
- Unrelated BM or PBSC donor ethnicity
- Unrelated BM or PBSC donor race
- BM or PBSC donor CMV serology
- BM or PBSC donor parity

Statistical Analysis

Rationale for a fixed effects censored data logistic regression model

One of the CIBMTR's goals for the transplant center-specific outcomes analysis is to calculate a fair and accurate predicted survival rate given a center's recipient case mix. To do this, a fixed effects censored data logistic regression model is used. The fixed-effects logistic regression model provides information about how the recipients actually treated in a particular center would have fared had they been transplanted at a "generic" transplant center within the CWBYCTP. This model assumes *no center effect*. In other words, it assumes that recipients are dying at the same uniform rate across all CWBYCTP transplant centers, after adjusting for covariates. The model also adequately accounts for recipients with incomplete follow-up at one year.

Every effort is made to update follow-up information on each recipient. Some recipients are indeed lost to follow-up, and their final survival status at one year is unknown. To address this

^{*} For PBSC and marrow transplants, high-resolution typing at HLA-A, -B, -C, and –DRB1 was used for the cases where it was available. For the remaining patients with PBSC and bone marrow graft sources, the best available matching information at HLA-A, -B, -C, and -DRB1 was used (Weisdorf et al. [12]). For single cord blood transplants, high-resolution typing at HLA-A, -B, -C, and –DRB1 was used for the cases where it was available. For the remaining patients with single cord blood graft source, low/intermediate-resolution typing at HLA-A and -B, and highresolution typing at -DRB1 only were considered. For multiple cord blood transplants, low/intermediate-resolution typing at HLA-A and -B, and high-resolution typing at -DRB1 only were considered. The match grade of the worstmatched unit was analyzed.

problem, the analysis only includes centers that demonstrated 90% completeness of follow-up, meaning that the one-year status was known for at least 90% of their transplanted recipients. However, there are still some recipients for whom survival status at one year is incomplete, although many recipients had follow-up done just prior to one year. If these recipients are excluded from the center-specific analysis, it may bias the survival estimates. A censored data version of logistic regression based on pseudo-values proposed by Andersen et al. [2], Klein and Andersen [3], and Klein et al. [4] addresses this issue. This method is a generalization of logistic regression that simplifies to logistic regression technique is used to estimate the fixed effects and predict the recipients' survival probabilities based on their patient characteristics alone. These predicted survival probabilities are then used to construct confidence limits for a center's survival probability according to the characteristics of the patients transplanted at that center. The actual survival observed at that center can be compared to these intervals to assess the performance of the center. This method is described in more detail below.

Details of fixed effects censored data logistic regression and confidence limits

Modeling for the center-specific survival analysis can be broken down into four steps, as outlined below.

I. Definition of pseudo-values

To compute the pseudo-value for recipient *i*, first compute the pooled sample Kaplan-Meier estimate of survival at one year based on the entire sample, $\hat{S}_p(1)$. Next compute the Kaplan-Meier estimate of survival at one year based on the entire dataset with observation *i* removed $\hat{S}_p^{(i)}(1)$. The *i*th pseudo-value is defined by $\hat{\theta}_i = n\hat{S}_p(1) - (n-1)\hat{S}_p^{(i)}(1)$. If there is no censoring, then the *i*th pseudo-value is simply the indicator that the *i*th recipient was alive at one year. These pseudo-values will then be used in a regression model using a logit link, similar to a standard logistic regression model, as described in the next section. The parameters of the regression model can be estimated using generalized estimating equations (GEE), which are implemented in PROC GENMOD in SAS.

II. Model building

Let $(Z_{i1},...,Z_{ip})$ denote the set of covariates in the final model for recipient i. First fit a fixed effects censored data logistic regression model with no center effect,

$$\varphi_i = \ln \frac{\theta_i}{1 - \theta_i} = \beta_0 + \sum_{l=1}^p \beta_l Z_{il} \ .$$

III. Predicted and observed survival

From the fitted logistic regression model, each recipient has an estimated survival rate

$$\hat{p}_i = \frac{\exp(\hat{\varphi}_i)}{1 + \exp(\hat{\varphi}_i)}$$

based on his or her risk characteristics. The predicted survival rate at center j based on recipient characteristics $E(S_j)$ is the average of the estimated survival rates for all recipients at center j,

$$E(S_j) = \left(\sum_{i \in C_j} \hat{p}_i\right) * \frac{1}{n_j}.$$

The observed one-year survival rate at center *j* can be computed using the Kaplan-Meier estimate of survival using the recipients at center *j*. This simplifies to the sample proportion of recipients alive when there is no censoring prior to one year present.

IV. Confidence Limits

Confidence limits are generated using a bootstrapping methodology. However, the bootstrap technique was modified slightly from previous years' reports to improve the coverage probabilities of the intervals, as described in Logan et al. [11] Previously, binary outcomes were generated for each individual to simulate the confidence limits; however, a more accurate prediction interval that controls the type I error rate can be obtained by resampling the residuals from the general linear model instead. Define the scaled Pearson residual for patient *i* by

$$r_i = \frac{\hat{\theta}_i - \hat{p}_i}{\sqrt{\hat{p}_i(1 - \hat{p}_i)}},$$

then the bootstrap re-sampling algorithm to generate a prediction interval for center j is as follows. For b=1 to 10,000:

- 1. Generate r_i^{*b} for patient *i* by sampling with replacement from the set of residuals $\{r_i, i = 1, ..., n\}$
- 2. Compute the bootstrap predicted value for patient *i* as

$$Y_{i}^{*b} = \hat{p}_{i} + r_{i}^{*b} \sqrt{\hat{p}_{i}(1-\hat{p}_{i})}$$

3. Compute the predicted center outcome for center *j* as

$$S_j^{*b} = \frac{1}{n_j} \sum_{i \in C_j} Y_i^{*b}$$

Then the 95% predicted confidence bounds for survival at center *j* are obtained by taking the 2.5th and 97.5th percentile of S_i^{*b} across the 10,000 bootstrap samples.

This confidence interval refers to the survival rate that might be observed at that center if there were no center effect and those recipients had been transplanted at any center in the network. The observed survival rate can be compared with this confidence interval to see if there is evidence of the center over-performing or under-performing the overall network.

Handling missing data for risk factors

Occasionally, data are not available for significant characteristics for subjects as reported by the centers. If there were sufficient numbers of such subjects, they were included in the multivariate modeling as a distinct category of the covariate. However, when the number of subjects with data not available for a variable is too small (generally less than 20 subjects) to fit the model as its own category, those subjects were imputed to the relevant highest frequency

category within the variable for categorical variables, and to the median value for ordinal or numeric variables (e.g. HCT-CI).

Results

Patient demographics

Demographics of the patient population are provided in the report, displayed by donor type according to unrelated vs. related donor.

Risk factors included in final multivariate model

The results of the multivariate model are presented in a set of tables where each variable and its associated odds ratio are described, along with 95% confidence limits. Factors included in the final model are:

- Recipient age
- Recipient race**
- Karnofsky / Lansky score at transplant
- Sorror HCT-CI
- Adult BMI group
- Pediatric BMI group**
- Recipient CMV status
- History of mechanical ventilation
- History of invasive fungal infection
- Prior autologous transplant**
- Disease and disease status/stage
- AML ELN risk group
- AML transformed from MDS/MPN
- Number of induction cycles for AML and in CR1**
- Time from diagnosis to transplant for AML and ALL in CR2 and CR3+/Relapse (used as surrogate for length of CR)
- ALL cytogenetic risk group**
- Philadelphia chromosome in ALL patients
- Number of induction cycles for ALL and in CR1**
- MDS IPSS-R risk score at HCT
- MDS therapy related
- MDS with predisposing condition**
- CLL and other chronic leukemia stage
- NHL subtype
- Sensitivity to chemotherapy in NHL
- Sensitivity to chemotherapy in HL
- Plasma cell disorder disease status

^{**} Maintained in the model due to clinical relevance but not statistically significant

- Year of transplant
- HLA matching by donor and graft type
- Donor/Recipient sex match (bone marrow or PBSC only)
- Donor age (unrelated bone marrow or PBSC donors only)
- Recipient median household income based on zip code

The model is similar to last year, except for the following:

- Recipient ethnicity and ALL T-cell lineage were not included in the model due to lack of statistical significance.
- Therapy related MDS was included in the model this year due to meeting the significance requirement.
- Donor age was broken down into 5-year increments rather than decades.
- Two new variables were added, BMI groups separately for adults and pediatric patients. These identified those underweight to test as a factor in the model. It was significant among the adult group only.

The Beta_0 intercept term for the model is made available in the formal report.

Center-specific results

Final center-specific results are presented, along with centers' historical performance in tables, and on the public website. Numbers of transplanted recipients at each center, actual (observed) survival at one year, predicted survival at one year, 95% confidence intervals for predicted survival, and performance status are displayed for each center. Centers whose actual survival is outside the 95% confidence limits for predicted survival have a "-1" in the performance status column if performing below the confidence limit. Centers with a "0" in the performance status column are performing as predicted. Most centers performed as predicted with respect to overall performance in previous years. Since the censored data logistic regression model assumes no center effect, centers with smaller numbers of transplants (e.g. N = 1 or 2) will *not* have their predicted survival proportion regress toward the network average. Rather, the confidence limits around the predicted survival at that center will simply be much wider than those of larger centers.

Results are also displayed for centers via a visual box-plot graphic. Centers are arranged by center number, while reading from left to right across these figures. The actual survival at each center is superimposed with each box plot (using the symbol '•') to give the reader an instantaneous picture of how close to under- or over-performing the center was.

Patients can find information about all U.S. transplant centers performing allogeneic transplants in the online U.S. Transplant Center Directory on <u>http://bethematch.org</u>. Listings are organized by state and can be found at <u>bethematch.org/tcdirectory/search</u>. Along with center outcomes, each listing includes a description of that center's program, contact information, the number of transplants performed over a specified time period and survival

statistics by patient's age, disease type and stage for both related and unrelated donor transplants. A link to the Transplant Center Directory can also be found on the Health Resources and Services Administration (HRSA) https://bloodstemcell.hrsa.gov/data/donation-and-transplantation-statistics website.

Because the outcome of interest is one-year survival, at least one year of follow-up time is required to be included in the analysis. Data are refreshed once a year. After the report on transplant center-specific survival rates is approved by HRSA, the Transplant Center Directory is repopulated with the new data.

Summary

A censored data logistic regression model is fitted to survival data for first unrelated and related donor hematopoietic cell transplants at U.S. centers. The model is adjusted for recipient age, recipient race, Karnofsky/Lansky score, Sorror HCT-CI, adult BMI group, pediatric BMI group, recipient CMV status, history of mechanical ventilation, history of invasive fungal infection, prior autologous transplant, disease/stage, AML ELN risk group, AML transformed from MDS or MPN, number of induction cycles for AML in CR1, interval from diagnosis to transplant in ALL and AML in CR2 and CR3+/Relapse, ALL cytogenetic risk group, Philadelphia positive-status in ALL, number of induction cycles for ALL in CR1, MDS IPSS-R risk score at HCT, therapy related MDS, MDS predisposing condition, CLL and other chronic leukemia disease status, NHL subtype, sensitivity to chemotherapy in NHL and HL, Plasma cell disorder disease status, year of transplant, donor type/graft type/HLA matching, BM or PBSC donor/recipient sex match, unrelated BM or PBSC donor age at transplant, and recipient median household income. The report on transplant center-specific survival rates helps to identify centers that may have under-performed or over-performed compared to the overall network of transplant centers during this specified time period.

References

- [1] National Marrow Donor Program, "Center-Specific Report," Internal NMDP Report, 1998.
- [2] Andersen PK, Klein JP and Rosthoj S, "Generalized linear models or correlated pseudo-observations with applications to multi-state models," *Biometrika*, vol. 90, pp. 15-27, 2003.
- [3] Klein JP and Andersen PK, "Regression modeling of competing risks data based on pseudo-values of the cumulative incidence function," *Biometrics,* vol. 61, pp. 223-229, 2005.
- [4] Klein JP, Logan BR, Harhoff M and Andersen PK, "Analyzing survival curves at a fixed point in time," *Statistics in Medicine*, vol. 26, pp. 4505-4519, 2007.
- [5] "Recommendations from the 2010 Center-Specific Outcomes Analysis Forum," http://www.cibmtr.org/Meetings/Materials/CSOAForum/Documents/Center_Forum_Summary2010.pdf, 2010.
- [6] Sorror ML, "How I assess comorbidities before hematopoietic cell transplantation," *Blood*, vol. 121, no. 15, pp. 2854-2863, 2013.
- [7] Döhner H, Estey E, Grimwade D, et al., "Diagnosis and management of AML in adults: 2017 ELN recommendations from an international expert panel," *Blood*, vol. 129, no. 4, pp. 424-447, 2017.
- [8] Moorman AV, Chilton L, Wilkinson J, et al, "A population-based cytogenetic study of adults with acute lymphoblastic leukemia," *Blood,* vol. 115, no. 2, pp. 206-214, 2010.
- [9] Greenberg PL, Tuechler H, Schanz J, et al., "Revised international prognostic scoring system for myelodysplastic syndromes," *Blood*, vol. 120, no. 12, pp. 2454-2465, 2012.
- [10] Palumbo A, Avet-Loiseau H, Olivia S, et al., "Revised international staging system for multiple myeloma: a report from International Myeloma Working Group," *Journal of clinical oncology*, vol. 33, no. 26, pp. 2863-2869, 2015.
- [11] Logan BR, Nelson G and Klein JP, "Analyzing center-specific outcomes in hematopoietic cell transplantation. Lifetime Data Analysis," *Lifetime Data Analysis*, vol. 14, pp. 389-404, 2008.
- [12] Weisdorf D, Spellman S, Haagenson M, et al., "Classification of HLA-matching for retrospective analysis of unrelated donor transplantation: revised definitions to predict survival. Biol Blood Marrow Transplant," *Biol Blood Marrow Transplant*, vol. 14, pp. 748-758, 2008.