

CLINICAL TRIALS AND OBSERVATIONS

KIR-favorable TCR- $\alpha\beta$ /CD19-depleted haploidentical HCT in children with ALL/AML/MDS: primary analysis of the PTCTC ONC1401 trial

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KEY POINTS

- Two-year DFS/OS of children with leukemia/MDS (MRD <0.01%) were 79%/82%, with rates of transplant-related mortality and chronic GVHD <10%.
- Non-radiation containing reduced-toxicity approaches led to superior DFS/OS compared with traditional myeloablative approaches.

We performed a prospective multicenter study of T-cell receptor $\alpha\beta$ (TCR- $\alpha\beta$)/CD19-depleted haploidentical hematopoietic cell transplantation (HCT) in children with acute leukemia and myelodysplastic syndrome (MDS), to determine 1-year disease-free survival (DFS) and compare 2-year outcomes with recipients of other donor cell sources. Fifty-one patients aged 0.7 to 21 years were enrolled; donors were killer immunoglobulin-like receptor (KIR) favorable based on ligand mismatch and/or high B content. The 1-year DFS was 78%. Superior 2-year DFS and overall survival (OS) were noted in patients <10 years of age, those treated with reduced toxicity conditioning (RTC) rather than myeloablative conditioning, and children with minimal residual disease <0.01% before HCT. Multivariate analysis comparing the KIR-favorable haploidentical cohort with controls showed similar DFS and OS compared with other donor cell sources. Multivariate analysis also showed a marked decrease in the risk of grades 2 to 4 and 3 to 4 acute graft versus host disease (aGVHD), chronic GVHD, and transplant-related mortality vs other donor cell sources. Ethnic and racial minorities accounted for 53% of enrolled patients, and data from a large cohort of recipients/donors screened for KIR showed that >80% of recipients had a KIR-favorable donor by our definition, demonstrating that this approach is broadly applicable to groups often unable to find donors.

This prospective, multicenter study showed improved outcomes using TCR- $\alpha\beta$ /CD19-depleted haploidentical donors using RTC for children with acute leukemia and MDS. Randomized trials comparing this approach with matched unrelated donors are warranted. This trial was registered at <https://clinicaltrials.gov> as #NCT02646839.

Introduction

The importance of strategic mismatch of killer immunoglobulin-like receptors (KIRs) between donor and recipients of allogeneic hematopoietic cell transplantation (HCT) has been debated extensively after early investigations in haploidentical HCT of patients with acute myeloid leukemia (AML) showed that specific combinations of KIR molecules in donors and recipients decreased relapse and lowered transplant-related mortality (TRM).¹⁻³ Although the effect has been absent or equivocal in

some studies,⁴⁻⁶ several studies have shown benefits in both pediatric and adult recipients of cells from a variety of donor sources.⁷⁻¹⁰ Methods of choosing optimal donors based on KIR differences have focused on 2 broad approaches: (1) the absence of KIR inhibitory ligands in a recipient with the presence of the KIR inhibitory molecule in the donor, allowing for the natural killer (NK) activity against recipient cancer cells to proceed, and (2) the presence of higher levels of KIR molecules that license or increase the likelihood of NK-cell activation against recipient cells.

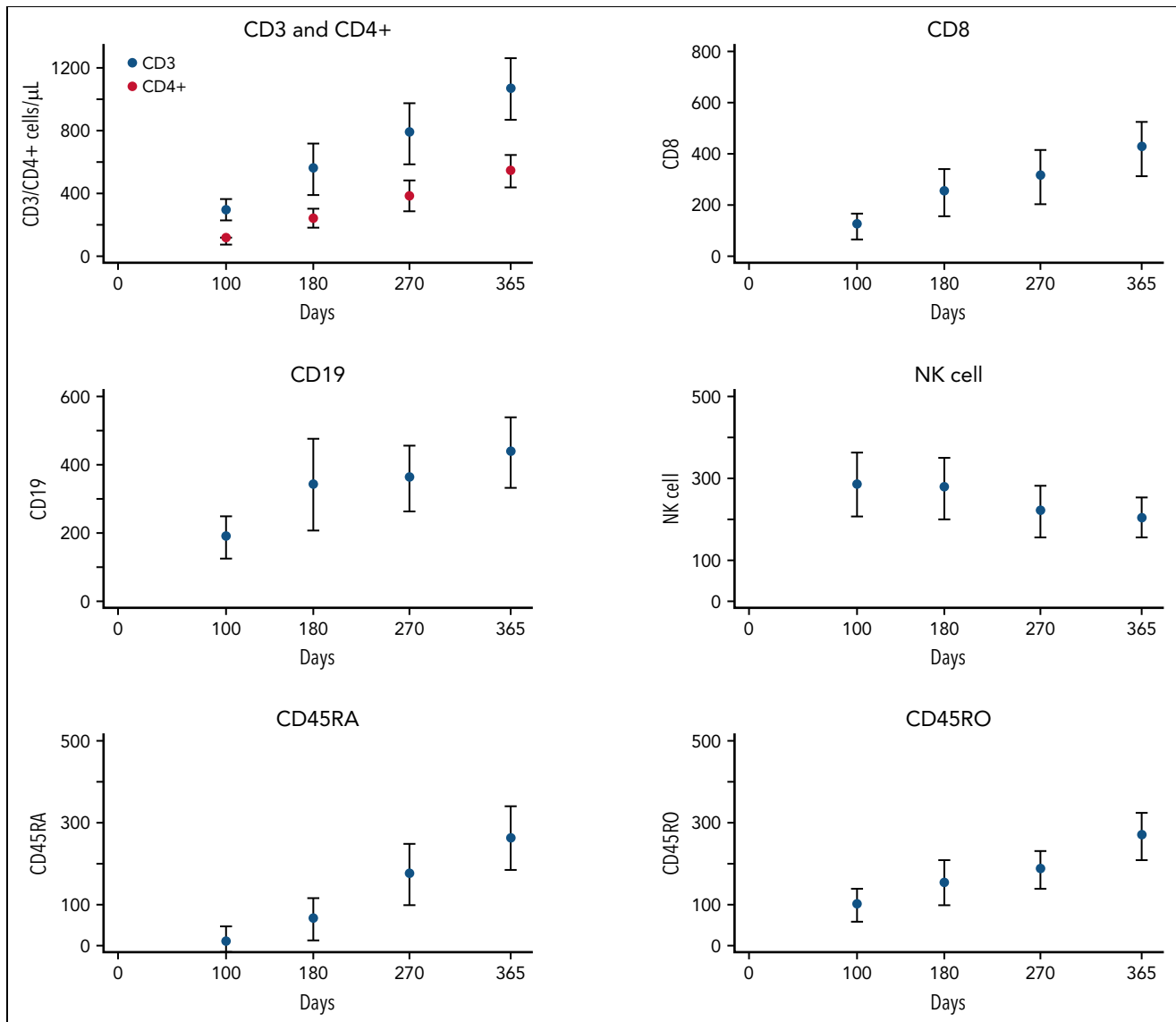


Figure 1. Lymphocyte recovery over the first year after transplant for CD3, CD4, CD8, CD19, NK cells, CD45RA, and CD45RO. The y-axes are expressed in cells per microliter, and the x-axes are days after HCT infusion.

The first category of KIR effect has been termed “missing ligand” and is routinely assessed by testing for the presence or absence of the C1, C2, and Bw4 inhibitory ligands on patients, while assessing donors for the presence of their inhibitory molecule binding partners 2DL2 and 2DL3, 2DL1, and 3DL1, respectively.^{11,12} There are many ways of assessing licensing of NK activity by KIR molecules. One example is a model put forward by Cooley who used a combination of types of KIR molecules (A vs B), as well as location of the KIR molecules (centromeric vs telomeric).⁹ They noted that in recipients of HCT for AML, those receiving donors with higher B content had the most protection against relapse. Donor/recipient missing ligand and donor B content according to Cooley score can be easily calculated (https://www.ebi.ac.uk/ipd/kir/donor_b_content.html) based on donor KIR typing and then used to choose donors for prospective trials.

The bulk of the HCT literature on KIR has focused on AML in adults, and KIR effects in adult patients with acute lymphocytic

leukemia (ALL) have usually been limited or absent.^{9,13} By contrast, multiple investigators have shown a beneficial effect of KIR mismatching in pediatric patients with ALL who undergo HCT,^{14,15} with a prominent role played by NK cells in the control of both diseases.^{16,17} Because a large portion of HCT for hematological malignancies in pediatrics occurs in patients with ALL, demonstration of a beneficial KIR effect in pediatric ALL would be important to the pediatric HCT field.¹⁸ Single-center work by Leung and colleagues at St Jude Children’s Hospital showed that when using KIR-favorable donors, survival outcomes with ex vivo T-cell-depleted haplotransplantation approached or exceeded those obtained using other donor types (matched sibling and unrelated).¹⁹ These outstanding outcomes in a pediatric population included all hematological malignancies, but a sizable percentage were transplanted for ALL.

With these preliminary data in hand, we hypothesized that in a multicenter setting, 1-year disease-free survival (DFS) after

Table 1. Patient demographics for both the phase 2 cohort and the CIBMTR comparator cohort

Characteristic	Phase 2 cohort (n = 51)	CIBMTR cohort (n = 1865)
Patients		
Median age at HCT, y (range)	11.73 (0.67-21.79)	12.24 (0.34-21.96)
Sex, n (%)		
Male	31 (61)	1098 (60)
Female	20 (39)	730 (40)
Donor, n (%)		
Median age, y (range)	35 (6-61)	23.4 (0-60)
Donor sex		
Male	29 (57)	1043 (57)
Female	22 (43)	769 (42)
Data missing	—	16 (1)
Donor/patient relations, n (%)		
Sex mismatch (F≥M)	12 (23.5)	446 (24.4)
Donor relationship, n (%)		
Parent; maternal	37 (72.5); 16 (43)	0
Sibling	12 (23.5)	527 (29)
Other related	2 (4)	0
Unrelated	0	886 (48)
CB	0	415 (23)
Disease status, n (%)		
ALL CR1	16 (31.4)	475 (26)
ALL CR2	14 (27.5)	487 (26.6)
AML CR1	11 (21.6)	617 (33.8)
AML CR2	5 (9.8)	249 (13.6)
MDS	5 (9.8)	0
Cell source		
Haplo (5/10, 6/10, 7/10, 8/10)	51	0
Matched sibling donor	0	527 (29)
MURD (9/10, 10/10)	0	670 (37)
MMURD (7/8)	0	216 (12)
CB 8/8	0	47 (3)
CB 7/8	—	75 (4)
CB ≤6/8	—	199 (11)
CB multidonor	—	82 (4)
CB missing	—	5 (<1)
CMV		
Patient		
Positive	27 (53)	1056 (58)
Negative	24 (47)	564 (31)
Data missing	0	208 (11)

Table 1 (continued)

Characteristic	Phase 2 cohort (n = 51)	CIBMTR cohort (n = 1865)
Donor		
Positive	27 (53)	734 (40)
Negative	24 (47)	886 (48)
Data missing	0	208 (11)
Conditioning regimens, n (%)		
MAC	27 (53)	1828
TBI	16 (31)	—
Busulfan	11 (22)	—
RTC	24 (47)	0
rATG/flu/mel/thio	22 (43)	—
TLI	2 (4)	—
NK alloreactivity		
Ligand mismatch (yes/no)	44/7	—
B-content value 0-1 vs ≥2	20/31	—
Race		
Black	5 (10)	141 (7.7)
Asian	9 (17.6)	151 (8.3)
White	24 (47)	1344 (73.5)
Other	13 (25.5)	192 (10.5)
Ethnicity		
Non-Hispanic	24 (47)	1196 (65.4)
Hispanic	23 (45.1)	554 (30.3)
Other	4 (7.8)	78 (4.3)
Follow-up		
Median follow-up, d (range)	609 (29-1772)	740 (42-1961)
Median cell dose infused/recipient weight (range) ×10⁶/kg		
CD34 ⁺ cells (n = 51)	12.19 (2.57-36.4)	—
TCR-αβ cells (n = 51)	0.02845 (0-0.418)	—
TCR-γδ cells (n = 51)	7.739 (0.1077-1810)	—
NK cells (n = 47)	41.13 (3.41-193.3)	—
CD20 ⁺ cells (n = 51)	0.06435 (0.007-2.756)	—

CMV cytomegalovirus; TLI, total lymphoid irradiation.

T-cell receptor-αβ (TCR-αβ)/CD19-depleted haploidentical HCT for children and young adults up to age 21 with complete response 1 (CR1)/CR2 ALL/AML/myelodysplastic syndrome (MDS) would be 80% and would compare favorably with other donor types if concomitant controls were used. This article describes primary outcomes for the ONC1401 trial run through the Pediatric Transplantation and Cellular Therapy Consortium (PTCTC).

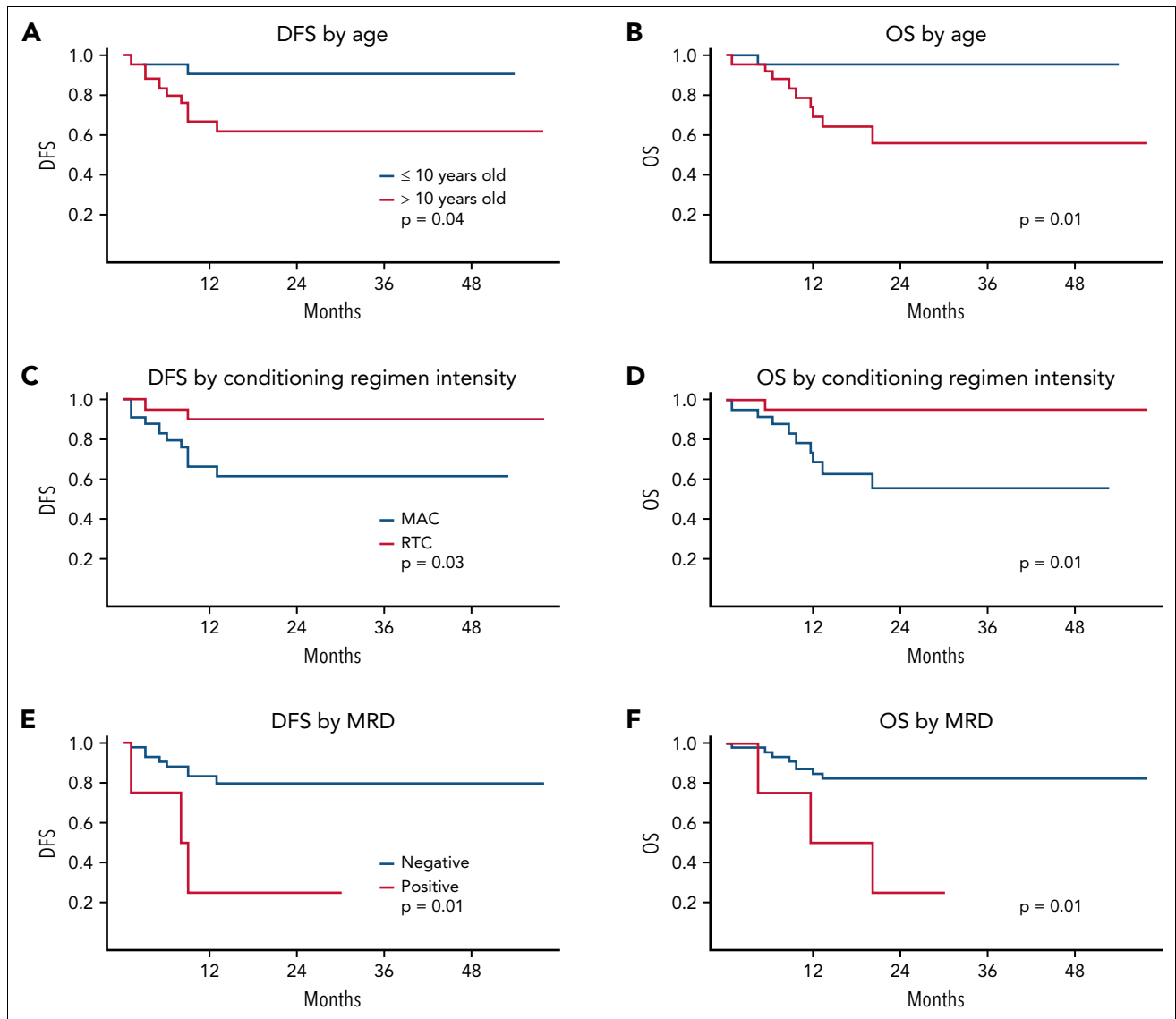


Figure 2. DFS and OS. Survival statistics by age (A-B), conditioning regimen (C-D), and MRD status (E-F).

Methods

PTCTC ONC1401 was a phase 2 trial conducted at 12 institutions of the PTCTC from January 2016 through July 2021. The study was approved by all local institutional review boards. Written informed consent was obtained from all parents and/or legal guardians in accordance with the Declaration of Helsinki.

Patient eligibility

Patients with ALL, AML, or MDS who were in CR1 or CR2 were eligible if they were <22 years of age with adequate organ function and had an eligible KIR-favorable haploidentical donor. The KIR-favorability assessment was performed in a single Clinical Laboratory Improvement Amendments–approved laboratory at Children’s Hospital of Philadelphia. KIR favorability was defined as B content according to the Cooley Scale of $\geq 2^9$ or inhibitory ligand mismatch by DNA typing or both.

Additional disease-specific criteria were as follows: for ALL, high-risk in CR1 eligible for HCT. Example CR1 indications:

induction failure (>5% blasts by morphology on postinduction bone marrow [BM]), minimal residual disease (MRD) by flow cytometry >0.01% after consolidation, hypodiploidy (<44 chromosomes), persistent or recurrent cytogenetic or molecular evidence of disease during therapy requiring additional therapy after induction to achieve remission (eg, persistent molecular BCR-ABL positivity). ALL in CR2 indications were B cell, early (≤ 36 months from initiation of therapy) BM relapse; late BM relapse with MRD >0.1% by flow cytometry after first induction; T-cell or Ph⁺ with BM relapse at any time; very early (<18 months from initiation of therapy) isolated extramedullary relapse (T cell or B cell). Indications for AML were high-risk AML defined as monosomy 5, del 5q, monosomy 7, M6, M7, t(6;9), FLT3-ITD, or patients who have $\geq 25\%$ blasts by morphology after induction, or failure to achieve CR after 2 courses of therapy. Also, patients with $\geq 0.1\%$ MRD or evidence of progressive extramedullary disease after induction chemotherapy. These patients achieved morphologic remission before transplant. Any patients with AML in CR2 were allowed. Indications for MDS: any 2001 World Health Organization classification

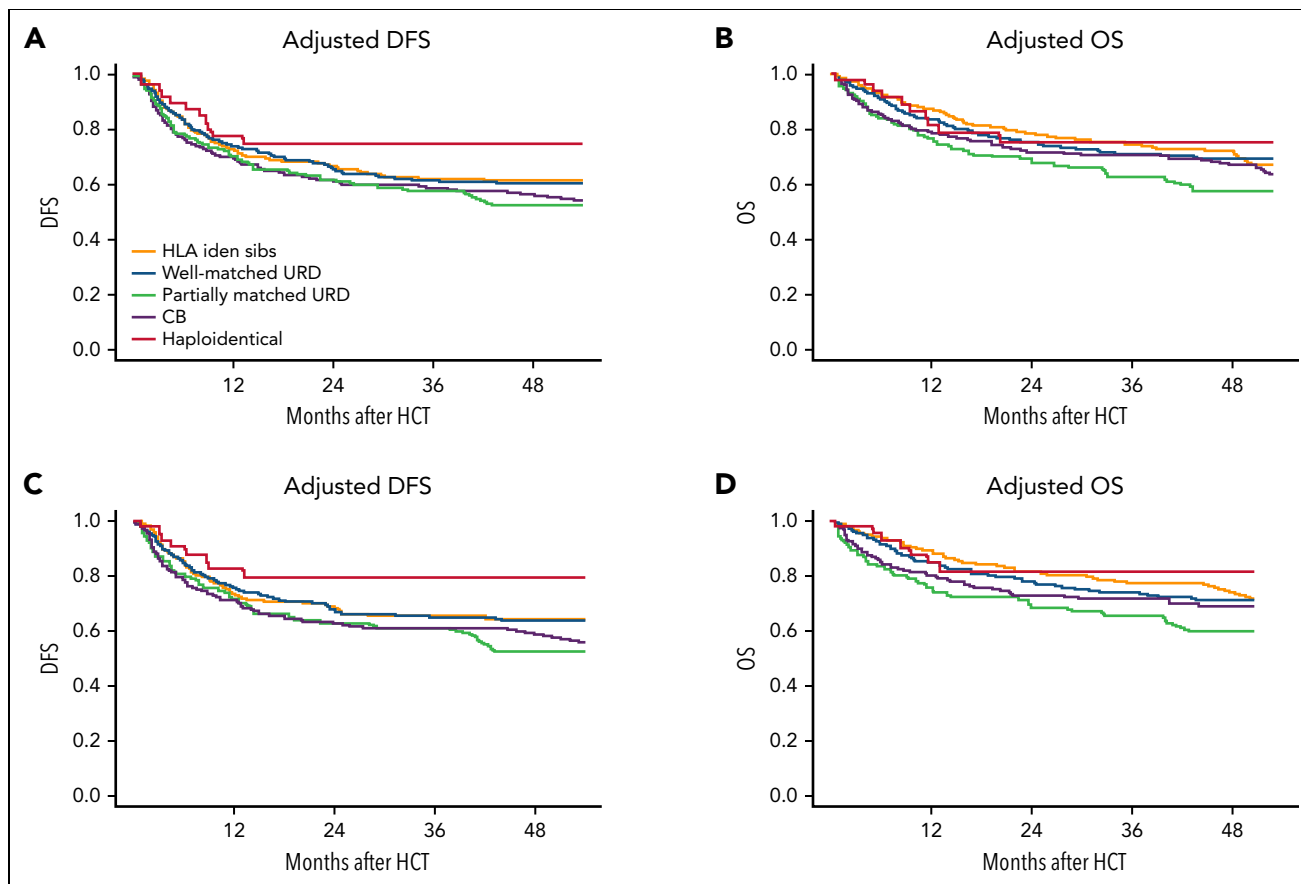


Figure 3. Adjusted DFS and OS. Adjusted statistics of the full phase 2 cohort vs CIBMTR cohorts (A-B) and the flow MRD negative before HCT phase 2 vs CIBMTR cohorts (C-D).

subtype was eligible (cytogenetic/molecular lesions of patients; supplemental Table 1, available on the *Blood* Web site); eligibility criteria (full study protocol) are included as supplemental Data.

Study procedures

Patients received a preparative regimen with either traditional myeloablative (MAC) or reduced toxicity (RTC) conditioning, less-intensive myeloablative approaches using agents associated with lower rates of toxicity. The 2 MAC regimen options included rabbit antithymocyte globulin (rATG; thymoglobulin) 3 mg/kg from day -12 to day -10 (total, 9 mg/kg), total body irradiation (TBI) 200 cGy twice daily from day -8 to day -6 (total, 1200 cGy; TBI-MAC), or busulfan from day -9 to day -6 (targeting an area under the curve of 60 to 95 mg/L per hour; Bu-MAC), with both regimens also including thiotepa (thio) 5 mg/kg on days -5 and -4 (total 10 mg/kg) and cyclophosphamide (cy) 60 mg/kg on days -3 and -2 (total 120 mg/kg). RTC regimen choices included (1) rATG 3 mg/kg from days -12 to -10 (total, 9 mg/kg), fludarabine (flu) 40 mg/m² from days -9 to day -5 (total, 200 mg/m²), thio 5 mg/kg twice daily on day -4 (total, 10 mg/m²), melphalan (mel) 70 mg/m² on days -3 and -2 (total, 140 mg/m²), or (2) total lymphoid irradiation 200 cGy twice daily on day -9, 200 cGy daily from day -8 to day -7 (total 800 cGy), flu 30 mg/m² from day -8 to day -4 (total 150 mg/m²), cy 60 mg/kg on day -6, thio 5 mg/kg twice daily on

day -4 (total, 10 mg/kg), and mel 70 mg/m² on days -3 and -2 (total 140 mg/m²). The choice of regimen was based on center, investigator, and patient preference.

All haploidentical grafts were peripheral blood stem cells (PBSCs) processed fresh using Miltenyi CliniMACS TCR- $\alpha\beta$ /CD19 depletion using infused graft cell processing targets as follows: CD34⁺ goal $\geq 10 \times 10^6$ /kg, TCR- $\alpha\beta$ ⁺CD3⁺ goal $< 1 \times 10^5$ /kg, CD19⁺ goal $< 1 \times 10^5$ /kg, with rituximab 375 mg/m² given on day +1 if CD19⁺ cells exceeded 1×10^5 /kg. CD34⁺ selection and a second infusion within a day was allowed to optimize the CD34⁺ cell dose. No pharmacological graft-versus-host disease (GVHD) prophylaxis was used. Granulocyte-colony stimulating factor (G-CSF) was used rarely, according to local practice, but only after days +14 to +21.

Statistical analysis

The primary objective of this phase 2 trial was to prospectively assess DFS (events: relapse and death) of KIR-favorable haploidentical HCT using ex vivo TCR $\alpha\beta$ ⁺CD3⁺/CD19⁺-depleted grafts in children with high-risk ALL, AML, and MDS. Secondary objectives included assessing rates of relapse, TRM, GVHD, event-free survival (EFS; events: relapse, death, and rejection), and overall survival (OS) and comparing these outcomes with concurrently enrolled children receiving myeloablative HCT from matched sibling, matched unrelated, 7 of 8 HLA-mismatched

Table 2. Combined analyses of the full and MRD-negative KIR cohorts with the full and MRD-negative CIBMTR cohorts for DFS/OS

Outcome	HR	95% CI, lower limit	95% CI, upper limit	P	Overall P	n
KIR cohort and CIBMTR cohort						
DFS						
Donor type						
Haploidentical	1.00	—	—	—	.1312	46
HLA-identical sibling	1.434	0.778	2.64	.2478		526
Well-matched URD	1.448	0.788	2.661	.2337		665
Partially matched URD	1.745	0.929	3.277	.0834		216
CB	1.728	0.937	3.189	.0799		412
Ethnicity						
Hispanic	1.00	—	—	—	.0305	569
Non-Hispanic	0.798	0.672	0.948	.0101		1215
Data missing	0.758	0.502	1.145	.1879		81
Disease						
AML	1.00	—	—	—	.0009	876
ALL	0.76	0.647	0.894	.0009		989
Disease status (≤8 mo)						
CR1	1.00	—	—	—	.0004	1114
CR2	1.426	1.171	1.737	.0004		751
Disease status (>8 mo)						
CR1	1.00	—	—	—	.0852	774
CR2	0.781	0.59	1.035	.0852		499
OS						
Donor type (≤4 mo)						
Haploidentical	1.00	—	—	—	.0001	46
HLA-identical sibling	2.024	0.274	14.975	.4897		527
Well-matched URD	2.67	—	19.485	.3327		670
Partially matched URD	4.783	—	35.521	.126		216
CB	5.567	0.769	40.279	.089		415
Donor type (>4 mo)						
Haploidentical	1.00	0.408	1.619	.556	.2449	45
HLA-identical sibling	0.813	0.463	1.813	.8015		491
Well-matched URD	0.916	0.585	2.439	.625		619
Partially matched URD	1.195	0.409	1.652	.5813		191
CB	0.822	0.409	1.652	.5813		356
Ethnicity						
Hispanic	1.00	—	—	—	.01	574
Non-Hispanic	0.756	0.618	0.925	.0066		1218
Data missing	0.58	0.34	0.989	.0454		82
Disease (≤6 mo)						
AML	1.00	—	—	—	.8471	882
ALL	1.03	0.765	1.385	.8471		992

Bold P-values indicate statistically significant results.

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Table 2 (continued)

Outcome	HR	95% CI, lower limit	95% CI, upper limit	P	Overall P	n
Disease (>6 mo)						
AML	1.00	—	—	—	<.0001	741
ALL	0.53	0.411	0.682	<.0001		824
Disease status (≤5 mo)						
CR1	1.00	—	—	—	.0003	1119
CR2	1.813	1.317	2.496	.0003		755
Disease status (>5 mo)						
CR1	1.00	—	—	—	.5978	1018
CR2	0.936	0.734	1.195	.5978		652
MRD⁻ KIR cohort and MRD⁻ CIBMTR cohort						
DFS						
Donor type					.1327	
Haploidentical	1.00	—	—	—		42
HLA-identical sibling	1.636	0.797	3.355	.1795		378
Well-matched URD	1.617	0.792	3.303	.1869		478
Partially matched URD	2.057	0.978	4.327	.0574		149
CB	1.995	0.971	4.098	.0601		298
Disease					.0126	
AML	1.00	—	—	—		609
ALL	0.782	0.645	0.949	.0126		736
OS						
Donor type (≤4 mo)					.0002	
Haploidentical	1.00	—	—	—		42
HLA-identical sibling	1.959	0.261	14.731	.5134		379
Well-matched URD	2.219	0.299	16.486	.4361		480
Partially matched URD	6.092	0.815	45.541	.0783		149
CB	5.046	0.69	36.886	.1108		300
Donor type (>4 mo)					.4106	
Haploidentical	1.00	—	—	—		41
HLA-identical sibling	0.87	0.371	2.043	.7497		354
Well-matched URD	1.15	0.497	2.658	.7441		443
Partially matched URD	1.374	0.564	3.344	.4841		126
CB	1.061	0.45	2.499	.8922		258
Ethnicity					.0314	
Hispanic	1.00	—	—	—		406
Non-Hispanic	0.728	0.567	0.935	.0129		886
Data missing	0.613	0.319	1.181	.1436		58
Disease (≤6 mo)						
AML	1.00	—	—	—	.4472	612
ALL	1.152	0.799	1.661	.4472		738

Bold P-values indicate statistically significant results.

Table 2 (continued)

Outcome	HR	95% CI, lower limit	95% CI, upper limit	P	Overall P	n
Disease (>6 mo)						
AML	1.00	—	—	—	<.0001	511
ALL	0.49	0.354	0.677	<.0001		611
Disease status (≤5 mo)						
CR1	1.00	—	—	—	.0081	784
CR2	1.686	1.145	2.482	.0081		566
Disease status (>5 mo)						
CR1	1.00	—	—	—	.0512	711
CR2	0.73	0.533	1.002	.0512		492

Bold P-values indicate statistically significant results.

unrelated, and cord blood cell sources. Of 1865 recipients in the CIBMTR (Center for International Blood and Marrow Transplant Research) cohort, MRD data were available on 1745 patients. Additional secondary objectives included assessment of outcomes by MRD status and identification of risk factors for poor outcome, including a comparison of preparative approaches.

Cumulative incidences of GVHD, relapse, and TRM were considered to accommodate for competing risks. Cox proportional hazard analysis for EFS, DFS, and OS and the proportional cause-specific hazards model for GVHD, relapse, and TRM were used to identify prognostic factors via forward stepwise selection. The proportional hazard assumption for each variable was examined by testing whether its coefficient was constant over time. Covariates with a $P < .05$ were considered significant. Interactions among significant covariates were also examined. The variables for the regression analysis include race, ethnicity, donor/recipient sex match, donor/recipient cytomegalovirus serostatus, disease type, disease status, MRD status, and performance status. Adjusted survival and cumulative incidences were calculated based on the final regression model for each outcome.^{9,10} Kaplan-Meier estimates and cumulative incidence were calculated for the analysis restricted to the clinical trial data. All statistical analyses were performed with SAS version 9.4 (SAS Institute, Cary, NC) and R version 4.0 (Vienna, Austria).

Results

Demographics of the KIR-favorable haploidentical and CIBMTR comparator cohorts

Table 1 shows details of the prospectively enrolled phase 2 KIR-favorable haploidentical cohort (KIR cohort) and the CIBMTR cohort. The median age of patients on the trial was 11.7 (range, 0.7-21.8) years, whereas the median donor age was 35 (6-61) years. Although most of the haploidentical donors were parents (maternal, 43%; paternal, 57%), 12 (24%) were siblings and 2 were other related donors. The majority (59%) of the patients underwent HCT for ALL with a fairly even mix of CR1 and CR2. Only 5 patients had MDS, all of whom had advanced MDS. Because of the low number of patients with MDS in the phase 2 cohort, we did not include them in the comparative analysis with the CIBMTR comparator cohort. The CIBMTR cohort included all transplants at US centers in patients <22 years of

age (median 12.2; range, 0.34-21.96) with ALL and AML, with HLA-identical sibling donors, well-matched (8 of 8 alleles) or partially matched (7 of 8 alleles) unrelated donors (URDs), or unrelated cord blood (UCB) during the period of the trial (2015 through 2020), and was limited to myeloablative procedures by CIBMTR consensus definition²⁰ for first HCT in CR1 or CR2.

The phase 2 cohort had substantially higher numbers of ethnic minorities compared with the CIBMTR cohort: non-White 53% vs 26.5% ($P < .0001$), Hispanic 45% vs 30% ($P = .02$; Table 1).

Engraftment and immune reconstitution/effect of rituximab

CD34⁺ cell doses were high in the KIR cohort, with a median 12×10^6 CD34⁺ cells/kg recipient weight. Neutrophil engraftment was rapid (median day +14; range, 8-19). The overall rate of rejection was low ($n = 4$; 7.8%). Rejection occurred mostly in myeloid patients (2 AML, 1 MDS, and 1 ALL) and all received the rATG/flu/thio/mel approach. Three of the patients were at high risk for rejection, 2 due to a prolonged period without chemotherapy before HCT (2 to 4 months), and 1 due to MDS with high-risk cytogenetics, no pre-HCT chemotherapy, and low cell dose (4.83×10^6 /kg CD34⁺/kg, below our target of 10×10^6 /kg). All were successfully engrafted after rescue HCT using a different donor (1 mismatched URD, 1 cord, and 2 haplo).

Immune reconstitution was notable for normalization of NK cell counts by day 100, with a mean of 282/μL (95% confidence interval [CI], 204-360) followed by stable NK counts through the first year. T- and B-cell counts were a mean CD3 and CD19 of 284/μL (95% CI, 284-354) and 184/μL (95% CI, 120-248) at day 100 and 554/μL (95% CI, 390-717) and 339/μL (95% CI, 202-476) at 6 months, with continued increases over the first year after HCT. CD45RA and CD45RO counts were a mean of 1/μL (95% CI, 0-44) and 95/μL (95% CI, 55-136) at day 100; 61/μL (95% CI, 17-103) and 150/μL (95% CI, 109-191) at 6 months; and 171/μL (95% CI, 128-214) and 183/μL (95% CI, 142-224) at 9 months after HCT (Figure 1).

An analysis of patients who received rituximab after infusion for CD20⁺ counts above 1×10^5 /kg in the depleted product (24 received/27 did not) showed no differences in time to cellular immune recovery or cessation of IVIg. GVHD and survival

Table 3. Comparative analyses of full and MRD-negative phase 2 with CIBMTR cohorts for GVHD, relapse, and TRM

Outcome	HR	95% CI, lower limit	95% CI, upper limit	P	Overall P	n
KIR cohort and CIBMTR cohort						
aGVHD 2-4						
Donor type					.0001	
Haploidentical	1.00	—	—	—		46
HLA-identical sibling	2.186	0.834	5.729	.1117		104
Well-matched URD	4.093	1.626	10.302	.0028		119
Partially matched URD	4.587	1.713	12.286	.0024		45
CB	4.907	2.00	12.039	.0005		254
aGVHD 3-4						
Donor type					.0062	
Haploidentical	1.00	—	—	—		46
HLA-identical sibling	2.633	0.316	21.95	.371		104
Well-matched URD	10.214	1.37	76.172	.0234		119
Partially matched URD	11.374	1.458	88.731	.0204		45
CB	8.129	1.11	59.526	.0391		254
Race						
White	1.00	—	—	—	.045	360
Black	2.087	1.206	3.61	.0086		79
Others	0.963	0.47	1.972	.9183		75
Data missing	0.834	0.33	2.109	.7008		54
Disease						
AML	1.00	—	—	—	.016	272
ALL	1.75	1.11	2.76	.016		296
cGVHD						
Donor type					<.0001	
Haploidentical	1.00	—	—	—		46
HLA-identical sibling	2.932	1.079	7.966	.0349		521
Well-matched URD	4.508	1.668	12.181	.003		659
Partially matched URD	5.108	1.862	14.01	.0015		212
CB	3.362	1.233	9.166	.0178		408
Ethnicity						
Hispanic	1.00	—	—	—	.0004	568
Non-Hispanic	0.657	0.53	0.814	.0001		1200
Data missing	0.627	0.377	1.041	.071		78
Race						
White	1.00	—	—	—	<.0001	1340
Black	2.115	1.567	2.856	<.0001		145
Others	1.633	1.21	2.204	.0013		159
Data missing	0.863	0.615	1.211	.3938		202

Shown are data for factors associated with GVHD, relapse, and TRM (in patients with ALL or AML). Bold P-values indicate statistically significant results. KPS, Karnofsky, performance score.

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Table 3 (continued)

Outcome	HR	95% CI, lower limit	95% CI, upper limit	P	Overall P	n
Relapse						
Donor type					.1447	
Haploidentical	1.00	—	—	—		46
HLA-identical sibling	1.794	0.84	3.835	.1313		526
Well-matched URD	1.519	0.712	3.241	.28		665
Partially matched URD	1.261	0.565	2.814	.5706		216
CB	1.738	0.808	3.737	.157		412
Disease					<.0001	
AML	1.00	—	—	—		876
ALL	0.626	0.519	0.755	<.0001		989
TRM					<.0001	
Donor type						
Haploidentical	1.00	—	—	—		46
HLA-identical sibling	0.749	0.263	2.135	.5885		526
Well-matched URD	1.223	0.441	3.393	.6995		665
Partially matched URD	2.93	1.047	8.2	.0406		216
CB	1.743	0.629	4.831	.2855		412
Ethnicity					.0223	
Hispanic	1.00	—	—	—		569
Non-Hispanic	0.658	0.483	0.897	.008		1215
Data missing	0.563	0.243	1.302	.1791		81
Disease					.0155	
AML	1.00	—	—	—		876
ALL	1.465	1.075	1.995	.0155		989
KPS					<.0001	
≥90%	1.00	—	—	—		1606
<90%	2.039	1.44	2.885	<.0001		259
MRD⁺ KIR cohort and MRD⁻ CIBMTR cohort						
Donor type					.0016	
Haploidentical	1.00	—	—	—		42
HLA-identical sibling	2.125	0.789	5.725	.1359		74
Well-matched URD	3.87	1.516	9.879	.0047		86
Partially matched URD	2.826	0.925	8.64	.0684		27
CB	4.485	1.812	11.099	.0012		177
Donor type					.004	
Haploidentical	1.00	—	—	—		42
HLA-identical sibling	1.817	0.189	17.477	.6052		74
Well-matched URD	12.588	1.688	93.848	.0135		86
Partially matched URD	7.675	0.856	68.855	.0687		27
CB	7.345	0.993	54.355	.0509		177

Shown are data for factors associated with GVHD, relapse, and TRM (in patients with ALL or AML). Bold P-values indicate statistically significant results. KPS, Karnofsky, performance score.

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Table 3 (continued)

Outcome	HR	95% CI, lower limit	95% CI, upper limit	P	Overall P	n
Disease						
AML	1.00	—	—	—	.0213	188
ALL	1.93	1.103	3.379	.0213		218
Donor type						
Haploidentical	1.00	—	—	—	.0002	42
HLA-identical sibling	2.525	0.922	6.919	.0716		374
Well-matched URD	4.224	1.556	11.469	.0047		471
Partially matched URD	4.617	1.66	12.844	.0034		147
CB	3.19	1.161	8.764	.0245		296
Ethnicity					.0013	
Hispanic	1.00	—	—	—		401
Non-Hispanic	0.647	0.499	0.839	.001		874
Data missing	0.455	0.229	0.905	.0247		55
Race						
White	1.00	—	—	—	.0001	965
Black	2.059	1.432	2.96	<.0001		104
Others	1.737	1.205	2.503	.0031		108
Data missing	1.015	0.692	1.487	.9405		153
Donor type						
Haploidentical	1.00	—	—	—	.0604	42
HLA-identical sibling	2.642	0.971	7.191	.0572		378
Well-matched URD	2.219	0.816	6.033	.1183		478
Partially matched URD	1.499	0.514	4.37	.4588		149
CB	2.506	0.914	6.869	.0741		298
Disease						
AML	1.00	—	—	—	<.0001	609
ALL	0.554	0.438	0.7	<.0001		736
Donor type						
Haploidentical	1.00	—	—	—	<.0001	42
HLA-identical sibling	0.575	0.193	1.71	.3195		378
Well-matched URD	1.022	0.363	2.873	.9677		478
Partially matched URD	3.105	1.097	8.794	.0329		149
CB	1.545	0.548	4.356	.4104		298
Disease						
AML	1.00	—	—	—	.0029	609
ALL	1.771	1.216	2.58	.0029		736
KPS						
≥90%	1.00	—	—	—	.0003	1164
<90%	2.141	1.419	3.231	.0003		181

Shown are data for factors associated with GVHD, relapse, and TRM (in patients with ALL or AML). Bold *P*-values indicate statistically significant results. KPS, Karnofsky, performance score.

outcomes were similarly not affected by rituximab infusion. One patient who did not receive rituximab prophylactically developed CNS posttransplant lymphoproliferative disorder that required rituximab, local radiation therapy, and Epstein-Barre virus-specific T-cells to control.

Survival in the KIR cohort

One-year DFS in the KIR cohort was 78% (95% CI, 66% to 89%), with a 2-year EFS, DFS, and OS of 69% (95% CI, 56% to 82%), 75% (95% CI, 63% to 88%), and 75% (95% CI, 63% to 88%), respectively. Univariate risk analysis showed that several factors were predictive of survival. Age ≤ 10 years, RTC vs MAC, and flow MRD $< 0.01\%$ resulted in significant improvements in survival ($P = .03$, $.03$, and $.02$, respectively; [Figure 2](#)). The protocol included 2 traditional MAC approaches and 2 RTC approaches to accommodate patients not eligible for traditional MAC; however, because the RTC approaches are myeloablative and had been the preferred approaches in the St Jude single-center data,¹⁹ many centers chose RTC over traditional MAC for patients eligible for both. The higher rate of failure in the traditional MAC regimens was a mixture of both relapse and TRM: 5 of 27 (18.5%; 95% CI, 6% to 38%) relapsed and 5 of 27 (18.5%; 95% CI, 6% to 38%) experienced TRM with MAC vs 3 of 24 (12.5%; 95% CI, 3% to 32%) experiencing relapse and 0% (95% CI, 0% to 14%) TRM with RTC approaches.

Although half of the flow MRD-positive patients were treated with the MAC and half with the RTC regimens, MRD-positive patients were rare ($n = 6$; 12%) and, as anticipated, did poorly with both approaches (4 of 6 relapsed, 3 of 6 died; [Figure 2E-F](#)). To remove possible confounding of the imbalance of high-risk MRD-positive patients to the of MAC vs RTC outcomes comparison, we performed a subanalysis of outcomes in patients who were flow MRD $< 0.01\%$ before HCT. Notably, 2-year DFS and OS for patients were 79% (95% CI, 66% to 92%) and 82% (95% CI, 69% to 94%), respectively ([Figure 3C-D](#)). Superior survival outcomes for RTC vs MAC held in the MRD-negative cohort, with 2-year DFS of 90% (95% CI, 77% to 100%) vs 60% (95% CI, 40% to 81%), and OS of 95% (95% CI, 86% to 100%) and 55% (95% CI, 32% to 79%) with RTC vs MAC, respectively ($P = .03$ for both). ALL patients were our largest population; 16 patients with ALL were treated with a TBI regimen and 14 non-TBI, all flow MRD $< 0.01\%$ at HCT. Treatment failed in 3 of the 16 patients conditioned with TBI (19%: 2 relapsed, 1 TRM) and in 2 of the 14 not treated with TBI (14%: 1 relapse, 1 TRM).

Analysis of survival outcomes in the KIR and CIBMTR cohorts

Multivariable analyses for survival end points comparing the KIR cohort with cell sources from the CIBMTR cohort is presented in [Table 2](#). A subanalysis of patients in both cohorts coming to HCT with flow MRD $< 0.01\%$ is also included in [Table 2](#). There was a trend toward improved DFS for KIR cohort recipients compared with recipients of partially matched unrelated donors (URDs) and UCB, which strengthened in the MRD-negative analysis (partially matched URD hazard ratio [HR], 2.06; 95% CI, 0.98-4.3; $P = .057$) and UCB HR 2.0 (95% CI, 0.97-4.1; $P = .06$). OS was improved the first 4 months after HCT compared with other donor cell sources in both the overall and

MRD-negative cohort analyses ($P = .0001$ and $.0002$, respectively), but was similar after 4 months. [Figure 3](#) shows the DFS and OS comparisons between the KIR cohort and recipients of other donor cell sources for both the full cohorts and the pre-HCT MRD-negative cohorts.

Other factors notable for being independently associated with improved DFS include disease status (CR1; $P = .0004$), presence of ALL vs AML ($P = .0009$), and non-Hispanic ethnicity ($P = .03$).

Analysis of other significant outcomes: acute GVHD, chronic GVHD, relapse, and TRM

[Table 3](#) shows multivariate analyses of the combined KIR and CIBMTR overall and MRD-negative cohorts for acute GVHD (aGVHD) grades 2 to 4, severe aGVHD grades 3 and 4, chronic GVHD (cGVHD), relapse, and TRM. Notably, there was a dramatic decrease in the occurrence of significant acute (grades 2-4), severe acute (grades 3-4), and cGVHD (overall $P = .0001$, $.0062$, and $< .0001$, respectively) in recipients of TCR- $\alpha\beta$ /CD19-depleted haploidentical HCT. Aside from comparisons of aGVHD with matched sibling donors, where the advantage is not definitive, HRs of 4- to 10-fold were noted in recipients of URD and UCB HCT. There was also a clear advantage for KIR cohort recipients in TRM compared with 7 of 8 mismatched URD recipients. Importantly, there was no difference noted in risk of relapse in the KIR cohort. [Figure 4](#) shows the CI of grades 2 to 4 aGVHD, cGVHD, relapse, and TRM comparisons between the KIR cohort (red line) and recipients of other donor cell sources for both the full cohorts and the pre-HCT MRD-negative cohorts.

Other notable independent risk factors for aGVHD (grades 3-4) include Black race ($P = .0086$) and ALL ($P = .016$). For cGVHD, risks included Hispanic ethnicity ($P = .0004$) and Black ($P < .0001$) or other race ($P = .0013$). For relapse, the other independent risk was having AML ($P < .0001$). For TRM, other risks included Hispanic ethnicity ($P = .008$), having ALL ($P = .016$), or having a low Karnofsky performance score ($P < .0001$).

Analysis of KIR favorability in a population of haploidentical transplant candidates in 2 models

We analyzed 294 pediatric patients and 646 possible donors (2.2 donors/patient; range, 1-8; standard deviation; 0.975; a superset of the 51 cases included in this report) to determine the likelihood of identifying a KIR-favorable donor by our definition. A total of 196 of 294 (66.7%) patients had ≥ 1 absent ligand and 188 of 196 (96%) of those had a favorable ligand-mismatched donor identified ([Table 4](#)). We evaluated the KIR haplotype B content of the potential donors and found that 193 of 646 (29.9%) had a favorable B content score⁹ of ≥ 2 ([Table 5](#)). However, when all 646 potential donors were considered in the context of the 294 patients, just under half (142 of 294; 48.3%) of the patients had at least 1 potential donor with a Cooley B-content score of ≥ 2 , because multiple donors were screened for a given patient ([Table 6](#)). Notably, 236 of 294 (80.3%) had either a potential donor with a favorable ligand mismatch or high B content (matching our eligibility criteria), and 91 of 294 (31.0%) of patients had both favorable mismatch(es) and high B content ([Table 4](#)).

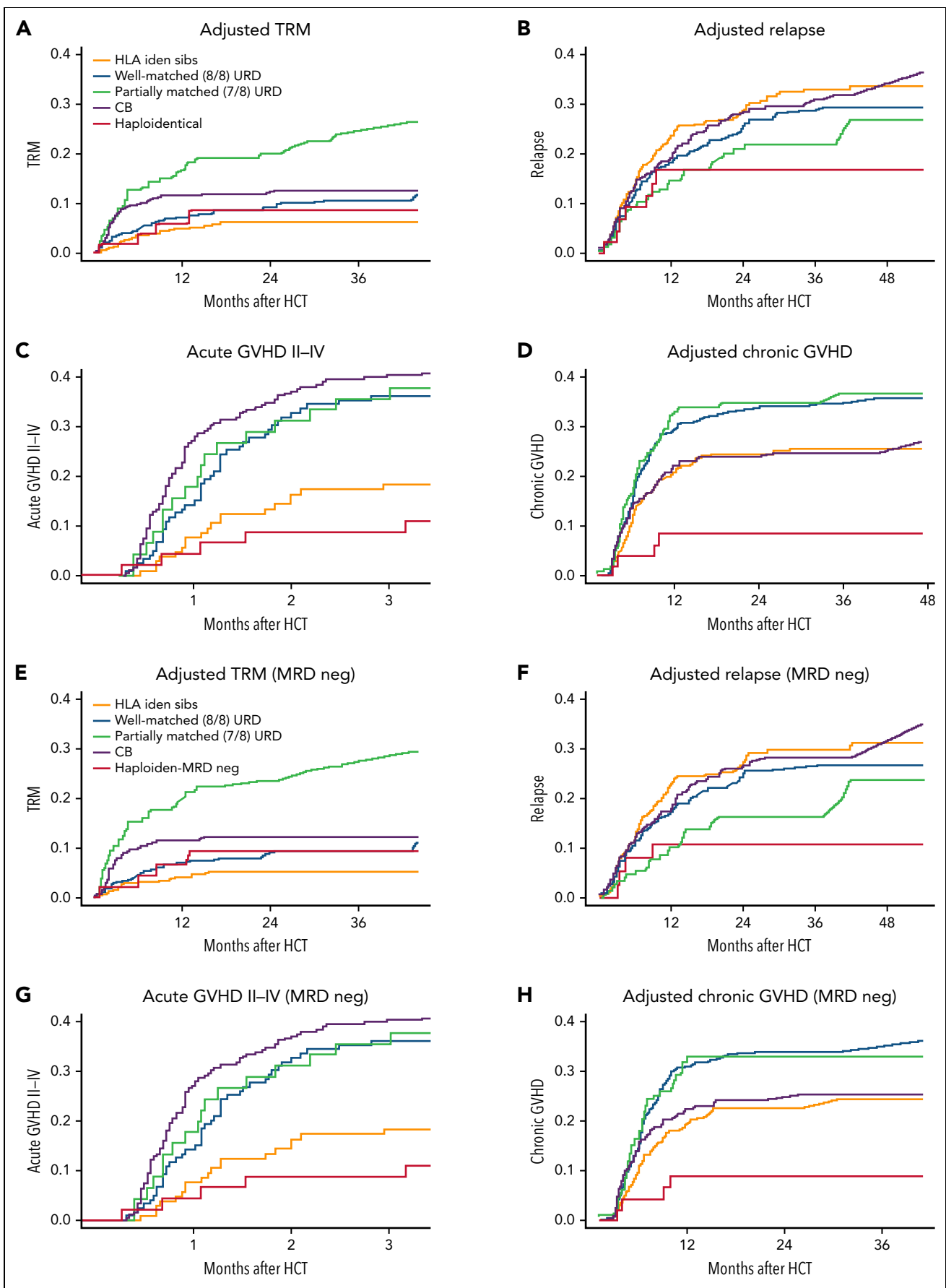


Figure 4. Comparative adjusted statistics for the study cohorts. TRM (A), relapse (B), acute GVHD grades 2 to 4 (C), and chronic GVHD (D) for all phase 2 and CIBMTR comparator patients. Adjusted TRM (E), relapse (F), aGVHD grades 2 to 4 (G), and cGVHD (H) for all phase 2 and CIBMTR comparator patients who were MRD-negative at the time of HCT.

Table 4. Results of KIR analysis for ligand mismatch and B content of 646 donors screened for 294 pediatric and adolescent and young adult recipients

	Patients screened n (%)	Missing at least 1 ligand n (%)	Missing C1 ligand n (%)	Missing C2 ligand n (%)	Missing Bw4 ligand n (%)
Possible mismatches (absent ligand in recipient)	294 (100)	196 (66.7)	51/196 (26.0)	111/196 (56.6)	74/196 (37.8)
Recipients with missing ligands who had donors available	—	188/196 (95.9)	49/51 (96.1)	107/111 (96.4)	68/74 (91.9)
Recipients with ≥ 1 mismatched donor at a given ligand	—	188/294 (63.9)	49/294 (16.7)	107/294 (36.4)	68/294 (23.1)
Recipients with a donor with ligand mismatch and Cooley score ≥2	91/294 (31.0)	—	—	—	—
Recipients with a donor that had ligand mismatch or Cooley score ≥2 or both	236/294 (80.3)	—	—	—	—

Discussion

Although outcomes for hematological malignancies in pediatrics have improved over time,^{21,22} allogeneic HCT continues to play an important role in salvage of high-risk and relapsed patients. Treatment with HCT after obtaining remission after first relapse of high-risk B-ALL and T-ALL and continues to be a standard.²³⁻²⁵ Chimeric antigen receptor (CAR) T-cell therapies have greatly improved outcomes in patients with B-ALL who have multiple relapses, but the therapy will fail in more than half of patients, leading eventually to a transplant,²⁶ and some argue that patients achieving remission with CAR T-cell therapy could benefit from consolidative HCT, either for all patients who have not had a previous HCT²⁷ or for patients with characteristics that put them at high risk of relapse.^{28,29} AML outcomes have shown, at best, modest improvement in the past few decades, and HCT for high-risk and relapsed disease is standard.³⁰ With a continued need for allogeneic HCT in children, URDs are often challenging to find, especially for minority patients,³¹ and COVID restrictions have placed a strain on donor acquisition, often forcing the use of PBSCs, which have been shown to lead to inferior outcomes in pediatric transplantation.^{32,33} With these concerns in mind, if haploidentical HCT approaches could be shown to be superior or, at

minimum, equivalent to URD approaches, it would be of major benefit to the pediatric HCT field. Our study addresses this need and is remarkable in that we achieved outstanding results, even with the preponderance of our patients being Hispanic or Black, demonstrating that this approach can be used for diverse populations.

In this prospective, multicenter trial we demonstrated that using an approach of KIR-favorable TCR- $\alpha\beta^+$ CD3⁺/CD19⁻-depleted haploidentical HCT, a 1-year DFS of ~80% could be realized and that DFS and OS were similar to CIBMTR outcomes when other cell sources were used. Furthermore, moderate and severe aGVHD, cGVHD, and TRM outcomes were clearly superior with our approach, and relapse risk was similar. The marked decrease in GVHD with maintenance of good outcomes was most likely the result of early licensed NK cell and/or $\gamma\delta$ T-cell activity. Our regimens involve administering rATG more distal to the graft infusion (starting at day -12), compared with other published TCR $\alpha\beta^+$ CD3⁺/CD19⁻-depleted approaches,³⁴ allowing less exposure of the infused graft to further depletion by rATG.³⁵ Our approach also uses no posttransplant immune suppression and results in rapid engraftment (median, day +14), which simplifies care and facilitates adding post-HCT cellular

Table 5. Number of potential donors with a particular B-content score from the screening cohort

B-content score	Donors, n (%)
0	213 (33.0)
1	240 (37.2)
2	145 (22.4)
3	36 (5.6)
4	12 (1.8)
Total	646 (100)

Table 6. Number of patients from the screening cohort that have a potential donor with minimum B-content score

Minimum B-content score for potential donor	Patients, n (%)
≥ 0	294 (100)
≥ 1	246 (83.6)
≥ 2	142 (48.3)
≥ 3	41 (14.1)
4	11 (4.8)

Screening cohort n = 294; potential donors n = 646.

therapies. It is notable that patients with pre-HCT by flow cytometry <0.01% had 2-year DFS and OS survival of 79% and 82%, respectively.

A challenge with haploidentical HCT has been high rates of rejection, and this has resulted in the use of intense approaches including the addition of thiotepa or melphalan to TBI or busulfan³⁶ (in our study, traditional MAC options were TBI/thio/cy and bu/thio/cy). In this trial, rejection was rare (7.8%), occurring mostly in patients at risk for rejection through MDS or prolonged periods (2-3 months) without chemotherapy who received our rATG/flu/thio/mel approach. Although all of our rejections were rescued with salvage HCT, for patients at high risk for rejection, clinicians could consider use of our total lymphoid irradiation-based RTC regimen or more traditional myeloablative approaches. Our data were striking, however, in that they indicate that, for most patients, more intense MAC approaches may be unnecessary and possibly detrimental, as a subanalysis of patients receiving RTC vs traditional MAC approaches on our study showed superior DFS and OS with the RTC approaches. This trial did not have a sufficient number of patients with MDS to draw conclusions regarding preparative approaches for the MDS population. Another provocative finding in this study is that outcomes of patients with ALL treated with TBI vs non-TBI approaches were very similar (16 TBI-conditioned patients: DFS, 81%; 1 NRM, 2 relapse; 14 non-TBI-treated patients: DFS, 86%; 1 NRM, 1 relapse). These results differ from the recent affirmation by Peters et al²³ of the superiority of TBI compared with treosulfan or busulfan when using matched sibling donors and URDs, but because our sample was small, a higher number of patients would be needed to confirm whether TBI could be omitted with this KIR-favorable haploidentical approach in MRD-negative patients with ALL, as studies addressing this issue to date have not been designed to answer this question.^{36,37}

With the outstanding outcomes noted using our approach in a multicenter trial setting, the question arises of how this approach using TCR $\alpha\beta^+CD3^+/CD19^-$ -depleted KIR-favorable haploidentical donors compares with haploidentical HCT with posttransplant cy. Studies in children and younger patients treated with MAC approaches have shown EFS ranging from 55% to 79% with OS of 72% to 85%.³⁸⁻⁴⁰ Rates of acute GVHD have been comparable, but rates of cGVHD have varied by stem cell source (higher with PBSC use) and have in general been higher than we report herein. Our TCR $\alpha\beta^+CD3^+/CD19^-$ -depletion approach is designed to use PBSCs, and generally allows high doses of CD34⁺ cells to be given, facilitating rapid engraftment with low rates of acute and chronic GVHD. Despite a modest cost for cell processing, with early discharge and low rates of aGVHD and cGVHD, costs may be lower than those for standard approaches. A planned prospective cost analysis of our study is underway. With these issues in mind, it is unclear whether TCR $\alpha\beta^+CD3^+/CD19^-$ -depleted haplo-HCT is better or worse than posttransplant cy, and prospective trials are needed to answer this question. Even though our data are very promising, it is unclear whether this approach could be better than 10 of 10 or 12 of 12 matched URD HCT. A randomized, prospective trial comparing outcomes of recipients of haploidentical donors with recipients of matched URDs is the only way to answer this question. Our study was planned

to lay the groundwork for such a trial and has provided key background data for Children's Oncology Group study ASCT2031, which will randomize haploidentical approaches with fully matched URDs (approved, opening in the fall of 2022).

In summary, our prospective, multicenter trial demonstrated that the use of KIR-favorable TCR $\alpha\beta^+CD3^+/CD19^-$ -depleted haploidentical HCT in children with hematological malignancies leads to rates of survival that meet or exceed those of other key donor cell sources, with rates of GVHD and TRM that are lower and rates of relapse that are similar. This approach is an excellent option for patients, especially if matched sibling or URD options are not available, given our analysis showing that 80.3% of screened recipients had potential donors with a favorable ligand mismatch or high B content. This means the approach is readily available to minority populations (45% of patients on the trial were of Hispanic ethnicity and 53% were Black, Asian, or other race) who otherwise have limited donor options.

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Authorship

Contribution: M.A.P., W.L., H.A.-A., D.M., and K.W.A., conceived and designed the study; M.A.P., N.J.B., N.L., E.A., A.F., M.S.C., J.-A.T., S.C., C.L.K., J.L.D., D.M., C.C.D., and H.A.-A. provided study material or patients; M.A.P., K.W.A., N.J.B., N.L., E.A., A.F., J.-A.T., S.C., C.L.K., J.L.D., D.M., C.C.D., and H.A.-A. collected and assembled the data; M.A.P., K.W.A., D.M., C.C.D., and H.A.-A. analyzed and interpreted the data; M.A.P., K.W.A., D.M., C.C.D., and H.A.-A. wrote the manuscript; and all authors provided final approval of the manuscript.

Conflict-of-interest disclosure: M.A.P. has served on advisory boards for Novartis, Mesoblast, Equillum, Medexus, and Vertex; has engaged in educational activities for Novartis; and has received study support from Miltenyi (not for this trial) and Adaptive. W.L. is an employee of Miltenyi. C.C.D. has been a consultant to and served on advisory boards of Jazz Pharmaceuticals, Alexion Inc, and Omeros Corporation. J.L.D. receives royalties from Omixon. D.M. is a consultant to, owns stock options in, and receives royalties from Omixon. S.C. has served on advisory boards of AbbVie, Viacord, and Alexion. H.A.-A. received study support from Adaptive. The remaining authors declare no competing financial interests.

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Footnotes

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Deidentified individual participant data that underlie the reported results will be made available 3 months after publication for a period of 5 years after the publication date. Proposals for access should be sent to michael.pulsipher@hci.utah.edu.

The online version of this article contains a data supplement.

There is a [Blood Commentary](#) on this article in this issue.

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