Autologous and allogeneic hematopoietic cell transplantation for diffuse large B-cell lymphoma–type Richter syndrome

Alex F. Herrera,1 Kwang Woo Ahn,2,3 Carlos Litovich,2 Yue Chen,3 Amer Assal,4 Qaiser Bashir,5 Rutheeu-Lu Bayer,6 Melanie Coleman,7 Zachariah DeFilippis,8 Noshra Farhadfar,9 Matthew Greenwood,10 Theresa Hahn,11 Mitchell Horwitz,12 Caron Jacobson,13 Samantha Jaglowski,14 Sylvie Lachance,15 Amelia Langston,16 Bassam Mattar,17 Richard T. Maziarz,18 Joseph McGuirk,19 Mohammad A. H. Mian,20 Sunita Nathan,21 Adrienne Phillips,22 Kevin Rakszawski,23 Henrik Sengelov,24 Shalini Shenoy,25 Robert Stuart,26 Craig S. Sauter,27,28 Mohamed A. Kharfan-Dabaja,29 and Mehdi Hamadani2

1Department of Hematology and Hematopoietic Cell Transplantation, City of Hope National Medical Center, Duarte, CA; 2Center for International Blood and Marrow Transplant Research, Department of Medicine, and 3Division of Biostatistics, Institute for Health and Equity, Medical College of Wisconsin, Milwaukee, WI; 4Columbia University Irving Medical Center, Department of Medicine, Bone Marrow Transplant and Cell Therapy Program, New York, NY; 5Department of Stem Cell Transplantation and Cellular Therapy, Division of Cancer Medicine, The University of Texas MD Anderson Cancer Center, Houston, TX; 6North Shore University Hospital, Northwell Health Cancer Institute, Long Island, NY; 7Indiana Blood & Marrow Transplantation, Indianapolis, IN; 8Hematopoietic Cell Transplant and Cellular Therapy Program, Massachusetts General Hospital, Boston, MA; 9Division of Hematology/Oncology, University of Florida College of Medicine, Gainesville, FL; 10Royal North Shore Hospital, St. Leonards, NSW, Australia; 11Department of Medicine, Roswell Park Comprehensive Cancer Center, Buffalo, NY; 12Duke University Medical Center, Durham, NC; 13Dana Farber Cancer Institute, Boston, MA; 14Division of Hematology, The Ohio State University Medical Center, Columbus, OH; 15Institut Universitaire d’Hématologie et Thérapie Cellulaire, Université de Montréal, Montréal, QC, Canada; 16Department of Hematology and Medical Oncology, Winship Cancer Institute of Emory University, Atlanta, GA; 17Ascension Via Christi Hospitals, Wichita, KS; 18Adult Blood and Marrow Stem Cell Transplant Program, Knight Cancer Institute, Oregon Health and Science University, Portland, OR; 19Division of Hematologic Malignancies and Cellular Therapeutics, The University of Kansas Cancer Center, Kansas City, KS; 20Georgia Cancer Center, Augusta University Medical Center, Augusta, GA; 21Section of Bone Marrow Transplant and Cell Therapy, Rush University Medical Center; Chicago, IL; 22New York Presbyterian Hospital at Cornell, Weill Cornell Medical College, New York, NY; 23Penn State Cancer Institute, Penn State Health; 24University Hospital, Rigshospitalet, Denmark; 25Washington University/St. Louis Children’s Hospital, St. Louis, MO; 26Hollings Cancer Center, Medical University of South Carolina, Charleston, SC; 27Memorial Sloan Kettering Cancer Center, Department of Medicine, New York, NY; 28Department of Medicine, Weill Cornell Medical College, New York, NY; and 29Division of Hematology-Oncology, Blood and Marrow Transplantation Program, Mayo Clinic, Jacksonville, FL

Richter syndrome (RS) represents a transformation from chronic lymphocytic leukemia/small lymphocytic lymphoma (CLL/SLL) to aggressive lymphoma, most commonly diffuse large B-cell lymphoma (DLBCL), which is associated with a dismal prognosis. Patients with DLBCL-RS have poor outcomes with DLBCL-directed therapy; thus, consolidation with hematopoietic cell transplantation (HCT) has been used, with durable remissions observed. Studies reporting HCT outcomes in patients with DLBCL-RS have been small, have not evaluated the prognostic impact of cytogenetic risk factors, and were conducted prior to the era of novel targeted therapy of CLL/SLL. We performed a Center for International Blood and Transplant Research registry study evaluating outcomes after autologous HCT (auto-HCT; n = 53) and allogeneic HCT (allo-HCT; n = 118) in patients with DLBCL-RS treated in the modern era. More auto-HCT recipients were in complete response (CR) at HCT relative to allo-HCT recipients (66% vs 34%), whereas a higher proportion of allo-HCT recipients had 17p deletion (33% vs 7%) and had previously received novel agents (39% vs 10%). In the auto-HCT cohort, the 3-year relapse incidence, progression-free survival (PFS), and overall survival (OS) were 37%, 48%, and 57%, respectively. Among allo-HCT recipients, the 3-year relapse incidence, PFS, and OS were 30%, 43%, and 52%, respectively. In the allo-HCT cohort, deeper response at HCT was associated with outcomes (3-year PFS/OS, 66%/77% CR vs 43%/57% partial response vs 5%/15% resistant; P < .0001 for both), whereas

Submitted 29 March 2021; accepted 6 June 2021; prepublised online on Blood Advances First Edition 8 September 2021; final version published 17 September 2021. DOI 10.1182/bloodadvances.2021004865.

Data sharing requests should be sent to Mehdi Hamadani (mhamadani@mcw.edu). The Center for International Blood and Transplant Research (CIBMTR) supports accessibility of research in accord with the National Institutes of Health Data Sharing Policy and the National Cancer Institute Cancer Moonshot Public Access and Data Sharing Policy. The CIBMTR only releases deidentified datasets that comply with all relevant global regulations regarding privacy and confidentiality. The full-text version of this article contains a data supplement. © 2021 by The American Society of Hematology
Introduction

About 2% to 10% of patients with chronic lymphocytic leukemia/small lymphocytic lymphoma (CLL/SLL) will ultimately develop Richter syndrome (RS), which is characterized by transformation to an aggressive lymphoma, most commonly diffuse large B-cell lymphoma (DLBCL).1–7 DLBCL-RS is associated with a poor prognosis, with estimated overall survival (OS) measured in months and a median OS ≤ 1 year.2–4,9–11 Prognostic factors for outcome after the development of RS include clinical features at the time of transformation (eg, performance status, tumor size, number of prior lines of therapy) and whether the RS is clonally related to the preceding CLL/SLL.4,5,8–11 Patients with DLBCL that is clonally unrelated to the preceding CLL/SLL (~20% of RS) have been reported to have favorable outcomes relative to patients with clonally related DLBCL-RS.5 Patients with DLBCL-RS are typically treated with DLBCL-directed chemotherapy, but response rates and long-term outcomes are lower than observed with de novo DLBCL.3,6,10–14 Because of the poor outcomes after standard DLBCL-directed therapy, consolidative hematopoietic cell transplantation (HCT) has been studied in patients who are chemotherapy sensitive and have produced durable remissions in a subset of patients.8,10,11,15–18 Autologous HCT (auto-HCT) and allogeneic HCT (allo-HCT) have been used for RS, with the largest study being a registry analysis from the European Society for Blood and Marrow Transplantation reporting 3-year relapse-free survival of 27% after allo-HCT (n = 26) and 45% after auto-HCT (n = 34).18 Although multiple studies have suggested improved outcomes for DLBCL-RS patients who undergo HCT, a small minority of patients with DLBCL-RS respond sufficiently to therapy to proceed to HCT. Nevertheless, based on these limited data, it is generally recommended that patients with DLBCL-RS who are responsive to RS-directed therapy, particularly clonally related DLBCL-RS, undergo consolidative HCT.16

The therapeutic landscape of CLL/SLL has shifted dramatically in the past decade. A range of effective targeted therapies, such as Bruton’s tyrosine kinase inhibitors, BCL-2 inhibitors, and phosphatidylinositol 3-kinase inhibitors, produce durable responses in a large proportion of patients, including traditionally high-risk patients with chromosome 17p deletion (del17p) or TP53 aberration.19–31 Of note, the use of novel agents for CLL/SLL does not appear to impact the incidence of RS.15,21,24,32,33 Multiple studies have recently demonstrated that the development of RS in a patient who previously received a novel agent for CLL/SLL tends to be highly aggressive and associated with short survival.29,34,35 Targeted agents have been evaluated as a treatment for DLBCL-RS with objective responses observed, although the median duration of response tends to be short with single-agent therapies.2,11,36,37

Although the modern treatment for CLL/SLL and DLBCL-RS has evolved, no study has comprehensively evaluated HCT outcomes for patients with RS in the modern era. Although the impact of prior exposure to novel therapies on allo-HCT outcomes has been studied in CLL/SLL,38 no study has evaluated the relationship between prior novel therapy exposure and HCT outcomes in patients with RS. In the limited studies performed to date, the primary indicator of outcome in patients with RS who undergo HCT was disease status at the time of transplantation, with patients in response to RS-directed therapy having more favorable outcomes.10,15 However, there has not been an analysis of the impact of cytogenetic risk factors on HCT outcomes in patients with RS. Because RS is rare and prospective data on the topic are not forthcoming, we used the Center for International Blood and Marrow Transplant Research (CIBMTR) registry to evaluate the outcomes after HCT in patients with RS-DLBCL. We included patients who had undergone auto-HCT or allo-HCT prior to and since the availability of novel agents and sought to evaluate the impact of clinical prognostic factors, cytogenetic abnormalities, and the use of novel agents on HCT outcomes.

Methods

This study used data from the CIBMTR, a working group of >380 transplantation centers worldwide that contributed detailed data on HCT to a statistical center at the Medical College of Wisconsin. Participating centers are required to report all HCTs consecutively, and compliance is monitored by on-site audits. Computerized checks for discrepancies, physicians’ review of submitted data, and on-site audits of participating centers ensured data quality. Observational studies conducted by the CIBMTR are performed in compliance with all applicable federal regulations pertaining to the protection of human research participants. Patients provided written informed consent for research. The institutional review boards of the Medical College of Wisconsin and the National Marrow Donor Program approved this study.

Patients

Eligible patients included adults ≥18 years of age with DLBCL-RS (DLBCL transformed from underlying CLL/SLL) who underwent their first auto-HCT or allo-HCT between January of 2007 and December of 2017. Patients with prior auto-HCT were eligible for inclusion in the allo-HCT cohort; in these cases, data from only the first allo-HCT procedure were eligible for inclusion. Patients with DLBCL transformed from other subtypes of lymphoma or patients with non-DLBCL subtype RS (ie, Hodgkin lymphoma or peripheral T-cell lymphoma) were not eligible for this analysis. Eligible patients identified from the registry, additional data were collected on the type and number of therapies (including novel agents) received for antecedent CLL/SLL vs DLBCL-RS, the presence of cytogenetic abnormalities (patients classified according to highest-risk abnormality as any time point), and IGHV mutation testing. Because IGHV mutation data were not available for >95% of patients, these data are not presented.

Study end points and definitions

Disease response at the time of HCT was determined with the standard criteria in use at the time of HCT, including the International Working Group criteria39 or the Lugano Classification.40 Using consensus criteria, the intensity of allo-HCT conditioning regimens was categorized as nonmyeloablative, reduced intensity, or myeloablative.41
Baseline and transplant characteristics are reported descriptively. The primary end point was OS, defined as death from any cause. Secondary end points included progression-free survival (PFS), nonrelapse mortality (NRM), and relapse/progression. For PFS, an event was defined as progression/reapse or death from any cause. NRM was defined as death without preceding lymphoma relapse/progression, with relapse considered a competing risk. Progression/reapse was defined as progressive lymphoma after HCT or lymphoma recurrence after a complete remission, with NRM considered a competing risk. Data regarding histology at relapse (ie, CLL/SLL vs DLBCL) were not available; therefore, any relapse was considered an event. Surviving patients were censored at the date of last follow-up. PFS and OS probabilities were calculated using Kaplan-Meier estimates.

Acute and chronic graft-versus-host disease (GVHD) were graded using established clinical criteria. Neutrophil recovery was defined as the first of 3 successive days with an absolute neutrophil count \( \geq 500 \) per microliter after post-HCT nadir. Platelet recovery was defined as the first of 3 consecutive days with a platelet count \( \geq 20,000 \) per microliter in the absence of platelet transfusion for 7 consecutive days. For neutrophil and platelet recovery, death without hematopoietic recovery, GVHD, relapse, and NRM were calculated to accommodate for competing risks. Cox proportional-hazard analysis for PFS and OS and cumulative incidences for relapse and NRM were used to identify prognostic factors, via forward stepwise selection, for the allo-HCT cohort. The proportional hazard assumption for each variable was examined by testing whether its coefficient is constant over time. The small sample size of the auto-HCT cohort precluded multivariate analysis. Center effect was tested with \( P \) value \( < .05 \) were considered significant. The variables considered in the multivariable regression analysis are shown in supplemental Table 1. Because the regression models for relapse, NRM, PFS, and OS identified 1 significant covariate for each outcome, Kaplan-Meier estimates for PFS and OS and cumulative incidences for relapse and NRM were calculated. All statistical analyses were performed using SAS version 9.4 (SAS Institute, Cary, NC).

Results

Baseline characteristics

One hundred and seventy-one patients with DLBCL-RS who underwent their first HCT between 2007 and 2017 with data reported to the CIBMTR were included. One hundred and eighteen patients received allo-HCT and 53 patients received auto-HCT. Baseline and HCT characteristics for all patients, organized according to type of HCT, are shown in Tables 1 and 2. Among auto-HCT patients, 7% had del17p aberration (although nearly half did not have available cytogenetic data); 5 of 53 (9%) had received a novel agent prior to HCT (all Bruton’s tyrosine kinase inhibitors), including 2 prior to and 3 after the RS; the median number of lines of therapy for the antecedent CLL/SLL was 2 (range, 1–5); the median number of lines of therapy for DLBCL-RS was 2 (range, 1–6); and 66% were in a CR prior to HCT. Six patients who underwent auto-HCT as their first HCT procedure subsequently underwent allo-HCT. Among allo-HCT patients, 33% had del17p; 46 of 118 (39%) had received a novel agent prior to HCT, including 11% prior to RS, 15% after RS, and 13% before and after RS; and 34% were in a CR prior to HCT. The majority of allo-HCT recipients received reduced-intensity/nonmyeloablative conditioning (77%), received calcineurin inhibitor-based GVHD prophylaxis without posttransplant cyclophosphamide (85%), and had a fully matched related or unrelated donor (63%). The median follow-up time in surviving patients in the auto-HCT and allo-HCT cohorts was 48 months (range, 4–98) and 49 months (range, 3–75), respectively.

Hematopoietic recovery, GVHD, NRM, and relapse

The 1-month cumulative incidence of neutrophil recovery among auto-HCT and allo-HCT recipients was 98% (95% confidence interval [CI], 89–100) and 96% (95% CI, 91–99), respectively, and the 100-day cumulative incidence of platelet recovery was 96% (95% CI, 86–100) and 93% (95% CI, 87–97), respectively (Table 3). Among allo-HCT recipients, the 180-day cumulative incidence of grade 2–4 and grade 3–4 acute GVHD was 39% (95% CI, 29–49) and 13% (95% CI, 7–21), respectively, and the 1-year incidence of cGVHD was 37% (95% CI, 27–46). The 1-year NRM and incidence of relapse among auto-HCT recipients were 10% (95% CI, 3–21) and 29% (95% CI, 17–42), respectively (Figure 1A–B; Table 3); they were 23% (95% CI, 16–31) and 23% (95% CI, 16–31) among allo-HCT recipients, respectively (Figure 2A–B; Table 3). Disease status at the time of allo-HCT was associated with 3-year adjusted incidence of relapse: 16% (95% CI, 6–29) among patients in CR at allo-HCT compared with 31% (95% CI, 18–44) in patients with partial response (PR) and 47% (95% CI, 22–69) in patients with resistant disease (Figure 3B; Table 4; \( P = .05 \)). There was not a significant difference in the 3-year adjusted incidence of relapse among allo-HCT recipients with del17p (37%; 95% CI, 22–51) vs those without del17p (32%; 95% CI, 19–44; \( P = .62 \); supplemental Table 2).

PFS and OS

The 1-year and 3-year PFS among auto-HCT recipients was 61% (95% CI, 47–74) and 48% (95% CI, 34–62), respectively, and the 1-year and 3-year OS was 73% (95% CI, 61–84) and 57% (95% CI, 43–70), respectively (Figure 1C–D; Table 3). Among allo-HCT recipients, the 1-year and 3-year PFS was 54% (95% CI, 45–63) and 43% (95% CI, 34–52), respectively, and the 1-year and 3-year OS was 69% (95% CI, 60–77) and 52% (95% CI, 43–61), respectively (Figure 2C–D; Table 3). Disease status at the time of allo-HCT was significantly associated with 3-year adjusted PFS: 66% (95% CI, 51–81) among patients in CR at allo-HCT compared with 43% (95% CI, 29–57) in patients with PR and 5% (95% CI, 0–15) in patients with resistant disease (\( P < .0001 \); Figure 3C; Table 4). Similarly, disease status at allo-HCT was associated with 3-year adjusted OS (\( P < .0001 \); Figure 3D; Table 3). However, the presence of del17p was not associated with adjusted PFS or OS (supplemental Table 2).

Statistical analyses

Baseline patient and HCT characteristics were described. Because there were notable differences in the characteristics of patients who received auto-HCT compared with patients who received allo-HCT (ie, more auto-HCT patients in a complete response [CR] at HCT, very few auto-HCT patients had received prior novel agents, and few patients with high-risk cytogenetics in the auto-HCT cohort) and because of the potential for selection bias in utilizing 1 transplant procedure subsequently underwent allo-HCT. Among allo-HCT patients, 33% had del17p; 46 of 118 (39%) had received a novel agent prior to HCT, including 11% prior to RS, 15% after RS, and 13% before and after RS; and 34% were in a CR prior to HCT. The majority of allo-HCT recipients received reduced-intensity/nonmyeloablative conditioning (77%), received calcineurin inhibitor-based GVHD prophylaxis without posttransplant cyclophosphamide (85%), and had a fully matched related or unrelated donor (63%). The median follow-up time in surviving patients in the auto-HCT and allo-HCT cohorts was 48 months (range, 4–98) and 49 months (range, 3–75), respectively.
Multivariable analysis

A multivariable regression model was constructed to evaluate for covariates associated with NRM, relapse, PFS, and OS in the larger allo-HCT cohort. Covariates are listed in supplemental Table 1. Receipt of 3 regimens for DLBCL-RS was associated with an increased risk for NRM (hazard ratio [HR], 4.05, 95% CI, 1.75-9.39; P = .001, Table 5). Disease status at allo-HCT was associated with poorer PFS (HR for PR, 1.97; 95% CI, 1.10-3.68; P = .03 and HR for resistant, 5.42; 95% CI, 2.73-10.79; P < .001), and poorer OS (HR for PR, 2.05; 95% CI, 1.04-4.06; P = .04 and HR for resistant, 6.63; 95% CI, 3.16-13.92; P < .001; Table 5). Other covariates were not significantly associated with outcomes, including presence.

Table 1. Baseline characteristics according to type of HCT

| Characteristics                        | Allo-HCT (n = 118) | Auto-HCT (n = 53) |
|----------------------------------------|--------------------|------------------|
| Age, median (range), y                 | 61 (26-73)         | 65 (30-79)       |
| Males                                  | 81 (69)            | 37 (70)          |
| White race                             | 96 (81)            | 47 (89)          |
| KPS                                    |                    |                  |
| ≥90                                    | 69 (58)            | 28 (53)          |
| Missing                                | 2 (2)              | 1 (2)            |
| HCT-CI                                 |                    |                  |
| 3+                                     | 41 (35)            | 32 (60)          |
| Missing                                | 8 (7)              | 2 (4)            |
| Prior auto-HCT                         | 4 (3)              | N/A              |
| Cytogenetic abnormalities              |                    |                  |
| 17p ± others                           | 39 (33)            | 4 (7)            |
| 11q deletion ± others (except 17p)     | 10 (9)             | 0                |
| Trisomy 12 ± others (except 17p and/or 11q) | 13 (11)          | 3 (6)            |
| None                                   | 13 (11)            | 13 (24)          |
| Deletion of 13q alone                  | 4 (3)              | 6 (11)           |
| Others                                 | 15 (13)            | 5 (9)            |
| Missing                                | 24 (20)            | 22 (42)          |
| Received novel agents before HCT*      |                    |                  |
| Bruton’s tyrosine kinase inhibitor      | 45 (38)            | 5 (9)            |
| Venetoclax                             | 7 (6)              | 0                |
| PI3K inhibitor                         | 6 (5)              | 0                |
| None                                   | 69 (59)            | 46 (87)          |
| Missing                                | 3 (2)              | 2 (4)            |
| Novel agent                            |                    |                  |
| Yes, before RS                         | 13 (11)            | 2 (4)            |
| Yes, after RS                          | 18 (15)            | 3 (6)            |
| Both                                   | 15 (13)            | 0                |
| Neither                                | 69 (58)            | 46 (87)          |
| Missing                                | 3 (3)              | 2 (4)            |
| Response to first line RS therapy      |                    |                  |
| CR                                     | 31 (26)            | 24 (45)          |
| PR                                     | 25 (21)            | 10 (19)          |
| Refractory disease                     | 41 (35)            | 8 (15)           |
| Not assessed                           | 21 (18)            | 11 (21)          |
| Prior therapy regimens received for CLL/SLL before RS, median (range), n | 2 (1-10) | 2 (1-5) |
| Prior therapy regimens received for RS, median (range), n | 2 (1-8) | 2 (1-6) |
| Time from CLL diagnosis to RS, median (range), mo | 30 (0-296) | 32 (0-166) |

Unless otherwise indicated, data are n (%).

HCT-CI, HCT Comorbidity Index; IGHV, immunoglobulin variable region heavy chain; PI3K, phosphatidylinositol 3-kinase; PS, Karnofsky Performance Status; ±, with or without.

*Total adds up to 130 rather than 118 because some patients received >1 type of novel therapy.
of del17p, conditioning intensity, receipt of prior novel agents (supplemental Table 3), and time between diagnosis of CLL/SLL and diagnosis of DLBCL-RS. No center effect on outcomes was observed.

### Causes of death

Disease relapse/progression was the most common cause of death after auto-HCT (48%) and allo-HCT (39%); causes of death by HCT type are listed in supplemental Table 4. Other frequent causes of death included GVHD (21%) and infection (11%) after allo-HCT, as well as organ failure (9%) after auto-HCT.

### Discussion

DLBCL-RS is a rare and aggressive lymphoma with limited treatment options; it is associated with dismal outcomes. In this registry study,
the largest study of HCT in RS and the first analysis to include a significant number of patients treated with novel targeted therapies, we demonstrate that auto-HCT or allo-HCT resulted in durable remission in nearly half of patients with DLBCL-RS. In the larger allo-HCT cohort, disease response status at the time of HCT was associated with incidence of relapse, PFS, and OS, whereas a higher number of RS-directed lines of therapy was associated with NRM. Among allo-HCT recipients, the presence of del17p or the receipt of a prior novel agent was not associated with outcomes. We did not compare outcomes after auto-HCT against outcomes after allo-HCT because of the differences in the characteristics of the study cohorts (ie, more patients in CR, very few patients had received prior novel agents, and few had high-risk cytogenetics in the auto-HCT cohort, including nearly half with missing cytogenetic data), as well as the possible biases in selecting 1 transplant approach vs the other; however, auto-HCT and allo-HCT appear to provide a proportion of treated patients with a durable remission.

Prior studies of HCT in patients with RS have been limited to small retrospective analyses, mostly single-center experiences of outcomes in all patients with RS, of which a small proportion underwent HCT.\(^\text{10,11,15,17}\) Our study confirmed the important finding from previous studies, including an European Society for Blood and Marrow Transplantation multicenter registry study from Cwynarski et al that disease status at the time of allo-HCT is associated with survival.\(^\text{15}\) The larger number of patients in our study allowed us to compare outcomes in patients with CR at allo-HCT against patients in PR or with resistant disease; indeed, patients with CR had decreased incidence of relapse and improved PFS and OS (3-year PFS and OS of 66% and 77%, respectively) compared with PR and resistant patients.

### Table 3. Outcomes of patients with DLBCL-RS undergoing auto-HCT or allo-HCT

| Outcomes               | Allo-HCT (n = 118) | Auto-HCT (n = 53) |
|------------------------|-------------------|------------------|
|                        | n                  | Prob, % (95% CI)  | n                  | Prob, % (95% CI)  |
| Neutrophil recovery    | 117                | 96 (91-99)       | 52                 | 98 (89-100)       |
| 1 mo                   |                    |                  |                    |                  |
| Platelet recovery      | 114                | 93 (87-97)       | 51                 | 96 (86-100)       |
| 100 d                  |                    |                  |                    |                  |
| Grade 2-4 acute GVHD   | 91                 | 34 (25-44)       | N/A                | N/A               |
| 100 d                  |                    |                  |                    |                  |
| Grade 3-4 acute GVHD   | 91                 | 12 (6-20)        | N/A                | N/A               |
| 6 mo                   |                    | 13 (7-21)        |                    |                  |
| Chronic GVHD           | 100                | 37 (27-46)       | N/A                | N/A               |
| 1 y                    |                    | 27 (19-36)       |                    |                  |
| 2 y                    |                    | 41 (31-51)       |                    |                  |
| NRM                    | 114                | 11 (6-17)        | 49                 | 4 (0-11)          |
| 100 d                  |                    | 23 (16-31)       |                    | 10 (3-21)         |
| 1 y                    |                    | 27 (19-36)       |                    | 15 (6-26)         |
| 3 y                    |                    | 41 (31-51)       |                    |                  |
| Relapse/progression*   | 114                | 23 (16-31)       | 49                 | 29 (17-42)        |
| 1 y                    |                    | 30 (22-39)       |                    | 37 (24-51)        |
| 3 y                    |                    | 54 (45-63)       |                    | 61 (47-74)        |
| PFS                    | 114                | 43 (34-52)       | 49                 | 48 (34-62)        |
| 1 y                    |                    | 43 (34-52)       |                    | 48 (34-62)        |
| 3 y                    |                    | 52 (43-61)       |                    | 57 (43-70)        |
| OS                     | 118                | 69 (60-77)       | 53                 | 73 (61-84)        |
| 1 y                    |                    | 52 (43-61)       |                    |                  |
| 3 y                    |                    | 55 (46-65)       |                    |                  |

N/A, not applicable; Prob, probability.

*Histology at relapse (CLL/SLL vs DLBCL-RS) was not available; therefore, any relapse was considered an event.

28 SEPTEMBER 2021 • VOLUME 5, NUMBER 18
Figure 1. Outcomes after auto-HCT. (A) NRM. (B) Relapse/progression. (C) PFS. (D) OS. Histology at relapse (CLL/SLL vs DLBCL-RS) was not available; therefore, any relapse was considered a relapse and PFS event.

Figure 2. Outcomes after allo-HCT. (A) NRM. (B) Relapse/progression. (C) PFS. (D) OS. Histology at relapse (CLL/SLL vs DLBCL-RS) was not available; therefore, any relapse was considered a relapse and PFS event.
with dismal outcomes, in our study, in which 39% of the allo-HCT cohort had received prior novel agents and 24% had received the novel agent prior to transformation to DLBCL-RS, receipt of a novel agent was not associated with outcome after allo-HCT. This parallels the observation that the type of prior novel agent received, the number of prior novel agents received, and the receipt of only novel agents vs receipt of novel agents and chemoimmunotherapy prior to allo-HCT are not associated with PFS or OS after allo-HCT in patients with CLL. There were too few patients who received prior novel agents (9%) or had del17p (10%) to assess their prognostic impact in the auto-HCT cohort, which may reflect a bias in the patients selected for auto-HCT.

Our study has several limitations, including the inherent limitations of potential selection bias and center-specific transplantation practices with a multicenter registry analysis. Because a prospective study in this population is not forthcoming, our registry study is important for

Table 4. Outcomes after allo-HCT according to disease status at HCT

| Outcomes               | CR (n = 40) | PR (n = 51) | Resistant (n = 20) |
|------------------------|-------------|-------------|-------------------|
|                        | n | Prob, % (95% CI) | n | Prob, % (95% CI) | n | Prob, % (95% CI) |
| Adjusted relapse*      | 40 | 13 (5-25) | 29 (17-42) | 32 (12-53) | .11          |
| 1 y                    | 49 | 18 (6-29) | 31 (18-44) | 47 (22-69) | .05          |
| Adjusted NRM           | 40 | 18 (5-31) | 23 (12-34) | 33 (14-52) | .47          |
| 1 y                    | 49 | 25 (10-41) | 25 (14-36) | 37 (16-58) | .61          |
| Adjusted PFS*          | 40 | 72 (58-86) | 47 (33-61) | 26 (7-46) | <.0001       |
| 1 y                    | 49 | 66 (51-81) | 43 (29-67) | 5 (0-15) | <.0001       |
| Adjusted OS            | 40 | 80 (67-92) | 67 (54-80) | 45 (23-67) | .02          |
| 1 y                    | 51 | 51 (36-65) | 51 (36-65) | 51 (36-65) | .61          |

Prob, probability.

*Histology at relapse (CLL/SLL vs DLBCL-RS) was not available; therefore, any relapse was considered an event.

Figure 3. Outcomes after allo-HCT according to disease status at allo-HCT. (A) NRM. (B) Relapse/progression. (C) PFS. (D) OS. Histology at relapse (CLL/SLL vs DLBCL-RS) was not available; therefore, any relapse was considered a relapse and PFS event.
understanding the outcomes and risk factors for treatment failure after HCT in DLBCL-RS. Any study of HCT in DLBCL-RS is subject to the reality that a selected minority of patients diagnosed with DLBCL-RS undergo HCT (10-12% in studies that report HCT as a proportion of all diagnosed RS patients\(^\text{10,11}\)), as well as that these patients may represent a biologically more favorable group. We did not have information about the clonal relationship between the DLBCL-RS and antecedent CLL/SLL, which has been identified as an important prognostic factor of outcome with DLBCL-RS.\(^\text{5}\) Clonally unrelated DLBCL-RS is associated with more favorable outcomes, and we cannot exclude that there may be patients with clonally unrelated disease included in this study. This especially may have been true in the auto-HCT cohort, given the low proportion of confirmed del17p and high proportion of CR at HCT in that group. TP53 disruption/del17p are less common in clonally unrelated DLBCL-RS.\(^\text{5}\) About 40% of patients in the allo-HCT cohort in this study had del17p, and outcomes were similar regardless of del17p status, suggesting that our results are applicable across DLBCL-RS patients undergoing allo-HCT. Information regarding the histology at relapse after HCT (DLBCL-RS vs the underlying CLL/SLL) was not available because of the limitations of the registry data. This limits the precision of our relapse and PFS estimates because some patients may have had relapse of CLL/SLL rather than DLBCL-RS. Post-HCT progression of CLL/SLL vs DLBCL-RS has starkly different clinical implications: relapsed DLBCL-RS would be expected to be more difficult to control and more likely to impact survival. Still, our results represent the most conservative “worst-case” estimate of outcomes specific to DLBCL-RS because all DLBCL-RS relapses were included.

Although our study evaluated patients with DLBCL-RS who were diagnosed and treated in the era of novel targeted therapy of CLL/SLL, there are newer immunotherapies that are being studied in DLBCL-RS and were not reflected in our study, including checkpoint inhibitors,\(^\text{52,53}\) bispecific antibodies,\(^\text{54}\) and chimeric antigen receptor–modified T cells.\(^\text{55}\) In future studies, it will be important to study the impact of these therapies on HCT outcomes in patients with DLBCL-RS. Future work should also address the optimal timing of HCT in relation to these newer immunotherapies, including chimeric antigen receptor–modified T cells. Our study provides important data on observed outcomes after HCT in the modern era that will hopefully guide decision making for transplant-eligible patients with DLBCL-RS who are responsive to therapy.

In conclusion, we observed that a sizable proportion of patients with DLBCL-RS who were transplanted in the modern era achieved a durable remission with auto-HCT or allo-HCT. Among recipients of allo-HCT, outcomes were independent of receipt of prior novel agents or cytogenetic risk, and deeper response at the time of transplantation was associated with decreased relapse and improved PFS and OS. Allo-HCT outcomes were dismal among treatment-resistant patients; these patients should be considered for alternative management approaches. Although a minority of patients with DLBCL-RS will respond to treatment and be considered for HCT, our data suggest

### Table 5. Multivariable regression analysis in allo-HCT cohort

|               | n   | HR  | 95% CI, lower limit | 95% CI, upper limit | P     | Overall P |
|---------------|-----|-----|---------------------|---------------------|-------|-----------|
| **NRM**       |     |     |                     |                     |       |           |
| Prior therapy regimens received, n |     |     |                     |                     |       |           |
| 1             | 51  | 1   |                     |                     |       | .01       |
| 2             | 27  | 1.38| 0.52                | 3.63                | .52   |           |
| ≥3            | 30  | 4.05| 1.75                | 9.39                | <.001 |           |
| Missing       | 6   | 0.96| 0.12                | 7.59                | .97   |           |
| **Relapse**   |     |     |                     |                     |       |           |
| Disease status |     |     |                     |                     |       |           |
| CR            | 40  | 1.00|                     |                     | <.001 |           |
| PR            | 49  | 2.65| 1.10                | 6.38                | .03   |           |
| Resistant     | 19  | 6.81| 2.49                | 18.59               | <.001 |           |
| Missing       | 6   | 0.87| 0.11                | 7.07                | .90   |           |
| **PFS**       |     |     |                     |                     |       |           |
| Disease status |     |     |                     |                     |       |           |
| CR            | 40  | 1.00|                     |                     | <.001 |           |
| PR            | 49  | 1.97| 1.07                | 3.61                | .03   |           |
| Resistant     | 19  | 5.42| 2.73                | 10.79               | <.001 |           |
| Missing       | 6   | 0.78| 0.18                | 3.39                | .74   |           |
| **OS**        |     |     |                     |                     |       |           |
| Disease status |     |     |                     |                     |       |           |
| CR            | 40  | 1.00|                     |                     | <.001 |           |
| PR            | 51  | 2.05| 1.04                | 4.06                | .04   |           |
| Resistant     | 20  | 6.63| 3.18                | 13.92               | <.001 |           |
| Missing       | 7   | 1.05| 0.23                | 4.70                | .95   |           |

*Histology at relapse (CLL/SLL vs DLBCL-RS) was not available; therefore, any relapse was considered an event.
that HCT should be considered for responding patients with DLBCL-RS, regardless of whether they have del17p or have received prior novel therapies, and, especially, for those in CR.

Acknowledgments

The CIBMTR is supported primarily by Public Health Service U24CA076518 from the National Institutes of Health National Cancer Institute, the National Heart, Lung, and Blood Institute, and the National Institute of Allergy and Infectious Diseases. It is also supported by the Health Resources and Services Administration (HHS/250201700006C and HHS/250201700007C) and the Office of Naval Research (N00014-20-1-2705 and N00014-20-1-2832). Additional federal support is provided by the National Institutes of Health National Cancer Institute (R01CA215134), the National Institutes of Health National Institute of Allergy and Infectious Diseases (R01AI128775 U01AI126612), the National Institutes of Health National Heart, Lung, and Blood Institute (R01HL130388), the National Institutes of Health National Eye Institute (UG1HL06924), and the Biomedical Advanced Research and Development Authority. Support is also provided by Be the Match Foundation, Boston Children’s Hospital, Dana Farber, St. Baldrick’s Foundation, Stanford University, the Medical College of Wisconsin the National Marrow Donor Program, and the following commercial entities: Actinium Pharmaceuticals, Inc., Adienne SA, AlloVir, Inc., Amgen, Inc., Angiocrine Bioscience, Astellas Pharma US, bluebird bio, Inc., Bristol Myers Squibb Co., Celgene Corp., CSL Behring, CytoSen Therapeutics, Inc., Daiichi Sankyo Co., Ltd., ExCellThera, Fate Therapeutics, Gamida-Cell Ltd., Genentech Inc., Incyte Corporation, Janssen/Johnson & Johnson, Jazz Pharmaceuticals, Inc., Kiadis Pharma, Kite, a Gilead Company, Kyowa Kirin, Legend Biotech, Magenta Therapeutics, Merck Sharp & Dohme Corp., Millennium, the Takeda Oncology Co., Miltenyi Biotec, Inc., Novartis Pharmaceuticals Corporation, Omeros Corporation, OncoImmune, Inc., Orca Biosystems, Inc., Pfizer, Inc., Pharmacyclics, LLC, Sanofi Genzyme, StemCyte, Takeda Pharma, Vor Biopharma, and Xenoskus BV.

The views expressed in this article do not reflect the official policy or position of the National Institutes of Health, the Department of the Navy, the Department of Defense, the Health Resources and Services Administration, or any other agency of the US government.

Authorship

Contribution: M. Hamadani, A.F.H., M.A.K., and C.S.S. conceived and designed the study; Y.C., K.W.A., C.L. collected and assembled data; Y.C., K.W.A., C.L., and M. Hamadani analyzed data; A.F.H. and M. Hamadani wrote the manuscript; and all authors interpreted data and revised the manuscript.

Conflict-of-interest disclosure: A.F.H. has received research funding from Bristol Myers Squibb, Merck, Genentech, Inc./F. Hoffmann-La Roche Ltd., Gilead Sciences, Seattle Genetics, AstraZeneca, and ADC Therapeutics; has acted as a consultant for Bristol Myers Squibb, Merck, Genentech, Inc./F. Hoffmann-La Roche Ltd., Gilead Sciences, Seattle Genetics, and Karyopharm and has had travel, accommodations, and expenses paid by Bristol Myers Squibb. M. Hamadani has received research support/funding from Takeda Pharmaceutical Company, Spectrum Pharmaceuticals, and Astellas Pharma; has acted as a consultant in Janssen, Incyte Corporation, ADC Therapeutics, Celgene Corporation, Omeros, Verastem, and MorphoSys; and has been a member of the Speaker’s Bureau for Sanofi Genzyme, AstraZeneca, and BeiGene. A.A. has received grants from Incyte Corporation, outside of the submitted work. J.M. reports other from AlloVir HCP, Juno Therapeutics, Inc., Gilead-Kite Pharmaceuticals, and Magenta Therapeutics, outside of the submitted work. M.A.K.-D. reports consultancy for Daichi Sankyo, outside of the submitted work. C.S.S. has received grants from Juno Therapeutics, Celgene, Bristol-Myers Squibb, Sanofi Genzyme, and Precision BioSciences and has received personal fees from Precision BioSciences, Sanofi Genzyme, Juno Therapeutics, Spectrum Pharmaceuticals, Novartis, Gennab, Kite, a Gilead Company, Celgene, Gamida Cell, Karyopharm Therapeutics, and GlaxoSmithKline, outside of the submitted work. R.T.M. has received grants, nonfinancial support, and other from Novartis and Bristol Myers Squibb; nonfinancial support and other from Kite Therapeutics; and other from Artiva Biotherapeutics, outside the submitted work. K.R. reports personal fees from Seattle Genetics, outside the submitted work. C.J. has received personal fees from Kite, Novartis, BMS/Celgene, Precision Biosciences, Nkarta, Lonza, AbbVie, and bluebird bio, outside the submitted work. The remaining authors declare no competing financial interests.

ORCID profiles: A.F.H., 0000-0002-9665-7415; K.W.A., 0000-0003-4567-8037; A.A., 0000-0003-4707-9265; O.B., 0000-0002-4086-6453; Z.D.F., 0000-0002-7994-8974; M.G., 0000-0002-3667-2613; S.J., 0000-0002-4335-2554; A.L., 0000-0003-0649-1915; J.M.G., 0000-0002-0539-4796; M.A.K.-D., 0000-0001-7394-5185; M.H., 0000-0001-5372-510X.

Correspondence: Mehdi Hamadani, Center for International Blood and Marrow Transplant Research, Medical College of Wisconsin, 9200 W. Wisconsin Ave, Suite C5500, Milwaukee, WI 53226; e-mail: mhamadani@mcw.edu.

References

1. Ben-Dali Y, Hleuhal MH, da Cunha-Bang C, et al. Richter’s transformation in patients with chronic lymphocytic leukaemia: a nationwide epidemiological study. Leuk Lymphoma. 2020;61(8):1435-1444.

2. Ding W. Richter transformation in the era of novel agents. Hematology (Am Soc Hematol Educ Program). 2018;2018(1):256-263.

3. Parikh SA, Kay NE, Shanafelt TD. How we treat Richter syndrome. Blood. 2014;123(11):1647-1657.

4. Parikh SA, Rabe KG, Call TG, et al. Diffuse large B-cell lymphoma (Richter syndrome) in patients with chronic lymphocytic leukaemia (CLL): a cohort study of newly diagnosed patients. Br J Haematol. 2013;162(6):774-782.

5. Rossi D, Spina V, Deambrogi C, et al. The genetics of Richter syndrome reveals disease heterogeneity and predicts survival after transformation. Blood. 2011;117(12):3391-3401.
34. Maddocks KJ, Ruppert AS, Lozanski G, et al. Etiology of ibrutinib therapy discontinuation and outcomes in patients with chronic lymphocytic leukemia. *JAMA Oncol.* 2015;1(1):80-87.

35. Jain P, Thompson PA, Keating M, et al. Long-term outcomes for patients with chronic lymphocytic leukemia who discontinue ibrutinib. *Cancer.* 2017;123(12):2268-2273.

36. Davids MS, Roberts AW, Seymour JF, et al. Phase I first-in-human study of venetoclax in patients with relapsed or refractory non-Hodgkin lymphoma. *J Clin Oncol.* 2017;35(8):826-833.

37. Tsang M, Shanafelt TD, Call TG, et al. The efficacy of ibrutinib in the treatment of Richter syndrome. *Blood.* 2015;125(10):1676-1678.

38. Roeker LE, Dreger P, Brown JR, et al. Allogeneic stem cell transplantation for chronic lymphocytic leukemia in the era of novel agents. *Blood Adv.* 2020;4(16):3977-3989.

39. Cheson BD, Pfistner B, Juweid ME, et al; International Harmonization Project on Lymphoma. Revised response criteria for malignant lymphoma. *J Clin Oncol.* 2007;25(5):579-586.

40. Cheson BD, Fisher RI, Barrington SF, et al; United Kingdom National Cancer Research Institute. Recommendations for initial evaluation, staging, and response assessment of Hodgkin and non-Hodgkin lymphoma: the Lugano classification. *J Clin Oncol.* 2014;32(27):3059-3068.

41. Bacigalupo A, Ballen K, Rizzo D, et al. Defining the intensity of conditioning regimens: working definitions. *Biol Blood Marrow Transplant.* 2009;15(12):1628-1633.

42. Przepiorka D, Weisdorf D, Martin P, et al. 1994 Consensus Conference on Acute GVHD Grading. *Bone Marrow Transplant.* 1995;15(6):825-828.

43. Shulman HM, Sullivan KM, Weiden PL, et al. Chronic graft-versus-host syndrome in man. A long-term clinicopathologic study of 20 Seattle patients. *Am J Med.* 1980;69(2):204-217.

44. Schetelig J, van Biezen A, Brand R, et al. Allogeneic hematopoietic stem-cell transplantation for chronic lymphocytic leukemia with 17p deletion: a retrospective European Group for Blood and Marrow Transplantation analysis. *J Clin Oncol.* 2008;26(31):5094-5100.

45. Dreger P, Döhner H, Ritgen M, et al; German CLL Study Group. Allogeneic stem cell transplantation provides durable disease control in poor-risk chronic lymphocytic leukemia: long-term clinical and MRD results of the German CLL Study Group CLL3X trial. *Blood.* 2010;116(14):2438-2447.

46. Krämer I, Stilgenbauer S, Dietrich S, et al. Allogeneic hematopoietic cell transplantation for high-risk CLL: 10-year follow-up of the GCLLSG CLL3X trial. *Blood.* 2017;130(12):1477-1480.

47. Epperla N, Ahn KW, Khanal M, et al. Impact of reduced-intensity conditioning regimens on outcomes in diffuse large B cell lymphoma undergoing allogeneic transplantation. *Biol Blood Marrow Transplant.* 2021;27(1):58-66.

48. Ghosh N, Ahmed S, Ahn KW, et al. Association of reduced-intensity conditioning regimens with overall survival among patients with non-Hodgkin lymphoma undergoing allogeneic transplant. *JAMA Oncol.* 2020;8(7):1011-1018.

49. Hamadani M, Saber W, Ahn KW, et al. Impact of pretransplantation conditioning regimens on outcomes of allogeneic transplantation for chemotherapy-unresponsive diffuse large B cell lymphoma and grade III follicular lymphoma. *Biol Blood Marrow Transplant.* 2013;19(5):746-753.

50. Shah NN, Hamadani M. Is there still a role for allogeneic transplantation in the management of lymphoma? *J Clin Oncol.* 2021;39(5):487-498.

51. Kharfan-Dabaja MA, Moukalled N, Reljic T, El-Asmar J, Kumar A. Reduced intensity is preferred over myeloablative conditioning allogeneic HCT in chronic lymphocytic leukemia whenever indicated: a systematic review/meta-analysis. *Hematol Oncol Stem Cell Ther.* 2018;11(2):53-64.

52. Armand P, Murawski N, Molin D, et al. Pembrolizumab in relapsed or refractory Richter syndrome. *Br J Haematol.* 2020;190(2):e117-e120.

53. Ding W, LaPlant BR, Call TG, et al. Pembrolizumab in patients with CLL and Richter transformation or with relapsed CLL. *Blood.* 2017;129(26):3419-3427.

54. Alderuccio JP, Mackrides N, Chapman JR, Vega F, Lossos IS. Rapid complete response to blinatumomab as a successful bridge to allogeneic stem cell transplantation in a case of refractory Richter syndrome. *Leuk Lymphoma.* 2018;60(1):230-233.

55. Kittai AS, Bond DA, William B, et al. Clinical activity of axicabtagene ciloleucel in adult patients with Richter syndrome. *Blood Adv.* 2020;4(19):4648-4652.