Cardiovascular Effects of CAR T Cell Therapy
A Retrospective Study

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ABSTRACT

BACKGROUND Anti-CD19 chimeric antigen receptor (CAR) T cell therapy holds great promise in the treatment of patients with hematologic malignancies. A high occurrence of cardiac dysfunction has been noted in children treated with CAR T cell therapy.

OBJECTIVES The aim of this study was to define the occurrence of major adverse cardiovascular events (MACE) in adult patients treated with CAR T cell therapy and assess the relationships among clinical factors, echocardiographic parameters, laboratory values, and cardiovascular outcomes.

METHODS Baseline clinical, laboratory, and echocardiographic parameters were collected in 145 adult patients undergoing CAR T cell therapy. MACE included cardiovascular death, symptomatic heart failure, acute coronary syndrome, ischemic stroke, and de novo cardiac arrhythmia. Baseline parameters associated with MACE were identified using Cox proportional cause-specific hazards regression analysis.

RESULTS Thirty-one patients had MACE (41 events) at a median time of 11 days (interquartile range: 6 to 151 days) after CAR T cell infusion. The median follow-up period was 456 days (interquartile range: 128 to 1,214 days). Sixty-one patients died. Cytokine release syndrome (CRS) occurred 176 times in 104 patients; the median time to CRS was 6 days (interquartile range: 1 to 8 days). The Kaplan-Meier estimates for MACE and CRS at 30 days were 17% and 53%, respectively. The Kaplan-Meier estimates for survival at 1 year was 71%. Multivariable Cox proportional cause-specific hazards regression analysis determined that baseline creatinine and grade 3 or 4 CRS were independently associated with MACE.

CONCLUSIONS Patients treated with CAR T cell therapy are at an increased risk for MACE and may benefit from cardiovascular surveillance. Further large prospective studies are needed to confirm the incidence and risk factors predictive of MACE. (J Am Coll Cardiol CardioOnc 2020;2:193–203) © 2020 The Authors. Published by Elsevier on behalf of the American College of Cardiology Foundation. This is an open access article under the CC BY-NC-ND license (http://creativecommons.org/licenses/by-nc-nd/4.0/).
ANTI-CD19 CHIMERIC ANTIGEN RECEPTOR (CAR) T CELL THERAPY, FIRST DEVELOPED AT THE UNIVERSITY OF PENNSYLVANIA, IS A TREMENDOUS ADVANCE IN THE FIELD OF HEMATOLOGIC MALIGNANCIES. THE PATIENT’S OWN T CELLS ARE ENGINEERED TO TARGET CD19, AN ANTIGEN FREQUENTLY AND HIGHLY EXPRESSED IN SOME B CELL MALIGNANCIES, NOTABLY ACUTE LYMPHOBLASTIC LEUKEMIA (ALL), CHRONIC LYMHPHOCYTIC LEUKEMIA (CLL), AND DIFFUSE LARGE B-CELL LYMPHOMA (DLBCL) (1, 2). IN CLINICAL TRIALS, CAR T CELL THERAPY HAS SHOWN A REMARKABLE RESPONSE RATE (70% TO 90%) AND DURABILITY OF REMISSION; IN A CHEMOTHERAPY-RESISTANT DLBCL POPULATION, 65% OF PATIENTS WERE RECURRENCE FREE AT 1 YEAR (3–5). TO DATE, THERE ARE 2 U.S. FOOD AND DRUG ADMINISTRATION-APPROVED CD19 IMMUNOTHERAPIES FOR REFRACTORY B-CELL ALL AND LARGE B-CELL LYMPHOMA, TISAGENECLEUCEL (KYMRIAH, NOVARTIS, BASEL, SWITZERLAND) AND AXICABTAGENE CILOLEUCEL (YESCARTA, KITE PHARMA, SANTA MONICA, CALIFORNIA).

A FREQUENT AND POTENTIALLY LETHAL SIDE EFFECT OF CAR T CELL THERAPY IS CYTOKINE RELEASE SYNDROME (CRS), WITH SYMPTOMS INCLUDING MILD TO HIGH FEVER, SEVERE HYPOTENSION, AND HYPOXIA, OCCURRING USUALLY WITHIN DAYS AFTER CAR T CELL THERAPY. IN A RECENT LARGE SYSTEMATIC REVIEW AND META-ANALYSIS, CRS WAS REPORTED IN MORE THAN ONE-HALF OF THE PATIENTS RECEIVING CAR T CELL THERAPY (6). CRS IS A SYSTEMIC INFLAMMATORY RESPONSE SYNDROME TRIGGERED IN PARTICULAR BY THE RELEASE OF LARGE AMOUNTS OF INTERLEUKIN-6 (7). THE RELEASE OF CYTOKINES INDUCES FEVER, VASCULAR LEAKAGE, AND POTENTIALLY DIRECT MYOCARDIAL INJURY.

CARDIOVASCULAR COMPLICATIONS AND POTENTIAL CARDIOTOXICITY OF CAR T CELL THERAPY ARE ESPECIALLY RELEVANT CONCERNS IN THIS PATIENT POPULATION, AS THEY MAY HAVE PRE-EXISTING CARDIAC DYSFUNCTION OR RISK FACTORS (8) AND/OR RECEIVED MULTIPLE PREVIOUS CYCLES OF CARDIOTOXIC CHEMOTHERAPY. CARDIOVASCULAR COMPLICATIONS OF CAR T CELL THERAPY HAVE BEEN REPORTED IN 2 RETROSPECTIVE STUDIES IN THE PEDIATRIC POPULATION (1, 9) AND RECENTLY IN A THIRD RETROSPECTIVE STUDY IN ADULTS (10). IN 39 CHILDREN WITH ACUTE LEUKEMIA, FITZGERALD ET AL. (9) REPORTED THAT MORE THAN ONE-THIRD OF PATIENTS DEVELOPED CARDIOVASCULAR COMPLICATIONS SUCH AS SHOCK OR CARDIOMYOPATHY. IN THE OTHER STUDY, BY BURSTEIN ET AL. (1), 24 PATIENTS AMONG 98 CHILDREN (24%) HAD HYPOTENSION REQUIRING INOTROPIC SUPPORT. OF THESE 24 PATIENTS, 10 (50% OF CHILDREN WHO UNDERWENT ECHOCARDIOGRAPHY) DEMONSTRATED LEFT VENTRICULAR SYSTOLIC DYSFUNCTION. THE CARDIOMYOPATHY WAS MOSTLY REVERSIBLE IN THESE CHILDREN. LEFT VENTRICULAR DYSFUNCTION IN THE PEDIATRIC HEART, HOWEVER, MAY BE DIFFERENT IN SEVERITY, REVERSIBILITY, AND PROGRESSION COMPARED WITH THE ADULT HEART.

IN THE PRESENT STUDY, WE DEFINED THE RATE OF OCCURRENCE AND THE NATURAL HISTORY OF CARDIOVASCULAR EVENTS IN ALL CONSECUTIVE ADULT PATIENTS TREATED WITH CAR T CELL THERAPY. CLINICAL, LABORATORY, AND ECHOCARDIOGRAPHIC PARAMETERS WERE COLLECTED TO INVESTIGATE THE ASSOCIATION BETWEEN THESE PARAMETERS AND THE CARDIOVASCULAR OUTCOMES OF PATIENTS TREATED WITH CAR T CELL THERAPY.

METHODS

IDENTIFICATION OF PATIENTS AND ENDPOINTS.

THE STUDY WAS CONDUCTED AT THE HOSPITAL OF THE UNIVERSITY OF PENNSYLVANIA (HUP) AND WAS APPROVED BY THE LOCAL INSTITUTIONAL REVIEW BOARD. ALL CONSECUTIVE ADULT PATIENTS (>18 YEARS OF AGE) WITH CD19+ MALIGNANCY (DLBCL, ALL, OR CLL) TREATED WITH EXPERIMENTAL AND COMMERCIAL CAR T CELL THERAPY AT HUP BETWEEN AUGUST 2010 AND JANUARY 2019 WERE IDENTIFIED. MAJOR ADVERSE CARDIOVASCULAR EVENTS (MACE) (11) WERE DEFINED AS CARDIOVASCULAR DEATH, SYMPTOMATIC HEART FAILURE, NONFATAL ACUTE CORONARY SYNDROME, NONFATAL ISCHEMIC STROKE, AND DE NOVO CARDIAC ARRHYTHMIA. TO IDENTIFY THE OCCURRENCE OF MACE, EACH CHART WAS REVIEWED INDIVIDUALLY (B.L.). ALL MACE WERE RECORDED AND DEFINED USING THE AMERICAN COLLEGE OF CARDIOLOGY/AMERICAN HEART ASSOCIATION OUTCOME DEFINITIONS OUTLINED FOR CLINICAL TRIALS BY HICKS ET AL. (11). SYMPTOMATIC HEART FAILURE WAS IDENTIFIED WHEN 3 OR MORE OF THE FOLLOWING 4 CRITERIA WERE MET: SYMPTOMS OF HEART FAILURE, CLINICAL SIGNS OF HEART FAILURE ON PHYSICAL EXAMINATION, LABORATORY OR IMAGING OR RADIOGRAPHIC FINDINGS OF HEART FAILURE (B-TYPE NATRIURETIC PEPTIDE OR N-TERMINAL PRO-B-TYPE NATRIURETIC PEPTIDE, KERLEY B-LINES OR PULMONARY EDEMA, PLEURAL EFFUSION, DECREASED LEFT VENTRICULAR EJECTION FRACTION [LVEF]), AND INITIATION OF NEW TREATMENT FOR HEART FAILURE (PHARMACOTHERAPEUTICS SUCH AS DIURETIC AGENTS AND/OR MECHANICAL SUPPORT). ONE OR MORE SYMPTOMS OF HEART FAILURE AND 2 OR MORE SIGNS ON PHYSICAL EXAMINATION WERE NECESSARY FOR THE DIAGNOSIS. SYMPTOMS OF HEART FAILURE WERE DEFINED AS DYSPNEA AT REST OR DURING EXERCISE, DECREASED EXERCISE CAPACITY, AND SYMPTOMS OF VOLUME OVERLOAD. CLINICAL SIGNS ON PHYSICAL EXAMINATION WERE DEFINED AS PERIPHERAL EDEMA, ASCITES IN THE ABSENCE OF HEPATIC DISEASE, PULMONARY CRACKLES OR RALES, INCREASED JUGULAR VENOUS PRESSURE, S3 GALLOP, AND SIGNIFICANT AND RAPID WEIGHT GAIN RELATED TO FLUID RETENTION. DE NOVO
cardiac arrhythmias such as atrial fibrillation were identified on electrocardiography. All cardiovascular events were adjudicated by 2 independent cardiologists blinded to all other clinical and quantitative echocardiographic data (B.L. and Y.K.). A third cardiologist (M.S.-C.) further adjudicated if any disagreement occurred between the 2 initial reviewers. Signs and symptoms of heart failure arising concurrently with sepsis were excluded from MACE.

Cardiovascular risk factors and pre-existing cardiovascular disease were extracted manually from each electronic medical record. All baseline characteristics were reported at the time of or most proximal to CAR T infusion. Cardiovascular disease was defined by pre-existing coronary artery disease, chronic heart failure, atrial fibrillation, or stroke. Cardiovascular risk factors such as hypertension, hypercholesterolemia, and diabetes were classified as present if both diagnosis and treatment were identified in the medical chart. Other cardiovascular risk factors, such as obesity and smoking history, were also identified. Noncardiac death was also reported and included death due to septic shock, multiple organ failure, or progression of cancer. To minimize loss of follow-up from death, every patient was also searched in a national necrology database. Vital signs and laboratory results were extracted from the HUP electronic medical record.

CRS events were reviewed and graded according to the latest American Society for Transplantation and
Cellular Therapy consensus grading for CRS (12). CRS is defined into 4 grades according to symptoms and clinical status. Grades 3 and 4 CRS include temperature ≥38°C, hypotension requiring at least 1 vasopressor, and/or hypoxia with high-flow oxygen supplementation, positive pressure, or intubation.

ECHOCARDIOGRAPHIC ANALYSIS. Echocardiographic images were extracted from the HUP database. Measurements were acquired by a single observer (B.L.) blinded to clinical data and the occurrence of MACE and reported from the average of 3 consecutive cardiac cycles using the recommendations from the American Society of Echocardiography. LVEF was calculated using the modified Simpson biplane method, and left atrial volume was calculated using the biplane method (13).

STATISTICAL ANALYSIS. Categorical data are expressed as percentages and continuous data as mean ± SD or median (interquartile range [IQR]). Normality was determined using the Kolmogorov-Smirnov test. Differences among patients with ALL, CLL, and DLBCL were determined using a 1-way analysis of variance or the Kruskal-Wallis test. Kaplan-Meier (KM) estimates at 30 days, 6 months, and 12 months were also performed for each cancer subtype and MACE, CRS, and survival rates. The cumulative incidence function was used to estimate the incidence of MACE and noncardiac death.

A Cox proportional cause-specific hazards regression analysis was used to determine the parameters associated with MACE in our population. All variables were tested using univariable analysis. Initially, clinically significant noncollinear variables with p values <0.10 by univariable analysis were entered into a multivariable Cox proportional cause-specific hazards regression. Multicollinearity was tested using the variance inflation factor. The decision was made to use a maximum of 4 variables (41 events, 1 parameter per 10 events) to avoid model instability. To confirm the results, multivariable Cox proportional cause-specific hazards regression was repeated with stepwise regression using the backward elimination method using all variables with p values <0.10 by univariable analysis. All variables were subsequently reentered individually into the final model, to assess for significance in the presence of the final model variables. Hazard ratios (HRs) are expressed with 95% confidence intervals (CIs). All CRS gradings were compared with CRS grade 0 as an indicator. Univariable and multivariable analysis were also confirmed using the Fine and Gray distribution hazard regression model in a competing risk regression model.

Data were analyzed using SPSS version 16.0 (SPSS, Chicago, Illinois) and R version ii386 3.5.0 (R Foundation for Statistical Computing, Vienna, Austria). A p value <0.05 was considered to indicate statistical significance.

RESULTS

CHARACTERISTICS OF PATIENTS TREATED WITH CAR T CELL THERAPY. Between August 2010 and January 2019, a total of 145 patients were identified.
Thirty-six patients (25%) were diagnosed with ALL, 43 patients with DLBCL (30%), and 66 patients (46%) with CLL. Baseline clinical characteristics, reported at the time of or most proximal to CAR T cell infusion, medications, and laboratory results are presented in Table 1. The median follow-up period was 456 days (IQR: 128 to 1,214 days; range: 5 to 3,103 days). There were 176 occurrences of CRS in 104 patients (72%), with a median time to CRS of 6 days (IQR: 1 to 8 days; range: 0 to 175 days). The KM estimates for CRS rate were 53% at 30 days, 64% at 6 months, and 71% at 12 months (Figure 1). Sixty-one patients died, 59 of them of noncardiac causes. The KM estimates for overall survival was 95% at 30 days, 81% at 6 months, and 71% at 12 months (Figure 2). Six patients (4%) were lost to follow-up at a median time of 391 days.

OCCURRENCE AND CHARACTERISTICS OF MACE DEVELOPED BY PATIENTS TREATED WITH CAR T CELL THERAPY. Thirty-one patients developed MACE, for a total of 41 events. All events were considered in the subsequent analyses. The median time to MACE was 11 days after CAR T cell infusion (IQR: 6 to 151 days; range 0 to 2,372 days). The KM estimates for MACE were 17% at 30 days, 19% at 6 months, and 21% at 12 months (Figure 3). The adjudication of all heart failure events is included in Supplemental Table 1. The median follow-up period of patients in the MACE group was 753 days (IQR: 215 to 1,714 days; range: 14 to 3,061 days). There were 22 heart failure events (1 of which was a stress-induced cardiomyopathy) in 21 patients (15%), 12 episodes of atrial fibrillation in 11 patients (7.5%), 2 events of other arrhythmias (supraventricular tachycardia, nonsustained ventricular tachycardia), 2 episodes of acute coronary syndrome, and 2 cardiac deaths. The cardiac deaths were a pulseless electric activity arrest and a massive pulmonary embolism leading to a ST-segment elevation myocardial infarction. Two patients had atrial fibrillation concurrent with the use of ibrutinib and thus were not included among those experiencing MACE. The cumulative incidence of MACE and noncardiac deaths in all patients is shown in Supplemental Figure 1.

ECHOCARDIOGRAPHIC CHARACTERISTICS OF PATIENTS TREATED WITH CAR T CELL THERAPY. Baseline echocardiographic findings are listed in Table 2. LVEF obtained both from echocardiography and
multiple-gated acquisition was available for 124 patients (86%) and averaged 61 ± 9%. Prior to CAR T cell infusion, 110 patients underwent echocardiography (78 of whom had full reports available that could be used for further analysis). The median time from echocardiography to CAR T cell infusion was 43 days (IQR: 21 to 80 days; range: 2 to 615 days) and was not associated with MACE. The mean baseline LVEF in patients who developed MACE was 62 ± 7% (echocardiography performed in 28 patients) and 61 ± 9% in patients who did not. Seventeen patients (55% of those with MACE) underwent repeat echocardiography when MACE occurred. The mean LVEF measured at the time of MACE was 49 ± 14%.

### Table 2: Echocardiographic Parameters at Baseline

| Parameter     | All Patients* (N = 78) | MACE (n = 16) | No MACE* (n = 62) |
|---------------|-----------------------|---------------|-------------------|
| Heart rate    | 75 ± 17               | 72 ± 8        | 75 ± 17           |
| LVEDD (cm)    | 4.65 ± 0.53           | 4.72 ± 0.17   | 4.62 ± 0.52       |
| LVESD (cm)    | 3.15 ± 0.52           | 3.20 ± 0.32   | 3.14 ± 0.52       |
| IVS (cm)      | 0.93 ± 0.18           | 0.97 ± 0.03   | 0.92 ± 0.18       |
| PWT (cm)      | 0.93 ± 0.14           | 0.93 ± 0.01   | 0.93 ± 0.14       |
| RV base (cm)  | 3.45 ± 0.53           | 3.45 ± 0.52   | 3.46 ± 0.54       |
| TAPSE (cm)    | 2.3 ± 0.4             | 2.3 ± 0.5     | 2.3 ± 0.4         |
| RV S’ (mm/s)  | 13.2 ± 3.1            | 13.5 ± 2.5    | 13.2 ± 2.7        |
| MV E          | 67.8 (59.8-81.0)      | 71.8 (63.2-83.1) | 67.6 (59.4-80.7) |
| MV a          | 61.0 (52.9-82.9)      | 56.4 (46.5-67.2) | 64.2 (54.8-87.4) |
| Mitral E/a ratio | 1.2 ± 0.5             | 1.3 ± 0.2     | 1.2 ± 0.5         |
| Mitral E/e’   | 8.7 ± 3.5             | 10.0 ± 1.1    | 8.3 ± 3.3         |
| PASP (mm Hg)  | 27 ± 6                | 30 ± 7        | 26 ± 6            |
| LAVI (cm²/m²) | 29.7 ± 11.0           | 35.1 ± 11.7   | 28.0 ± 10.1       |
| LVEF, Simpson method (%) | 62.1 ± 7.2 | 61.8 ± 7.5 | 62.3 ± 6.5 |

Values are mean ± SD or median (interquartile range). *Patients lost to follow-up and those with noncardiac death prior to MACE are included.

IVS = interventricular septal thickness; LAVI = left atrial volume index; LVEDD = left ventricular end-diastolic diameter; LVESD = left ventricular end-systolic diameter; MV = mitral valve; PASP = pulmonary artery systolic pressure, PWT = posterior wall thickness, RV = right ventricular; TAPSE = tricuspid annular plane systolic excursion; other abbreviations as in Table 1.
(HR: 1.15 for each 1-unit increase; 95% CI: 1.00 to 1.31; p = 0.046) and larger indexed left atrial volume (HR: 1.04 for each 1 ml/m² increase; 95% CI: 1.00 to 1.08; p = 0.075). Analyses of diastolic function using the 2016 guidelines (14) indicated that 31 patients had normal diastolic function (40%), 22 had grade I diastolic dysfunction (28%), 1 had grade II dysfunction (1%), 3 had grade III dysfunction (4%), and 21 had indeterminate diastolic function (27%). There was no association between the presence of diastolic dysfunction and MACE (p = 0.866).

Baseline creatinine, statin treatment, and CRS grading were chosen for the multivariable competing risk analysis according to their clinical relevance, effect size in univariable models, and limited to these variables to avoid multicollinearity. The multivariable competing risk regression analysis revealed that baseline creatinine (HR: 15.54 for each 1 mg/dl increase; 95% CI: 3.67 to 65.86; p < 0.001) and grade 3 CRS (HR: 8.42; 95% CI: 3.48 to 20.40; p < 0.001) or grade 4 CRS (HR: 29.86; 95% CI: 9.80 to 90.94; p < 0.001) were independently associated with MACE (Table 4). These results were confirmed with the stepwise regression using the backward elimination method and by individually adding variables to the final model. The same results were found with the Fine and Gray method (Supplemental Table 5). Statin use at baseline was not independently associated with MACE.

**DISCUSSION**

The present study demonstrates that adult patients with CD19⁺ malignancy treated with CAR T cell therapy are at risk for MACE, mainly symptomatic heart failure, most of which occur within weeks of the infusion (Central Illustration). The variables independently associated with MACE in our study included baseline creatinine and grades 3 and 4 CRS.

Between August 2010 and January 2019, a total of 145 patients were identified. The median age (60 years), male distribution (74%), and distribution of malignancies (25% ALL, 30% DLBCL, and 46% CLL) are representative of the current population receiving CAR T cell therapy (15,16). The KM estimates for overall survival were 95% at 30 days, 81% at 6 months, and 71% at 12 months (Figure 2). The mortality in our cohort is less than previously described, likely because of the improving expertise and expanding applications to patients with less severe conditions and to patients who have received less intensive chemotherapy regimens (4,15,17).

In a recent publication from our group in 450 patients with acute myeloid leukemia or ALL, only 3% of patients with ALL had heart failure (18). Another study investigating patients with hematologic cancers treated with anthracyclines reported that 4% of patients with lymphoma developed symptomatic heart failure at a median time of 523 days after the initiation of anthracycline therapy (19). Our study, with a prevalence of 15% for symptomatic heart failure at a median time of 11 days, suggests a higher heart failure incidence than previously described, underlining the effect of CAR T cell treatment.

**Table 3** Univariable Cox Proportional Cause-Specific Hazards Regression

| Variable                              | Hazard Ratio (95% CI) | p Value |
|---------------------------------------|-----------------------|---------|
| Age                                   | 1.01 (0.98-1.04)      | 0.431   |
| Sex (male)                            | 1.73 (0.66-4.51)      | 0.265   |
| Hypertension                          | 1.40 (0.68-2.89)      | 0.362   |
| Diabetes mellitus                     | 1.70 (0.59-4.86)      | 0.326   |
| Dyslipidemia                          | 1.58 (0.77-3.25)      | 0.212   |
| Congestive heart failure              | 1.17 (0.16-8.60)      | 0.879   |
| Coronary artery disease               | 2.07 (0.79-5.42)      | 0.139   |
| Atrial fibrillation                   | 2.83 (1.08-7.43)      | 0.035   |
| Stroke                                | 2.33 (0.55-9.44)      | 0.250   |
| Chronic kidney disease                | 2.08 (0.63-6.88)      | 0.228   |
| Smoking                               | 1.35 (0.92-1.97)      | 0.122   |
| Radiation                             | 0.66 (0.23-1.90)      | 0.441   |
| Cancer subtypes                       | 0.75 (0.44-1.27)      | 0.278   |
| Transplantation                       | 0.92 (0.35-2.43)      | 0.872   |
| Aspirin                               | 3.13 (1.09-8.99)      | 0.034   |
| ACE inhibitors/ARBs                   | 1.23 (0.50-3.01)      | 0.648   |
| Beta-blockers                         | 1.35 (0.58-3.15)      | 0.488   |
| Calcium channel blockers              | 0.86 (0.26-2.84)      | 0.806   |
| Statins                               | 2.29 (1.11-4.73)      | 0.025   |
| Diuretic agents                       | 0.97 (0.23-4.06)      | 0.964   |
| Oral anticoagulation                  | 0.88 (0.32-2.45)      | 0.898   |
| Insulin                               | 5.70 (1.70-19.08)     | 0.005   |
| Ferritin                              | 1.00 (1.00-1.00)      | 0.634   |
| CRS grade                             | 2.10 (1.47-2.98)      | <0.001  |
| 1                                     | 0.25 (0.03-1.86)      | 0.176   |
| 2                                     | 0.39 (0.17-0.91)      | 0.029   |
| 3                                     | 3.11 (1.55-7.09)      | 0.002   |
| 4                                     | 9.79 (3.96-24.23)     | <0.001  |
| LVEF                                  | 1.00 (0.96-1.05)      | 0.778   |
| Systolic blood pressure               | 0.99 (0.97-1.01)      | 0.346   |
| Diastolic blood pressure              | 0.95 (0.91-0.99)      | 0.007   |
| Heart rate                            | 1.00 (0.98-1.02)      | 0.868   |
| High                                  | 0.83 (0.69-0.99)      | 0.035   |
| WBC count                             | 1.00 (0.99-1.01)      | 0.619   |
| Platelet count                        | 0.99 (0.99-1.00)      | 0.027   |
| Creatinine                            | 4.30 (1.19-15.59)     | 0.026   |
| GFR                                   | 0.99 (0.98-1.00)      | 0.181   |
| Mitral E/e                             | 1.15 (1.00-3.1)       | 0.046   |
| PASP                                  | 1.05 (0.93-1.18)      | 0.150   |
| Diastolic dysfunction                 | 1.11 (0.32-3.85)      | 0.866   |
| Left atrial volume index              | 1.04 (1.00-1.08)      | 0.075   |
| Interventricular septal thickness     | 3.13 (0.19-50.91)     | 0.422   |

Cl = confidence interval; CRS = cytokine release syndrome; other abbreviations as in Tables 1 and 2.
The cardiovascular profile of the patients treated with CAR T cell cells was slightly worse compared with that noted in the general population with respect to the baseline rates of heart failure, hypertension, and coronary artery disease (20–23). According to age, <2% of persons 40 to 59 years of age in the general population are reported to have heart failure, compared with 3% in our study. The prevalence of coronary disease for those 40 to 59 years of age in the general population is about 6%, which is lower than in our study (10%) (22). The prevalence of hypertension is estimated at 32% in the general population, compared with 36% in our study (20,24). The prevalence of diagnosed diabetes mellitus is approximately 10%, slightly higher than in our study (9%) (21). The rate of current or previous smoking may appear elevated (41%) in our study. However, when taking into account current smokers only (12 of 145 patients [8%]), the prevalence of current smoking was less than the current rate of smoking in the US (14%) (25). The difference in proportion of previous heart failure or coronary disease could be attributed to prior cancer therapy (e.g., 60% anthracycline use, 23% radiation, 21% stem cell transplantation). Similarly, atrial fibrillation (8%) was more prevalent in our study than in the general population (<2% younger than 65 years) and may be explained by the use of ibrutinib (for CLL, diagnosed in 46% of our patients).

### Table 4

| Hazard Ratio (95% CI) | p Value |
|----------------------|---------|
| Statin 1.83 (0.88–3.81) | 0.105 |
| Creatinine 15.54 (3.67–65.86) | <0.001 |
| CRS grade 1 0.49 (0.06–3.74) | 0.489 |
| CRS grade 2 0.95 (0.33–2.71) | 0.917 |
| CRS grade 3 8.42 (3.48–20.40) | <0.001 |
| CRS grade 4 29.86 (9.80–90.94) | <0.001 |

Abbreviations as in Table 3.

As the median time to major adverse cardiovascular events (MACE) was 5 days later than the median time to cytokine release syndrome (CRS) onset, this suggests that CRS and its treatments can at least contribute to the occurrence of MACE in patients treated with anti-CD19 chimeric antigen receptor T cell therapy. IQR = interquartile range (25th to 75th quartile).
Cancer subtype does not seem to influence the risk for MACE (HR: 0.75; 95% CI: 0.44 to 1.28; p = 0.278), despite the different cardiovascular profiles of patients. Patients with CLL and DLBCL were older than patients with ALL and had increased cardiovascular risk, but their previous and current treatment may be less aggressive than that of patients with ALL. KM estimates for each cancer subtype and MACE, CRS, and survival were also performed at 30 days, 6 months, and 12 months (see Supplemental Figures 2 to 4).

In our study, 104 patients (72%) had at least 1 CRS episode, which is higher than the previously documented rate (55.3%) in a recent large systematic review and meta-analysis (6). We used the most current and broadest CRS consensus definition to identify the episodes (12), which may explain the increased detection of CRS.

The use of statins, insulin, and aspirin and higher baseline creatinine levels were each associated with MACE and likely reflect patients with a high cardiovascular risk profile and comorbidities pre-treatment. The analysis of baseline echocardiographic characteristics revealed that there was also a trend toward diastolic dysfunction and high left ventricular filling pressures and MACE. However, diastolic function categories as defined using the 2016 guidelines (14) were not associated with MACE; this result could be explained by the paucity of moderate or severe diastolic dysfunction and the relatively high frequency of indeterminate diastolic function.

Interestingly, baseline LVEF was not associated with MACE, underlining the importance of analyzing baseline diastolic function indexes (E/e’, left atrial volumes) in these patients. Although prior atrial fibrillation and the use of aspirin and insulin were associated with MACE, they were not included in further analysis, as the proportion of patients treated with aspirin and insulin or with previous atrial fibrillation was low.

The occurrence of grades 3 and 4 CRS was strongly associated with MACE. It is possible that CRS may result in depressed myocardial function, explaining the high rate of heart failure. It is also possible that patients with CRS receive large quantities of intravenous fluids that can worsen the volume overload state. The exact CRS treatment was left to the discretion of the treating physicians but usually comprises intensive fluid resuscitation followed by vasopressors. The median time to MACE was 5 days later than the median time to CRS onset, suggesting that CRS and its treatments may contribute to MACE in patients treated with CAR T cell therapy.

**STUDY LIMITATIONS.** The data were obtained retrospectively. As with all retrospective studies, MACE could have been misclassified because of reporting errors or loss to follow-up. However, each medical chart was carefully reviewed, and signs and symptoms of MACE were individually adjudicated. Also, our overall loss to follow-up was small (6 patients [4%]). To decrease loss to follow-up, every patient was researched using a national necrology database. Thus, it is unlikely that deaths were missed. CIs for many of our risk estimates were also wide, and as such, these results need to be confirmed in additional studies.

Given that patients were referred from around the world to receive CAR T cell therapy, prior anthracycline use could have been underreported, and it was not possible to determine the total radiation or anthracycline dose previously received. It is widely acknowledged that total radiation or anthracycline dose is directly associated with MACE; the association of anthracycline dose or radiation with MACE could have been underestimated.

**CLINICAL RELEVANCE.** As CAR T cell therapy use becomes more available and accessible, it is crucial to understand not only the benefits but the possible side effects and the time frame in which they can occur. In contrast to radiation or other potentially cardiotoxic chemotherapies, no guidelines are currently available for screening or surveillance of cardiac function of patients treated with CAR T cell infusion. Understanding the incidence and the natural history of treatment-induced side effects will allow better screening and follow-up for these patients. To this effect, an extensive monitoring program in the United Kingdom is under way (26).

**CONCLUSIONS**

To our knowledge, this is the largest retrospective study conducted in adult patients treated with CAR T cell therapy evaluating the occurrence of and risk factors associated with adverse cardiovascular events. Using multivariable Cox proportional cause-specific hazards regression analysis, we determined that baseline creatinine and grade 3 or 4 CRS were independently associated with MACE. CAR T cell therapy has opened a new field in the treatment of hematologic malignancies and is now under investigation for other indications, including solid tumors,
acute myeloid leukemia, and multiple myeloma (27–29). Given that CAR T cell use will only increase in the future and that no recommendations for monitoring and follow-up of left ventricular function and MACE currently exist (30), prospective studies are needed to ascertain the incidence and predictors of MACE.

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APPENDIX For supplemental tables and figures, please see the online version of this paper.