Allogeneic transplant and CAR-T therapy after autologous transplant failure in DLBCL: a noncomparative cohort analysis

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Key Points

- CIBMTR prognostic score predicts PFS and OS of patients with DLBCL receiving axicabtagene ciloleucel treatment after a prior autoHCT failure.
- CIBMTR high/very high-risk score marks an adverse risk cohort where novel immunotherapy or relapse prevention approaches are warranted.

Allogeneic transplant (alloHCT) and chimeric antigen receptor modified (CAR-T) cell therapy are potentially curative options of diffuse large B-cell lymphoma (DLBCL) relapsing after an autologous (auto)HCT. Although the Center for International Blood and Marrow Transplant Research (CIBMTR) prognostic model can predict outcomes of alloHCT in DLBCL after autoHCT failure, corresponding models of CAR-T treatment in similar patient populations are not available. In this noncomparative registry analysis, we report outcomes of patients with DLBCL (≥18 years) undergoing a reduced intensity alloHCT or CAR-T therapy with axicabtagene ciloleucel during 2012 to 2019 after a prior auto-HCT failure and apply the CIBMTR prognostic model to CAR-T recipients. A total of 584 patients were included. The 1-year relapse, nonrelapse mortality, overall survival (OS), and progression-free survival for CAR-T treatment after autoHCT failure were 39.5%, 4.8%, 73.4%, and 55.7%, respectively. The corresponding rates in the alloHCT cohort were 26.2%, 20.0%, 65.6%, and 53.8%, respectively. The 1-year OS of alloHCT recipients classified as low-, intermediate- and high/very high-risk groups according to the CIBMTR prognostic score was 73.3%, 59.9%, and 46.3%, respectively (P = .002). The corresponding rates for low-, intermediate-, and high/very high-risk CAR-T patients were 88.4%, 76.4%, and 52.8%, respectively (P < .001). This registry analysis shows that both CAR-T and alloHCT can provide durable remissions in a subset of patients with DLBCL relapsing after a prior autoHCT. The simple CIBMTR prognostic score can be used to identify patients at high risk of treatment failure after either procedure. Evaluation of novel relapse mitigation strategies after cellular immunotherapies are warranted in these high-risk patients.
Introduction

Diffuse large B-cell lymphoma (DLBCL) is a curable disease, and approximately 50% to 60% of patients do not require further treatment after an initial anthracycline and rituximab-containing regimen. The current standard-of-care for fit patients with relapsed/refractory DLBCL, includes high-dose chemotherapy and autologous hematopoietic cell transplant (autoHCT) consolidation in those with disease responding to salvage therapies or chimeric antigen receptor modified T-cell (CAR-T) treatment of those with disease refractory to at least 2 prior therapy lines. The outcomes for patients with DLBCL relapsing after autoHCT have been historically poor, with median survival of generally less than 1 year. Depending on prior treatments, patients with DLBCL relapsing after an autoHCT, in the modern era, can potentially benefit from a number of approved agents (eg, polatuzumab, tafasitamab, selinexor, or loncastuximab tesirine), but these options are generally not expected to provide durable disease control. Cellular immunotherapies directed against defined lymphoma-specific antigens (eg, anti-CD19 CAR-T treatment) or against undefined tumor antigens after allogeneic (allo)HCT (ie, the graft-versus-lymphoma effect), are potentially curative in DLBCL, even after failure of high-dose therapy and autoHCT. Because of the favorable outcome data reported for CAR-T, acceptable toxicity profile, and feasibility in patients with refractory relapse, the application of alloHCT in patients with DLBCL with a failed prior autoHCT has witnessed a rapid decline. Although this shift in practice is also endorsed by recent expert consensus, large studies providing contemporary, supportive evidence for this change in patient management are lacking. In addition, although prognostic models able to predict outcomes of alloHCT in DLBCL after autoHCT failure have been developed, to our knowledge, corresponding models of CAR-T treatment in similar patient populations are not available. A single prognostic model predictive of outcomes of both CAR-T and alloHCT performed after autoHCT failure may help identify clinical settings where 1 type of cellular therapy may be preferable over the other.

Using the Center for International Blood and Marrow Transplant Research (CIBMTR) registry, we retrospectively report alloHCT and CAR-T treatment outcomes in patients with DLBCL experiencing relapse after prior autoHCT. In addition, we applied the previously published CIBMTR prognostic score to the current patient cohorts to identify patients subsets at high risk for treatment failure with modern cell-based therapy approaches.

Methods

Data source

The CIBMTR is a collaborative working group of more than 380 international transplant and cellular immunotherapy centers managed by the Medical College of Wisconsin (MCW) and the National Marrow Donor Program. Detailed information on basic patient characteristics, demographics, clinical variables, and outcome data are contributed by members to the statistical center at the MCW. Participating centers are required to report and frequently update the information on all hematopoietic cell transplants. Cellular therapy data for any CAR-T recipients are collected longitudinally from 130 participating centers in the United States and Canada. Integrity and quality of data are monitored at different levels, including on-site audits and computerized checking for discrepancies. All studies conducted by the CIBMTR comply with applicable regulations to protect research participants.

Patients

Adult patients with DLBCL (age ≥ 18 years), undergoing a first alloHCT or CAR-T therapy during 2012 and 2019, with history of a failed prior autoHCT, were included in this analysis. The CAR-T cohort was limited to treatment with axicabtagene ciloleucel (axi-cel). The alloHCT cohort was limited to patients receiving reduced-intensity (RIC) or nonmyeloablative (NMA) conditioning platforms. Eligible donors for the alloHCT cohort included matched sibling, 8/8 matched unrelated (allele-level match at HLA-A, -B, -C, and -DRB1), or related haploidentical donors. Graft-versus-host disease (GVHD) prophylaxis was limited to conventional calcineurin inhibitor–based or posttransplant cyclophosphamide-based (± calcineurin inhibitor and mycophenolate mofetil) approaches. Patients receiving ex vivo graft manipulation (eg, CD34 selection), or those with history of DLBCL transforming from indolent histologies were excluded.

Definitions and end points

The intensity of alloHCT conditioning regimens was categorized as RIC/NMA using the consensus criteria. Disease response at the time of HCT was determined using the International Working Group criteria in use during the era of this analysis. Progression-free survival (PFS), was defined as the time from either alloHCT or CAR-T treatment to relapse/progression or death from any cause. Overall survival (OS) was defined as the time from treatment to death from any cause, the cumulative incidence of nonrelapse mortality (NRM) was defined as death without preceding disease progression, and the cumulative incidence of relapse/progression was defined as the time from treatment to relapse or disease progression.

Acute and chronic GVHD were graded using established clinical criteria. Neutrophil recovery was defined as the first of 3 successive days with an absolute neutrophil count ≥ 500/µL after post-HCT or CAR-T nadir. Platelet recovery was defined as the first of 3 consecutive days with a platelet count of 20 000/µL or higher in the absence of platelet transfusion for 7 consecutive days. For neutrophil and platelet recovery, death without the event was considered a competing risk. For patients receiving CAR-T, the cumulative incidence of cytokine release syndrome (CRS) and immune effector cell–associated neurotoxicity syndrome (ICANS) were calculated. CRS and ICANS were graded using the American Society of Transplant and Cellular Therapy grading criteria.

Statistical analysis

Baseline patient, alloHCT, and CAR-T characteristics were assessed using the Pearson χ² test for categorical variables and the Kruskal-Wallis test for continuous variables. Because there were notable differences in the characteristics of patients who received alloHCT compared with patients who received CAR-T (ie, alloHCT patients were younger, had better performance status, and more frequently had chemosensitive disease) and because of the potential for selection bias in using 1 approach vs the other, we did not compare outcomes between the alloHCT and CAR-T cohorts. The Kaplan-Meier estimator was used to analyze OS and PFS. The cumulative incidence rates of hematopoietic recovery, NRM, and...
relapse/progression rates were calculated while accounting for competing events. Four patients after alloHCT relapse underwent CAR-T therapy, and 5 relapsing patients after CAR-T received alloHCT. For OS analyses, these 9 subjects receiving alternative cellular therapy were not censored. In addition, we aimed to apply the previously published CIBMTR prognostic score (shown to predict PFS and OS for DLBCL undergoing alloHCT after a failed prior autoHCT)\(^\text{18}\) to the current cohort of a more contemporary population of patients with DLBCL undergoing either a RIC/NMA alloHCT or CAR-T treatment after a failed autoHCT. The details of CIBMTR prognostic score were previously published.\(^\text{18}\) Briefly the model consists of 3 adverse prognostic factors, including Karnofsky performance score (KPS) < 80 (4 points), autoHCT to alloHCT or CAR-T interval < 1 year (2 points), and chemoresistant disease at alloHCT or CAR-T (5 points). This CIBMTR prognostic model classifies patients into 4 groups: low risk (0 points), intermediate risk (2-5 points), high risk (6-9 points), or very high risk (11 points). Because of the small sample size of the very high-risk group (alloHCT = 9; CAR-T = 5), it was combined with the high-risk group for this analysis.

### Table 1. Characteristics of adult patients with DLBCL who received CAR-T or alloHCT

| Characteristic                              | CAR-T (N = 181) | alloHCT (N = 403) | P    |
|--------------------------------------------|----------------|------------------|------|
| Median age, y (range)                      | 61.0 (21.9-80.0) | 56.7 (18.5-72.9) | <.01*|
| Male sex, no. (%)                          | 117 (64.6)      | 261 (64.8)       | 1.00†|
| Karnofsky performance score 90-100        | 79 (43.6)       | 243 (60.3)       | <.01†|
| HCT Comorbidity Index ≥ 3, no. (%)        | 73 (40.3)       | 189 (46.9)       | .04‡ |
| Missing                                    | 9 (5.0)         | 7 (1.7)          |      |
| Median time from diagnosis to CAR-T or alloHCT (mo) | 34.1 (2.7-282.3) | 40.8 (3.2-340.5) | .12* |
| >12 mo                                     | 175 (96.7)      | 384 (95.3)       |      |
| Median time from prior autoHCT to CAR-T or alloHCT (mo) | 14.6 (3.2-268.7) | 15.5 (1.2-196.9) | .58‡ |
| >12 mo                                     | 97 (53.6)       | 208 (51.6)       |      |
| Graft source, no. (%)                      | N/A             | N/A              |      |
| Bone marrow                                | N/A             | 40 (9.9)         |      |
| Peripheral blood                           | N/A             | 363 (90.1)       |      |
| Donor type, no. (%)                        | N/A             | N/A              |      |
| HLA-identical sibling                      | N/A             | 189 (46.9)       |      |
| Matched unrelated donor                    | N/A             | 153 (38.0)       |      |
| Haploidentical related                     | N/A             | 61 (15.1)        |      |
| Year of CAR-T or alloHCT, no. (%)          | N/A             | N/A              | <.01‡|
| 2012-2014                                  | 0 (0.0)         | 201 (49.9)       |      |
| 2015-2017                                  | 2 (1.1)         | 171 (42.4)       |      |
| 2018-2019                                  | 179 (98.9)      | 31 (7.7)         |      |
| Disease status at CAR-T/alloHCT, no. (%)   | N/A             | N/A              | <.01‡|
| Complete remission                         | 9 (5.0)         | 220 (54.6)       |      |
| Partial remission                          | 39 (21.5)       | 103 (25.8)       |      |
| Resistant/untreated relapse                | 122 (67.4)      | 50 (12.4)        |      |
| Unknown                                    | 11 (6.1)        | 30 (7.4)         |      |
| Lymphodepleting chemotherapy, no. (%)     | N/A             | N/A              |      |
| Cyclophosphamide                           | 1 (0.6)         | N/A              |      |
| Cyclophosphamide + fludarabine             | 180 (99.4)      | N/A              |      |
| Bridging therapy use, no. (%)              | 35 (19.3)       | N/A              | N/A  |
| CIBMTR prognostic score, no. (%)           | N/A             | N/A              |    <.01|
| Low (score 0)                              | 54 (29.8)       | 212 (52.6)       |      |
| Intermediate (score 2,4,5)                 | 78 (43.1)       | 148 (36.7)       |      |
| High/very high (score 6,7,9,11)            | 49 (27.1)       | 43 (10.7)        |      |
| Follow-up, median (range) (mo)             | 13.0 (1.0-27.7) | 51.8 (0.2-98.6)  |      |

N/A, not applicable.
*Kruskal-Wallis test.
†Fisher exact test.
‡Fisher exact test via Mon.
Results

Baseline characteristics

We identified 584 patients with DLBCL who received either CAR-T treatment (n = 181) or an alloHCT (n = 403) after prior autoHCT failure. Baseline characteristics are summarized in Table 1. The median age of CAR-T recipients was 56.7 years, and 79 patients (43.6%) had a KPS of 90 to 100, and most had refractory disease (n = 122; 67.4%). Fludarabine and cyclophosphamide were used for lymphodepletion in 180 patients. Thirty-five patients received bridging treatment between the apheresis session(s) and start of lymphodepletion. The alloHCT cohort was significantly younger (median age, 56.7 years) and had a significantly greater proportion of patients with better performance status (KPS 90-100 = 261 [60.3%]) and chemosensitive disease (80.2%). Peripheral blood and matched siblings were the most common graft (90.1%) and donor (46.9%) source, respectively.

CAR-T outcomes

The median follow-up of CAR-T survivors was 13 months (range, 1.0-27.7; Table 2). The rates of grade ≥ 3 CRS and ICANS were 9.9% and 20.9%, respectively. The day 28 cumulative incidence of neutrophil recovery and day 100 incidence of platelet recovery was 89.7% (95% CI = 84.7-93.8) and 86.7% (95% CI = 81.2-91.4), respectively. At 1 year, the cumulative incidence rates of relapse/progression and NRM were 39.5% (95% CI = 32.1-47.2) and 4.8% (95% CI = 2.1-8.6), respectively (Figure 1A-B). The probability of 1-year OS and DFS was 73.4% (95% CI = 66.4-79.9) and 55.7% (95% CI = 48.0-63.2), respectively (Figure 1C-D).

Allogeneic HCT outcomes

The median follow-up of alloHCT survivors was 51.8 months (range, 0.2-27.7; Table 2). The day 180 cumulative incidence rates of grade 2 to 4 and 3 to 4 acute GVHD were 28.7% (95% CI = 18.9-39.7) and 8.2% (95% CI = 3.0-15.6), respectively. The cumulative incidence of chronic GVHD at 1 year was 35.6% (95% CI = 30.9-40.4). The day 28 cumulative incidence of neutrophil recovery and day 100 incidence of platelet recovery was 96.8% (95% CI = 94.8-98.3) and 92.1% (95% CI = 89.2-94.6), respectively. At 1 year, the cumulative incidence rates of relapse/progression and NRM were 26.2 (95% CI = 21.9-30.7) and 20.0% (95% CI = 16.2-24.2), respectively (Figure 1E-F). The probability of 1-year OS and DFS was 65.6% (95% CI = 60.9-70.2) and 53.8 (95% CI = 48.8-58.7), respectively (Figure 1G-H).

CIBMTR prognostic score

In the CAR-T cohort 54 (29.8%), 78 (43.1%) and 40 (27.1%) patients were classified into low-, intermediate- and high/very high-risk groups according to the CIBMTR prognostic score, respectively (Table 1). The 1-year PFS probabilities for the low-, intermediate- and high/very high-risk groups were 62.6% (95% CI = 55.9-69.1), 46.6% (95% CI = 38.4-54.9), and 34.1% (95% CI = 20.6-49.2), respectively (P < .001; Figure 2C; Table 3). The 1-year OS probabilities in similar order were 88.4% (95% CI = 78.4-95.6), 76.4% (95% CI = 65.4-85.8), and 52.8% (95% CI = 35.5-66.9), respectively (P < .001; Figure 2D; Table 3).

Table 2. Outcomes of CAR-T and alloHCT

| Characteristic                  | CAR-T          | alloHCT        |
|--------------------------------|----------------|----------------|
| CRS grade 1-5, no. (%)         | 149 (82.3)     | N/A            |
| CRS grade 3-5, no. (%)         | 18 (9.9)       | N/A            |
| Neurotoxicity grade 1-4, no. (%)| 112 (61.9)     | N/A            |
| Neurotoxicity grade 3-4, no. (%)| 38 (20.9)      | N/A            |
| Neutrophil recovery, % (95% CI)| Day 28 89.7 (84.7-93.8) | 96.8 (94.8-98.3) |
| Platelet recovery, % (95% CI)  | Day 100 86.7 (81.2-91.4) | 92.1 (89.2-94.6) |
| Grade 2-4 acute GVHD, % (95% CI)| Day 180 N/A | 28.7 (18.9-39.7) |
| Grade 3-4 acute GVHD, % (95% CI)| Day 180 N/A | 8.2 (3.0-15.6) |
| Chronic GVHD, % (95% CI)       | 1 y 73.4 (66.4-79.9) | 65.8 (60.9-70.2) |
| OS, % (95% CI)                 | 1 y 55.7 (48.0-63.2) | 53.8 (48.8-58.7) |
| PFS, % (95% CI)                | 1 y 39.5 (32.1-47.2) | 26.2 (21.9-30.7) |
| Relapse/progression, % (95% CI)| 1 y 4.8 (2.1-8.6) | 20.0 (16.2-24.2) |
| NRM, % (95% CI)                |                 | N/A, not applicable. |

The CIBMTR prognostic score classified 212 (52.6%), 148 (36.7%), and 43 (10.7%) of alloHCT patients into low-, intermediate- and high/very high-risk groups, respectively (Table 1). The 1-year PFS probabilities for the low-, intermediate-, and high/very high-risk groups were 62.6% (95% CI = 55.9-69.1), 46.6% (95% CI = 38.4-54.9), and 34.1% (95% CI = 20.6-49.2), respectively (P < .001; Figure 2C; Table 3). The 1-year OS probabilities in similar order were 73.3% (95% CI = 67.1-79.1), 59.9% (95% CI = 51.9-67.7), and 46.3% (95% CI = 31.5-61.5), respectively (P = .002; Figure 2D; Table 3).

Subset analysis

In an exploratory subset analysis, we looked at the outcomes of patients with refractory or untreated relapse (n = 172). The 1-year PFS and OS of patients receiving CAR-T with refractory or untreated relapse (n = 122) was 51.5% and 71%, respectively. The corresponding estimates for alloHCT patients (n = 50) were 37.6 and 49%, respectively.

Cause of death

During the follow-up, 199 patients (49.3%) from the alloHCT group and infections (9.1%; n = 5) and second cancers (5.5%; n = 3) new myeloid malignancies) in the CAR-T group.
Discussion

In this retrospective analysis using the CIBMTR registry, we describe the outcomes of patients with DLBCL relapsing after an autoHCT and subsequently undergoing either CAR-T therapy or alloHCT. We also applied the CIBMTR prognostic score to these patient cohorts to identify patients at high risk of treatment failure.

Our noncomparative cohort analysis shows the following after autoHCT failure: (a) CAR-T treatment in a cohort of mostly refractory patients provides 1-year PFS of 55% with a 5% NRM risk, (b) alloHCT in a group of predominantly chemosensitive patients produces a 1-year PFS of 54% with a 20% NRM risk, and (c) the CIBMTR prognostic score group of high/very high-risk disease identifies a cohort of patients at particularly high risk of therapy failure after either CAR-T therapy or alloHCT.

At the onset, it is important to acknowledge that the cohort of patients getting CAR-T treatment and alloHCT in our study are fundamentally different. For example, patients receiving CAR-T were older, less fit, and mostly had refractory disease. Patients in the alloHCT cohort were younger, had a better performance score, and most had disease responding to salvage treatments. Most CAR-T procedures were done during 2018 to 2019, whereas alloHCT spanned a wider time period (2012-2019). These important differences warrant caution while interpreting even an adjusted comparison across the 2 cohorts (as shown in supplemental Table 1). The short median follow-up of CAR-T-treated patients means that the long-term outcomes cannot be analyzed using the current dataset. Chemoresponse is not a prerequisite for offering CAR-T therapy, meaning a patient relapsing after autoHCT may be able to receive this treatment without salvage attempts. The large real-world analyses have also shown feasibility of this treatment in older individuals with comorbid conditions. In contrast, establishing chemosensitivity and careful patient selection are critical for alloHCT outcomes. To reduce heterogeneity, the CAR-T cohort in the current analysis was limited to axi-cel, as it represented the predominant commercial product reported to the CIBMTR registry during the analysis period.

There is a paucity of data directly comparing outcomes of alloHCT vs CAR-T treatment in DLBCL. Dregger et al compared outcomes of CAR-T and alloHCT in relapse/refractory DLBCL by retrospectively applying the intention-to-treat principle. The analysis (not limited to the post-autoHCT failure setting) included 41 and 60 patients for whom CAR-T cells and alloHCT were intended, respectively. In both cohorts, virtually all patients had active disease at the time of indication. The 1-year estimates for NRM, relapse, PFS, and OS for CAR-T cells vs alloHCT were 3% vs 21% (P = .04), 59% vs

Figure 1. Outcomes of CAR-T cell therapy after autologous transplant failure. (A) Progression/relapse. (B) NRM. (C) OS. (D) PFS. Outcomes of alloHCT after autoHCT failure. (E) Progression/relapse. (F) NRM. (G) OS. (H) PFS.
In our analysis, we demonstrated that the CIBMTR prognostic score can predict PFS and OS of both alloHCT and CAR-T in patients with DLBCL with a prior autoHCT failure. The main advantage of this model is the ease of its use, because it uses easily available clinical variables. We acknowledge that additional variables (eg, metabolic tumor volumes, CAR-T cell fitness, composition and expansion, bilirubin level, lactate dehydrogenase, Eastern Cooperative Oncology Group performance score) have been shown to predict outcomes of CAR-T therapy but are not generally available in the CIBMTR registry. It possible a that a prognostic model incorporating such variables may have better discriminatory ability in

Table 3. PFS and OS after CAR-T cell or alloHCT according to the CIBMTR prognostic score

|                      | Low risk (score 0) | Intermediate risk (score 2, 4, 5) | High/very high risk (score 6, 7, 9, 11) | Overall P |
|----------------------|--------------------|----------------------------------|----------------------------------------|-----------|
| CAR-T cohort         |                    |                                  |                                        |           |
| 1-y OS (95% CI)      | 88.4 (78.4-95.6)   | 76.4 (65.4-86.8)                 | 52.8 (38.5-66.9)                       | <.001     |
| 1-y PFS (95% CI)     | 75.8 (62.9-86.6)   | 54.3 (42.3-66.1)                 | 34.9 (21.9-49.1)                       | <.001     |
| alloHCT cohort       |                    |                                  |                                        |           |
| 1-y OS (95% CI)      | 73.3 (67.1-79.1)   | 59.9 (51.9-67.7)                 | 46.3 (31.5-61.5)                       | .002      |
| 1-y PFS (95% CI)     | 62.6 (55.9-69.1)   | 46.6 (38.4-54.9)                 | 34.1 (20.6-49.2)                       | <.001     |
post-autoHCT CAR-T recipients. Of note, the 1-year OS of alloHCT patients in the low- (73%), intermediate- (60%), and high/very high-risk (46%) groups of the current analysis (era 2012-2019) compare favorably against the corresponding 1-year rates in the original publication (era 2000-2012: low risk, 63%; intermediate risk, 52%; high/very high risk, 17%-38%). This difference is possibly because of general improvements in HCT supportive care, advances in GVHD prevention and treatment, and the restriction of the current cohort to RIC regimens. The CIBMTR prognostic score of high/very high-risk disease identifies a cohort of patients at particularly high risk of therapy failure after both CAR-T therapy or alloHCT. Effective approaches to mitigate risk of therapy failure after alloHCT or CAR-T treatment of such patients is clearly an area of unmet need, and investigation of novel maintenance or consolidation approaches in this setting are warranted.

In patients with DLBCL with an autoHCT failure, both alloHCT and CAR-T are available options. Although not a direct comparison, the current analysis and the prior report by Dreger et al do not suggest an inferiority of CAR-T treatment in this setting. Indirect comparison of the outcomes of various risk groups identified by the CIBMTR score also does not identify a subgroup where 1 treatment is clearly superior over the other. In the absence of such data, CAR-T treatment has now practically emerged as the preferred cell-based therapy option in post-autoHCT failures, likely because of its safety and documented efficacy in refractory disease. The CIBMTR and European Society for Blood and Marrow Transplantation consensus guidelines, however, identify rare circumstances where alloHCT may be considered as the first cellular immunotherapy in relapsed/refractory DLBCL (eg, in patients with refractory cytopenias with or without signs of therapy-related myelodysplasia, cases with CAR-T manufacturing failure, or in regions or centers where CAR-T treatment is not available). In addition, limited data suggest that alloHCT may salvage a subset of CAR-T recipients with evidence of persistent or recurrent disease. Because, in such patients, higher-intensity conditioning regimens are associated with higher NRM and inferior OS, it is important to use lower-intensity conditioning platforms to minimize the risk of procedure-related morbidity and mortality.

In conclusion, this noncomparative, retrospective, registry analysis shows that both CAR-T and alloHCT can provide durable remissions in a subset of patients with DLBCL relapsing after prior autoHCT. The profiles of patients benefiting from these approaches are distinct. The simple-to-use CIBMTR prognostic score can be used to identify patients at high risk of treatment failure after either procedure in clinical practice or incorporated in the design of trials evaluating novel relapse mitigations strategies after cellular immunotherapies in patients with DLBCL with a prior autoHCT failure.

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Authorship

Contribution: M.H., F.L.L., A.K.G., and M.P. conceived and designed the study; X.O. collected and assembled the data; X.O. and S.K. analyzed the data; M.H. prepared the first draft of the manuscript; and all authors interpreted the data and revised the manuscript.

Conflict-of-interest disclosure: M.H. received research support/consulting from Takeda Pharmaceutical Company, Spectrum Pharmaceuticals, and Astellas Pharma; consulted for Janssen, Incyte Corporation, ADC Therapeutics, Celgene Corporation, Omeros, Verastem, and MorphoSys; and was on the speaker's bureau for Sanofi Genzyme, AstraZeneca, and Beigene. A.K.G. received research funding from Bristol Myers Squibb, Pharmacics, Gilead, Genentech, AstraZeneca, Pfizer, Teva, Takeda, Acorete, IgM, H-Mab, Agios, and Merck and consulted for Abbvie, Genentech, Janssen, AstraZeneca, Pharmacics, Bristol Myers Squibb, Amgen, Morphosys, TG Therapeutics, Kite Pharma, Adaptive, SeaGen, Epizyme, Gilead, ADCT, Incyte, Karyopharm, Actinium, Asana bio, Nurix, and Aptevio. F.L.L. was a scientific advisor for Allogene, Amgen, Bluebird Bio, BMS/Celgene, Calib, Cellular Biomedicine Group, GammaDelta Therapeutics, Iovance, Kite Pharma, Janssen, Legend Biotech, Novartis, Wugen, and Umoja; received research funding from Kite Pharma (institutional), Alogene (institutional), Novartis (institutional); and holds several patents (held by the institution in his name; unlicensed) in the field of cellular immunotherapy. S.C. was a scientific advisor for Allogene, Acorete, Cellularity, CareDx, Kiadis Pharma, Milteny Biotech, and Molmed and received research funding from Milteny and Kiadis Pharma. P.B.D. sat on an advisory board for Kite (Gilead). J.G. received institutional research support from CCSG and SCCA Swim Across America and received consulting fees from JMP, Eusapharma, Larvol, and Multerra Bio. The remaining authors declare no competing financial interests.

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1. Sehn LH, Salles G. Diffuse large B-cell lymphoma. *N Engl J Med*. 2021;384(9):842-858.

2. Philip T, Guglielmi C, Hagenbeek A, et al. Autologous bone marrow transplantation as compared with salvage chemotherapy in relapses of chemotherapy-sensitive non-Hodgkin's lymphoma. *N Engl J Med*. 1995;333(23):1540-1545.

3. Shah NN, Ahn KW, Litovitch C, et al. Is autologous transplant in relapsed DLBCL patients achieving only a PET+ PR appropriate in the CAR T-cell era [correction published in *Blood*. 2021;137(20):2854-2855]? *Blood*. 2021;137(10):1416-1423.

4. Hamadani M, Hari PN, Zhang Y, et al. Early failure of frontline rituximab-containing chemo-immunotherapy in diffuse large B cell lymphoma does not predict futility of autologous hematopoietic cell transplantation. *Biol Blood Marrow Transplant*. 2014;20(11):1729-1736.

5. Gisselbrecht C, Glass B, Mounier N, et al. Salvage regimens with autologous transplantation for relapsed large B-cell lymphoma in the rituximab era. *J Clin Oncol*. 2010;28(27):4184-4190.

6. Abramson JS, Palomba ML, Gordon LI, et al. Lisocabtagene maraleucel for patients with relapsed or refractory large B-cell lymphomas (TRANSCEND NHL 001): a multicentre seamless design study. *Lancet*. 2020;396(10254):839-852.

7. Jacobson CA, Hunter BD, Redd R, et al. Axicabtagene ciloleucel in the non-trial setting: outcomes and correlates of response, resistance, and toxicity. *J Clin Oncol*. 2020;38(27):3095-3106.

8. Locke FL, Ghobadi A, Jacobson CA, et al. Long-term safety and activity of axicabtagene ciloleucel in refractory large B-cell lymphoma (ZUMA-1): a single-arm, multicentre, phase 1-2 trial. *Lancet Oncol*. 2019;20(1):31-42.

9. Neelapu SS, Locke FL, Bartlett NL, et al. Axicabtagene ciloleucel CAR T-cell therapy in refractory large B-cell lymphoma. *N Engl J Med*. 2017;377(26):2531-2544.

10. Schuster SJ, Svoboda J, Chong EA, et al. Chimeric antigen receptor T cells in refractory B-cell lymphomas. *N Engl J Med*. 2017;377(26):2545-2554.

11. Epperla N, Badar T, Szabo A, et al. Postrelapse survival in diffuse large B-cell lymphoma after therapy failure following autologous transplantation. *Blood Adv*. 2019;3(11):1661-1669.

12. Sehn LH, Herrera AF, Flowers CR, et al. Polatuzumab vedotin in relapsed or refractory diffuse large B-cell lymphoma. *J Clin Oncol*. 2020;38(2):155-165.

13. Kalakonda N, Maerevoet M, Cavallo F, et al. Selinexor in patients with relapsed or refractory diffuse large B-cell lymphoma (SADAL): a single-arm, multinational, multicentre, open-label, phase 2 trial. *Lancet Haematol*. 2020;7(7):e511-e522.

14. Caimi PF, Al W, Alderuccio JP, et al. Loncastuximab tesirine in relapsed or refractory diffuse large B-cell lymphoma (LOTIS-2): a multicentre, open-label, single-arm, phase 2 trial. *Lancet Oncol*. 2021;22(6):790-800.

15. Hamadani M, Radford J, Carlo-Stella C, et al. Final results of a phase 1 study of loncastuximab tesirine in relapsed/refractory B-cell non-Hodgkin lymphoma. *Blood*. 2021;137(19):2634-2645.

16. Salles G, Duell J, Gonzalez Barca E, et al. Tafasitamab plus lenalidomide in relapsed or refractory diffuse large B-cell lymphoma (L-MIND): a multicentre, prospective, single-arm, phase 2 study. *Lancet Oncol*. 2020;21(7):978-988.

17. Dreger P, Sureda A, Ahn KW, et al. PTCy-based haploidentical vs matched related or unrelated donor reduced-intensity conditioning transplant for DLBCL. *Blood Adv*. 2019;3(3):360-369.

18. Fenske TS, Ahn KW, Graff TM, et al. Allogeneic transplantation provides durable remission in a subset of DLBCL patients relapsing after autologous transplantation. *Br J Haematol*. 2016;174(2):235-248.

19. Hamadani M, Benson DM, Jr., Hofmeister CC, et al. Allogeneic stem cell transplantation for patients with relapsed chemorefractory aggressive non-Hodgkin lymphomas. *Biol Blood Marrow Transplant*. 2009;15(5):547-553.

20. 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.

21. Urbano-Ispizua A, Pavletic SZ, Flowers ME, et al. The impact of graft-versus-host disease on the relapse rate in patients with lymphoma depends on the histological subtype and the intensity of the conditioning regimen. *Biol Blood Marrow Transplant*. 2015;21(10):1746-1753.

22. Rezvani AR, Norasetthada L, Gooley T, et al. Non-myeloablative allogeneic haematopoietic cell transplantation for relapsed diffuse large B-cell lymphoma: a multicentre experience. *Br J Haematol*. 2008;143(3):395-403.

23. Thomson KJ, Morris EC, Bloor A, et al. Favorable long-term survival after reduced-intensity allogeneic transplantation for multiple-relapse aggressive non-Hodgkin’s lymphoma. *J Clin Oncol*. 2009;27(3):426-432.

24. van Kampen RJ, Canals C, Schouten HC, et al. Allogeneic stem-cell transplantation as salvage therapy for patients with diffuse large B-cell non-Hodgkin’s lymphoma relapsing after an autologous stem-cell transplantation: an analysis of the European Group for Blood and Marrow Transplantation Registry. *J Clin Oncol*. 2011;29(10):1342-1348.

25. Dreger P, Fenske TS, Montoto S, Pasquini MC, Sureda A, Hamadani M; European Society for Blood and Marrow Transplantation (EBMT) and the Center for International Blood and Marrow Transplant Research (CIBMTR). Cellular immunotherapy for refractory diffuse large B cell lymphoma in the chimeric antigen receptor-engineered T cell era: still a role for allogeneic transplantation? *Biol Blood Marrow Transplant*. 2020;26(4):e77-e85.

26. 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.
27. 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.

28. 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.

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

30. 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.

31. Lee DW, Santomasso BD, Locke FL, et al. ASTCT consensus grading for cytokine release syndrome and neurologic toxicity associated with immune effector cells. *Biol Blood Marrow Transplant.* 2019;25(4):625-638.

32. Locke FL, Rossi JM, Neelapu SS, et al. Tumor burden, inflammation, and product attributes determine outcomes of axicabtagene ciloleucel in large B-cell lymphoma. *Blood Adv.* 2020;4(19):4898-4911.

33. 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;6(7):1011-1018.

34. 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.

35. Dreger P, Dietrich S, Schubert ML, et al. CAR T cells or allogeneic transplantation as standard of care for advanced large B-cell lymphoma: an intent-to-treat comparison. *Blood Adv.* 2020;4(24):6157-6168.

36. Dean EA, Mhaskar RS, Lu H, et al. High metabolic tumor volume is associated with decreased efficacy of axicabtagene ciloleucel in large B-cell lymphoma. *Blood Adv.* 2020;4(14):3268-3276.

37. Nastoupil LJ, Jain MD, Feng L, et al. Standard-of-care axicabtagene ciloleucel for relapsed or refractory large B-cell lymphoma: results from the US Lymphoma CAR T Consortium. *J Clin Oncol.* 2020;38(27):3119-3128.

38. Zeiser R, Blazar BR. Acute graft-versus-host disease: biologic process, prevention, and therapy. *N Engl J Med.* 2017;377(22):2167-2179.

39. 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. *Transplant Cell Ther.* 2021;27(1):58-66.

40. Shadman M, Gauthier J, Hay KA, et al. Safety of allogeneic hematopoietic cell transplant in adults after CD19-targeted CAR T-cell therapy. *Blood Adv.* 2019;3(20):3062-3069.