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Regular Article

TRANSPLANTATION

High-resolution HLA matching in hematopoietic stem cell transplantation: a retrospective collaborative analysis

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Key Points

- HLA mismatches at the allele and antigen level (possibly with the exception of HLA-DQB1) should be treated equally in donor selection.
- HLA mismatches at >1 locus (including HLA-DQB1) have additive detrimental effects.

To validate current donor selection strategies based on previous international studies, we retrospectively analyzed 2646 transplantations performed for hematologic malignancies in 28 German transplant centers. Donors and recipients were high resolution typed for HLA-A, -B, -C, -DRB1, and -DQB1. The highest mortality in overall survival analysis was seen for HLA-A, -B, and DRB1 mismatches. HLA-DQB1 mismatched cases showed a trend toward higher mortality, mostly due to HLA-DQB1 antigen disparities. HLA incompatibilities at >1 locus showed additive detrimental effects. HLA mismatching had no significant effect on relapse incidence and primary graft failure. Graft source had no impact on survival end points, neither in univariate nor in multivariate analysis. Higher patient age, advanced disease, transplantations before 2004, patient C2C2 killer cell immunoglobulin-like receptor (KIR)-ligand phenotype, and unavailability of a national donor adversely influenced outcomes in multivariate analysis. Our study confirms the

association of HLA-A, -B, -C, and -DRB1 incompatibilities with adverse outcome in hematopoietic stem cell transplantation (HSCT). The relevance of HLA-DQB1 disparities in single mismatched transplantations remains unclear. Similar hazard ratios for allele and antigen mismatches (possibly with an exception for HLA-DQB1) highlight the importance of allele level typing and matching in HSCT. The number of incompatibilities and their type significantly impact survival. (*Blood.* 2013;122(18):3220-3229)

Introduction

Hematopoietic stem cell transplantation (HSCT) from unrelated volunteer donors is the most widely used treatment alternative when no suitable sibling donor is identified. HSCT is potentially curative for a variety of malignant diseases of the hematopoietic system and certain life-threatening nonmalignant conditions. Transplantation of hematopoietic stem cells is a field of constant evolution and change. Major improvements have been the introduction of reduced intensity conditioning regimes (RIC), which decrease toxicity, and as a consequence transplant-related mortality (TRM), as well as the introduction of high-resolution HLA typing for donor selection.^{1,2} As HSCT becomes a safer therapy for many patients, increasing numbers of transplantations are performed worldwide. In this context, 3 main trends can be observed over the last decade: (1) the proportion of older transplant patients increases;

(2) peripheral blood stem cells (PBSCs) are presently the main graft source; and (3) the demand for unrelated donors is increasing.^{3,4}

Notwithstanding these changes, the paramount criterion for donor selection has remained HLA matching and, despite some variation in detail, a complete match at all relevant loci is unanimously recommended. However, the diversity of HLA remains a challenge in donor search and selection, sometimes forcing acceptance of mismatched donors. The aim of our study was to validate current donor selection strategies based on previous international studies and to investigate whether allelic HLA mismatches differentially impact outcome of HSCT compared with antigen mismatches at individual loci. As killer cell immunoglobulin-like receptor (KIR)-ligand interactions were reported to influence HSCT outcome, these were considered as well.^{5,6}

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HIGH-RESOLUTION HLA-MATCHING IN HSCT 3221

Table 1. Patient characteristics

Category	n	Percent
Number of patients	2646	100
Number of centers	28	
Age category		
18-29	333	12.6
30-39	348	13.2
40-49	554	20.9
50-59	770	29.1
60-69	577	21.8
70-79	64	2.4
Diagnosis		
AML	810	30.6
ALL	350	13.2
AL	135	5.1
CML	211	8.0
MDS	486	18.4
NHL	437	16.5
MM	217	8.2
Ethnicity		
White	2636	99.6
Asian	7	0.3
African	3	0.1
Disease stage		
Early	1047	39.6
Intermediate	922	34.8
Advanced	677	25.6
Conditioning regimen		
Myeloablative	1692	63.9
Reduced intensity	954	36.1
Stem cell source		
BM	346	13.1
PBSC	2300	86.9
Donor-recipient sex match		
Male-male	1227	46.4
Male-female	664	25.1
Female-male	362	13.7
Female-female	393	14.9
Year of transplanation		
1997-2000	299	11.3
2001-2005	589	22.3
2006-2010	1758	66.4

AML, acute myeloid leukemia; ALL, acute lymphoblastic leukemia; AL, unclassified acute leukemia; CML, chronic myeloid leukemia; MDS, myelodysplastic syndrome; NHL, non-Hodgkin lymphoma; MM, multiple myeloma; BM, bone marrow.

Patients and methods

Patients

Adult patients diagnosed with a malignant hematological disorder and for whom the unrelated donor search was performed in the search unit in Ulm were included in this analysis. All patients received a first allogeneic transplant using bone marrow or PBSCs without T-cell depletion between 1997 and 2010 (Table 1). Retransplanted patients were censored at the time of second transplantation. Disease stage definitions were adopted from a previous report of the European Group for Blood and Marrow Transplantation (EBMT) study group defining the EBMT risk score for outcome after HSCT.⁷ Myeloablative conditioning was defined as total body irradiation ≥ 10 Gy and/or cyclophosphamide ≥ 120 mg/m² and/or busulfan ≥ 16 mg/kg.^{8,9} Patients treated with less intense conditioning regimens were considered RIC. All patients received T cell–replete grafts. Patient and donor consents for HLA typing and for the analysis of clinical data were obtained in accordance with the Declaration of Helsinki. The study was approved by the ethical review board of the University of Ulm (project 263/09).

HLA typing

HLA typing of donors and patients carried out after May 2005 required testing for HLA-A, -B, -C, -DRB1, and -DQB1 at high resolution (n = 1804; 68.2%). Confirmatory typings for patients and donors from searches completed before May 2005 (n = 842; 31.8%) included low-resolution typing for HLA-A and -B and high-resolution typing for HLA-DRB1 and -DQB1. In these cases, HLA class I high-resolution typing was carried out retrospectively.

Exons 2 and 3 were analyzed for HLA class I typing, and exon 2 only for HLA class II typing. High-resolution HLA typing was performed by sequencing analysis (sequence-based typing) for HLA class I, and by the polymerase chain reaction (PCR)-sequence-specific oligonucleotide probes method or sequence-based typing for HLA class II. Ambiguities involving the most frequent null (not expressed) alleles were resolved if necessary by the PCR-sequence-specific primer technique.¹⁰ Like in studies performed by the Center for International Blood and Marrow Transplant Research (CIBMTR),¹¹⁻¹³ differences of the first 2 digits were classified as antigen mismatches irrespective of their vector¹⁴; other differences were classified as allele mismatches with some exceptions, which, as in the CIBMTR studies, were grouped according to their serologic reactivity.¹⁵

To better characterize patients without a suitable donor in the national registry, patients with rare HLA phenotypes were identified using high-resolution 5-locus HLA-haplotypes, derived from German registry donors.^{16,17} Patients were assigned to 2 groups: if the patient HLA phenotype was explainable by any pair of haplotypes available in the registry haplotype database, the phenotype was considered common; other HLA phenotypes, consisting of at least one rare putative haplotype, where the patient's HLA type could not be formed from haplotypes listed in the database, were considered as rare.

Definitions

Overall survival (OS) was defined as time to death of any cause and was censored at the time of last follow-up. Disease-free survival (DFS) was defined as time to relapse of primary disease or death from any cause and was also censored at the time of last follow-up. Relapse incidence (RI) was defined as the cumulative probability of relapse at any given time point; TRM was defined as mortality in complete remission of disease. Primary graft failure (PGF) was defined as failure to engraft 28 days after transplantation, with engraftment defined as absolute neutrophil count $>0.5 \times 10^9$ /L for the first of ≥ 3 consecutive days.⁸

Statistical analysis

For univariate analysis of OS, the Kaplan-Meier method and log-rank testing were used. Multivariate analysis of OS and DFS was performed using extended Cox models.¹⁸ For TRM and RI, competing risks analysis was used.¹⁹ PGF was analyzed using logistic regression. Statistical models covered covariates defining the EBMT risk score: patient age, disease stage, time to transplantation, and donor-recipient gender combination.' In addition to these, HLA matching status, patient and donor cytomegalovirus (CMV) status, year of transplantation, conditioning regimen intensity, donor origin (national vs international), and treatment with anti-thymocyte globulin (ATG) were evaluated. Missing data for ATG and CMV status were treated as separate categories in multivariate analysis (missing: ATG treatment 19.0%, and CMV status 34.9%).²⁰ To account for possible natural killer cell alloreactivity, the KIR-ligand incompatibility model (predicted from the HLA phenotype in HLA-B and -C mismatched cases), as well as the patient HLA-C phenotype (C1 group, Asn80; C2 group, Lys80), were included in the statistical models.^{5,6} Adjustments for diagnosis and center effect were made.^{7,21} Subanalysis of single HLA mismatches was performed, but as numbers in some subgroups were too small for a robust analysis, combined models, including all HLA mismatches, were calculated as well. Confounding effects for presence of multiple HLA mismatches were considered in these models and adjusted.²² All models were checked for interactions and proportional hazards assumption. Violations of the proportional hazards assumption by the disease stage and conditioning regimen intensity were adjusted using time-dependent covariates. Because of multiple comparisons, the significance level in multivariate analyses was set to .01.

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Locus	n	HR	95% CI	Р
os				
Complete match	1511	1.00		
HLA-A MM	282	1.43	1.19-1.72	<.001
HLA-B MM	283	1.52	1.20-1.93	<.001
HLA-C MM	620	1.35	1.17-1.56	<.001
HLA-DRB1 MM	126	1.42	1.10-1.82	.006
HLA-DQB1 MM	191	1.23	1.00-1.51	.050
DFS				
Complete match	1511	1.00		
HLA-A MM	282	1.25	1.05-1.50	.014
HLA-B MM	283	1.32	1.05-1.66	.016
HLA-C MM	620	1.32	1.15-1.51	<.001
HLA-DRB1 MM	126	1.29	1.01-1.63	.040
HLA-DQB1 MM	191	1.18	0.97-1.43	.110
TRM				
Complete match	1509	1.00		
HLA-A MM	282	1.51	1.16-1.98	.003
HLA-B MM	283	1.50	1.19-1.89	<.001
HLA-C MM	620	1.45	1.19-1.76	<.001
HLA-DRB1 MM	126	1.57	1.12-2.21	.009
HLA-DQB1 MM	191	1.28	0.96-1.71	.091

Table 2. Analysis of the impact of HLA mismatches	in all
transplantations after multivariate modeling	

Analysis of OS, DFS, and TRM for any mismatch at each locus. Significance level set to P < .01. Significant results are bold.

Results

The patient cohort characteristics are summarized in Table 1. A total of 2646 patients were included in this analysis, with a median post-transplant follow-up time of 24 months. Transplantations were performed between 1997 and 2010. Median patient age was 51 years (range: 18-75 years). The conditioning regime was myeloablative in 63.9% of the cases. Most patients received PBSCs as a stem cell source (86.9%).

os

Analysis of OS showed higher risk for death if any HLA mismatch was present (Table 2). Highest hazard ratios (HRs) were seen for HLA-A, -B, and -DRB1 mismatches. Classification of mismatches in allele or antigen mismatches (Table 3) showed that the HLA-B allele and HLA-C antigen mismatches were most strongly associated with adverse outcome. Comparison of allele vs antigen mismatches within each locus showed slightly lower estimates for the HLA-C allele and HLA-DQB1 allele mismatches. These differences were not statistically significant (HLA-C allele vs antigen mismatch: HR 0.96, confidence interval [CI] 0.70-1.30, P = .780; HLA-DQB1 allele vs antigen mismatch: HR 0.79, CI 0.54-1.17, P = .240). HLA-DQB1 mismatching did not reach statistical significance, although a trend toward higher risk could be seen (HR 1.23, CI 1.00-1.51, P = .050; Table 2). Separate analysis of allele and antigen mismatches showed that an impact of HLA-DQB1 mismatching might be limited to antigen differences (HR 1.39, CI 1.04-1.85, P = .028; Table 3).

Subanalysis of single mismatches (Table 4) revealed similar HRs for HLA class I mismatches compared with the model including multiple mismatches (Table 3).

Analysis of mismatch combinations showed HRs increasing with the numbers of HLA mismatches, whereas classification in "allele or antigen mismatch" for single and double mismatches did not add any predictive value (Table 5) regarding mortality risk. Univariate analysis also did not reveal any significant difference between single allelic and antigenic HLA-A, -B, -C, or -DRB1 mismatches as shown in Figure 1 (loci combined, P = .924), whereas the difference between 8/8 and 7/8 matched transplantations was highly significant (P < .001). HLA-DQB1 mismatches were not included in this analysis because multivariate analysis indicated a trend toward worse outcome for HLA-DQB1 antigen differences compared with allelic incompatibilities for this locus.

If double mismatched cases (including HLA-DQB1 mismatches) were grouped according to the HLA class combination of their mismatches (Table 6), a higher HR was seen for combined class I and class II mismatches vs double HLA class I or double HLA class II mismatches, respectively. The direct comparison between combined HLA class I+II mismatches vs double HLA class I or double HLA class I or double HLA class I mismatches was, however, not statistically significant (HR 1.29, CI 0.88-1.90, P = .190).

DFS

In general, HRs for events in DFS analysis were lower than in OS. However, the observed HR-patterns were similar, with the highest impact seen for HLA-B and the lowest for HLA-DQB1 mismatches. The trend for adverse outcome of HLA-DQB1 mismatching was again mostly due to HLA-DQB1 antigen mismatches, whereas DQB1 allele mismatching had no significant impact on DFS.

Most predictive of an adverse outcome were HLA-C antigen mismatches (HR 1.32, CI 1.14-1.52, P < .001), whereas for HLA-C allele mismatches, such an association was not found (HR 1.20, CI 0.93-1.55, P = .150). The number of mismatches was a strong predictor for DFS, whereas antigen or allele differences did not show differential patterns. Classification according to combinations of class I and class II mismatches showed the highest risk for combined class I and class II mismatches and lower HRs for double class I or double class II mismatches.

TRM

The observed HRs for TRM were generally higher than those found for DFS and OS. HLA-A, -B, and -DRB1 mismatches had the highest impact, whereas the lowest impact was seen for HLA-DQB1 mismatches. The high impact of HLA mismatching on TRM was also seen in the analysis of HLA mismatch combinations. An already high risk for TRM in transplantations with single HLA allele or antigen mismatched donors increased further if ≥ 2 mismatches were present.

Classification of double mismatches according to the HLA class showed highest TRM for combined class I/class II mismatches and lower ratios for double class I or double class II mismatches. A direct comparison between these subgroups did, however, not reach statistical significance (HR 1.45, CI 0.88-2.38, P = .140).

RI and PGF

Univariate and multivariate analysis showed no statistically significant effect of HLA mismatching on relapse incidence, neither for all loci combined nor for individual locus mismatches (relapse incidences for all loci combined: 10/10 matched, 33.7%; 9/10 matched, 32.0%, $\leq 8/10$ matched, 27.1%; P = .512). Interestingly, C2C2 KIR-ligand status in patients was associated with a slightly increased relapse rate (HR 1.27, CI 1.03-1.57, P = .028).

PGF rates were higher for \geq 2 HLA mismatches, but this difference was not statistically significant after correction for graft source

Locus	n	HR	95% CI	I P	
os					
Complete match	1511	1.00			
HLA-A allele MM	121	1.37	1.04-1.80	.026	
HLA-A antigen MM	164	1.41	1.12-1.77	.003	
HLA-B allele MM	237	1.48	1.15-1.91	.003	
HLA-B-antigen MM	50	1.48	1.00-2.20	.052	
HLA-C allele MM	106	1.28	0.98-1.67	.072	
HLA-C antigen MM	528	1.34	1.15-1.57	<.001	
HLA-DRB1 allele MM	108	1.46	1.12-1.90	.005	
HLA-DRB1 antigen MM	18	1.12	0.57-2.17	.750	
HLA-DQB1 allele MM	107	1.10	0.84-1.45	.480	
HLA-DQB1 antigen MM	85	1.39	1.04-1.85	.028	
DFS					
Complete match	1511	1.00			
HLA-A allele MM	121	1.22	0.94-1.59	.140	
HLA-A antigen MM	164	1.23	0.98-1.53	.070	
HLA-B allele MM	237	1.33	1.04-1.69	.024	
HLA-B-antigen MM	50	1.20	0.81-1.77	.370	
HLA-C allele MM	106	1.20	0.93-1.55	.150	
HLA-C antigen MM	528	1.32	1.14-1.52	<.001	
HLA-DRB1 allele MM	108	1.38	1.07-1.77	.014	
HLA-DRB1 antigen MM	18	0.87	0.45-1.69	.670	
HLA-DQB1 allele MM	107	1.06	0.81-1.37	.690	
HLA-DQB1 antigen MM	85	1.34	1.02-1.77	.036	
TRM					
Complete match	1509	1.00			
HLA-A allele MM	121	1.22	0.80-1.86	.360	
HLA-A antigen MM	164	1.64	1.21-2.22	.002	
HLA-B allele MM	237	1.62	1.26-2.08	<.001	
HLA-B-antigen MM	50	1.07	0.61-1.89	.810	
HLA-C allele MM	106	1.55	1.09-2.21	.015	
HLA-C antigen MM	528	1.38	1.12-1.69	.002	
HLA-DRB1 allele MM	108	1.68	1.17-2.40	.005	
HLA-DRB1 antigen MM	18	0.71	0.22-2.30	.570	
HLA-DQB1 allele MM	107	1.27	0.88-1.82	.200	
HLA-DQB1 antigen MM	85	1.25	0.82-1.89	.300	

Table 3. Analysis of the impact of HLA mismatches in all	
transplantations after multivariate modeling	

Analysis of OS, DFS, and TRM with allele and antigen mismatches separated. Significance level set to P < .01. Significant results are bold.

n, number of individuals within the respective category; MM, mismatch.

and treatment with growth factors (PGF rate 10/10 matched: 4.2%, 9/10 matched: 3.4%, $\leq 8/10$ matched: 8.2%, P = .080). No differential impact on PGF was seen if the graft was obtained from a national or an international donor.

Clinical predictors

HRs for 10/10, 9/10, and 8/10 HLA mismatch groups are shown in Table 7 in context with other clinical predictors. 8/10 matched pairs showed a significantly higher risk than the 9/10 group (8/10 vs 9/10—OS: 1.47, CI 1.19-1.82, P < .001; DFS: 1.47, CI 1.20-1.80 P < .001, TRM: 1.54, CI 1.18-2.02, P = .001). Besides HLA match, adverse outcomes were associated with advancing disease stage and patient age. Transplantations before 2004 had worse outcomes. Transplantations performed with international donors had a worse outcome compared with transplantations with national donors. Male patients with female international donors showed an even higher HR in analysis of OS (HR 1.51, CI 1.18-1.92, P < .001). This effect was independent from HLA mismatching, as well as the year of transplantation, and can also be shown in a subset of transplantations performed with 10/10 matched donors (P = .047; Figure 2).

Predicted KIR-ligand incompatibility in HLA-C mismatched cases did not influence outcomes, but a patient HLA-C2C2 phenotype was associated with worse outcomes, both in matched and mismatched cases (Table 7).

HLA phenotype classification indicated a significantly higher proportion of patients with rare HLA phenotypes among recipients of grafts from an international donor compared with patients with a national donor (n = 71 of 555, 12.8% vs n = 85 of 2091, 4.1%; χ^2 ; P < .001). No significant impact was seen for ATG treatment, CMV status, and graft source on survival outcomes.

Discussion

We confirmed that a distinction between allele and antigen mismatches (as indicators for the amount of structural disparities between HLA molecules) does not correlate with differential outcome. It is rather the presence or absence of any HLA mismatch at the allele level that must be considered as clinically important for donor selection.^{23,24} This supports the current practice of allele-level HLA typing and matching for HSCT. Among allelic differences, permissive mismatch combinations might exist. This applies especially to HLA-C, where HRs for single-allele mismatches are constantly lower across different studies compared with single-antigen mismatched cases. Further work and significantly higher numbers of transplants are necessary to elucidate such combinations.²⁵

An adverse effect of a patient C2C2 KIR-ligand phenotype was found, which is in accordance with previous findings.^{6,26,27}

In our study, patients transplanted with donors from the national registry showed better survival than patients with international donors. Based on the analysis of PGF, factors determining transplant product quality do not appear to be a reason for this observation. Usually, an international donor is only selected if no well-matched national donor is available. Our findings indicate that the fraction of patients with rare HLA phenotypes is significantly higher in patients who received grafts from international donors. Adverse outcome has been reported for patients with rare HLA phenotypes by Jöris et al.²⁸ Differences in the extended haplotype (ie, immunologically relevant loci on chromosome 6 beyond the routinely typed classical HLA loci) may influence outcome even if patients and donors are 10/10 matched as suggested by Petersdorf and Hansen.²⁹ Such differences may be more frequent in patients with rare HLA phenotypes and when donors from a different ethnic background are selected.³⁰ In addition, non-HLA polymorphisms may influence transplant outcome,^{31,32} and further diversity may be added if donors from a different immunogenetic background are selected. The national vs international donor effect may be a phenomenon that specifically applies to the German setting, because for most of the patients, a national donor can be found, possibly segregating the remaining patients as an immunogenetic risk population. The situation is quite different in countries with smaller donor registries or a more HLA-diverse population, which predominantly use international donors. Additional risk may be conferred by prolonged searches, possibly leading to progression of the disease. This finding, although significant, does not alter search algorithms as the effect of a single HLA mismatch is stronger than the hazard added by selecting an international donor (Table 7). Thus, a matched international donor would still be preferred over a single mismatched national donor.

In our analysis, we found a trend toward higher risk for HLA-DQB1 mismatches, which could be mainly attributed to HLA-DQB1 antigen differences. This did not reach statistical significance, 3224 FÜRST et al

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Table 4. Subanalysis of	single mismatc	hed transplan	tations after
multivariate modeling			

Locus	n	HR	95% CI	Р
os				
Matched	1511	1.00		
A-allele-MM	61	1.44	1.02-2.04	.039
A-antigen-MM	121	1.31	1.00-1.72	.047
B-allele-MM	76	1.53	1.11-2.09	.009
B-antigen-MM	30	1.79	1.13-2.83	.014
C-allele-MM	67	1.10	0.78-1.56	.600
C-antigen-MM	300	1.27	1.06-1.53	.009
DR-allele-MM	46	1.17	0.75-1.82	.500
DR-antigen-MM	7	NA	NA	NA
DQ-allele-MM	54	0.76	0.48-1.21	.250
DQ-antigen-MM	41	1.42	0.93-2.17	.100
DFS				
Matched	1511	1.00		
A-allele-MM	61	1.25	0.90-1.76	.190
A-antigen-MM	121	1.16	0.89-1.50	.270
B-allele-MM	76	1.33	0.98-1.81	.067
B-antigen-MM	30	1.39	0.88-2.21	.160
C-allele-MM	67	1.12	0.81-1.55	.480
C-antigen-MM	300	1.25	1.05-1.48	.012
DR-allele-MM	46	1.14	0.75-1.72	.550
DR-antigen-MM	7	NA	NA	NA
DQ-allele-MM	54	0.79	0.52-1.20	.260
DQ-antigen-MM	41	1.32	0.88-1.97	.180
TRM				
Matched	1509	1.00		
A-allele-MM	61	1.33	0.80-2.23	.280
A-antigen-MM	121	1.53	1.07-2.20	.021
B-allele-MM	76	1.66	1.10-2.52	.017
B-antigen-MM	30	1.60	0.85-2.99	.140
C-allele-MM	67	1.34	0.85-2.12	.210
C-antigen-MM	300	1.29	0.99-1.69	.064
DR-allele-MM	46	1.53	0.86-2.73	.150
DR-antigen-MM	7	NA	NA	NA
DQ-allele-MM	54	1.03	0.59-1.81	.910
DQ-antigen-MM	41	1.16	0.57-2.40	.680

Impact of single allele or antigen HLA MMs per locus on OS, DFS, and TRM. Significance level set to P < .01. Cases with more than one HLA mismatch were excluded from this analysis. Significant results are bold

NA, number too low for analysis

as numbers of HLA-DOB1 mismatches were relatively small due to linkage disequilibrium with well-matched HLA-DRB1. Furthermore, DRB*3,4,5 polymorphisms in non-haplotype-matched unrelated HSCT might confer additional risk in the presence of an HLA-DQB1 mismatch.³³ Future studies in an extended cohort and typing for HLA-DRB*3,4,5 may allow a better understanding of the relative impact of HLA-DQB1 allele or antigen mismatches.

Due to weak linkage disequilibrium between DPB1 and the HLA loci we analyzed, the fraction of donors matched for DPB1 or with a permissive or nonpermissive mismatch should be very similar in all subgroups of our study population.³⁴ Therefore, we may assume that relative risk estimates obtained in our analysis are still accurate, although we could not include the HLA-DPB1 matching status in our multivariate analysis.

Analysis of mismatch combinations revealed higher HRs with increasing numbers of disparities and suggests additive effects of these incompatibilities.³⁵ In general, the impact of HLA mismatching on TRM is stronger and the effect on DFS is smaller compared with HRs observed in OS. Similarly to other studies, there was no significant effect of HLA incompatibilities on relapse incidence.^{12,13} The competing event relapse (or death from relapse) therefore slightly

blurs the analysis of OS and in particular of DFS, leading mostly to lower HRs in these analyses within the corresponding mismatch groups. PGF was not significantly associated with HLA mismatching in our study cohort. Findings in nonmalignant diseases differ regarding this issue.³⁶ The graft source (bone marrow vs PBSCs) did not influence survival end points, which is consistent with a previous observation in patients with acute leukemia.37 The cohort composition with respect to graft source (limited to bone marrow and PBSCs) in large studies probably does not impair comparability of results.^{12,13} The vector of mismatches was not considered, because no significant differences in outcome for the end points analyzed in our study were reported if HLA mismatches were grouped accordingly.¹⁴

We saw higher HRs for patients with 2 incompatibilities when these involved a combined HLA class I+II mismatch, contrary to a double class I or II mismatch situation. Although the direct comparison of these 2 groups did not reach statistical significance, this finding is of interest, as the majority of HLA class II disparities in our study cohort affected HLA-DQB1. The pattern of HRs for double HLA mismatch combinations in our study suggests that despite its controversial impact in single mismatch cases, HLA-DQB1 appears to have a significant effect in double mismatched situations and should be considered accordingly in donor selection. The higher risk for combined HLA class I and II mismatches is descriptive at present, as direct comparison with double HLA class I or double HLA class II mismatches did not reach statistical significance. This may be due to insufficient numbers in our study and should be re-evaluated in a larger study. Nevertheless, pending such a study, it might be advisable to avoid combined HLA class I and II incompatibilities if alternatives are at hand.

In our analysis, HLA-A, and particularly HLA-B, mismatches, were strongly associated with adverse outcome. Comparisons between different other large studies, which were conducted by the

Table 5. Subanalysis of mismatch combinations after multivariate modeling

Mismatch combination	n	HR	95% CI	Р
os				
Complete match	1511	1.00		
Single allele MM	304	1.21	1.01-1.44	.044
Single antigen MM	499	1.30	1.12-1.51	<.001
Double allele MM	46	1.70	1.15-2.53	.008
Double MM allele + antigen	139	2.04	1.54-2.69	<.001
Double antigen MM	62	1.76	1.25-2.48	.001
Triple MM	61	2.56	1.71-3.85	<.001
Multiple MM	24	3.11	1.70-5.67	<.001
DFS				
Complete match	1511	1.00		
Single allele MM	304	1.11	0.93-1.31	.240
Single antigen MM	499	1.21	1.05-1.39	.009
Double allele MM	46	1.60	1.09-2.33	.015
Double MM allele + antigen	139	1.75	1.34-2.27	<.001
Double antigen MM	62	1.75	1.27-2.41	<.001
Triple MM	61	2.03	1.37-3.03	<.001
Multiple MM	24	2.17	1.22-3.88	.009
TRM				
Complete match	1509	1.00		
Single allele MM	304	1.46	1.14-1.87	.003
Single antigen MM	499	1.34	1.08-1.66	.007
Double allele MM	46	2.08	1.21-3.57	.008
Double MM allele + antigen	139	2.16	1.57-2.99	<.001
Double antigen MM	62	1.90	1.18-3.07	.009
Triple MM	61	2.94	1.91-4.51	<.001
Multiple MM	24	2.93	1.66-5.16	<.001

Analysis of OS, DFS, and TRM in different mismatch groups. Multiple mismatches, >3 HLA mismatches. Significance level set to P < .01. Significant results are bold.

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Figure 1. OS of patients with 8/8 matched donors (solid line), patients with single allele mismatched donors at loci HLA-A, -B, -C, or -DRB1 (dashed line), and single antigen mismatched donors at loci HLA-A, -B, -C, or -DRB1 (dotted line). HLA-DQB1 status was omitted from this analysis because multivariate modeling indicated a possible differential impact of HLA-DQB1 allele and antigen mismatches. Comparison single allele vs single antigen mismatch, P = .924; comparison 8/8 matched vs 7/8 matched log-rank, P < .001.



CIBMTR, did not always show similar results across these cohorts.¹¹⁻¹³ Reasons for such differences may be statistical limitations due to sample size or variations regarding patient characteristics, especially patient age, diagnosis, follow-up-period, changes in treatment over time, and, in particular, the ethnic composition of the different study cohorts.

The relatively high number of HLA-C antigen mismatches in our study can be explained mainly by the fact that HLA typing and matching requirements before May 2005 in Germany considered HLA-A and -B at low resolution only. Retrospective typing disclosed therefore a considerable number of allele and antigen mismatches for HLA-C. In addition, confirmatory typing in general is more likely to show mismatches at the HLA-C locus, as the majority of unrelated donors are usually not typed for HLA-C on registration.

Table 6. Subanalysis of double HLA mismatch combinations after multivariate modeling

Mismatch combination	n	HR	95% CI	Р
os				
Complete match	1511	1.00		
Double MM HLA class I+II	67	2.20	1.60-3.03	<.001
Double MM only HLA class I	162	1.68	1.27-2.22	<.001
Double MM only HLA class II	18	1.83	1.04-3.22	.036
DFS				
Complete match	1511	1.00		
Double MM HLA class I+II	67	2.03	1.50-2.74	<.001
Double MM only HLA class I	162	1.55	1.19-2.02	.001
Double MM only HLA class II	18	1.57	0.89-2.75	.120
TRM				
Complete match	1509	1.00		
Double MM HLA class I+II	67	2.69	1.75-4.14	<.001
Double MM only HLA class I	162	1.90	1.39-2.61	<.001
Double MM only HLA class II	18	1.52	0.63-3.67	.360

Analysis of OS, DFS, and TRM in different mismatch groups. Significance level set to P < .01. Significant results are bold.

HLA-B antigen mismatches are rare because if a mismatch search had to be initiated, often an HLA-A mismatch was preferred over one for HLA-B. This is due to the strong HLA-B/C linkage disequilibrium, which would often reveal an additional HLA-C mismatch if an HLA-B mismatch was selected for further testing.³⁸

In all transplants included in this analysis, HLA class II molecules were high-resolution tested at the time of confirmatory typing, explaining the low number of HLA-DRB1 mismatches observed, in particular for HLA-DRB1 antigens, as such mismatches were totally avoided if possible. Similar HLA mismatch patterns were observed in other large studies.¹¹⁻¹³ In contrast to these studies, the vast majority of patients in our cohort were transplanted in recent years (2006-2010; 66.4%). As a consequence, most of the patients included in our study were transplanted with PBSCs as a graft source (86.9%). Retrospective studies published thus far by the CIBMTR investigated the impact of HLA matching on HSCT outcome, mostly including patients transplanted before 2007 with bone marrow^{11,12} or PBSCs only¹³ and with patient diagnoses being restricted to AML, ALL, CML, and MDS. Our study is currently the first to analyze the impact of high-resolution HLA matching on outcome of HSCT in a large European cohort including >2500 transplantations.

In previous studies, the problem of possible confounding of multiple HLA mismatches was addressed by performing subanalyses of single HLA mismatches. We performed such an analysis in our cohort and present the results in Table 4. In our view, the advantages of elimination of confounding in such a subanalysis are offset by the limitations conferred by multiple comparisons and low numbers within certain subgroups. Comparison of the single mismatch subanalysis (Table 4) with the model including all incompatible cases (Table 3) demonstrates good consistency of estimates in the groups with substantial numbers (n > 50), showing that it is feasible to create survival models with inclusion of cases with >1 mismatch, possibly leading to more robust estimates. 3226 FÜRST et al

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Table 7. Effects of clinical predictors on survival endpoints

			OS		DFS			TRM				
Predictor	n	HR	95% CI	Р	n	HR	95% CI	Р	n	HR	95% CI	Р
HLA match												
10/10	1511	1.00			1511	1.00			1509	1.00		
9/10	803	1.26	1.11-1.43	<.001	803	1.17	1.03-1.32	.012	803	1.38	1.15-1.66	<.001
8/10	247	1.86	1.51-2.29	<.001	247	1.71	1.40-2.09	<.001	247	2.13	1.64-2.77	<.001
Disease stage												
Early	1047	1.00			1047	1.00			1047	1.00		
Intermediate	922	1.46	1.26-1.70	<.001	922	1.50	1.31-1.73	<.001	921	1.60	1.29-1.98	<.001
Advanced	677	2.18	1.87-2.54	<.001	677	2.17	1.88-2.51	<.001	676	1.97	1.59-2.43	<.001
Patient age (years)												
<40	734	1.00			734	1.00			733	1.00		
40-55	951	1.39	1.19-1.62	<.001	951	1.30	1.13-1.50	<.001	950	1.60	1.27-2.00	<.001
>55	961	1.76	1.48-2.10	<.001	961	1.56	1.33-1.82	<.001	961	1.85	1.43-2.40	<.001
Donor-recipient gender												
Male-male	1227	1.00			1227	1.00			1227	1.00		
Male-female	362	1.13	0.95-1.34	.170	362	1.15	0.98-1.35	.087	362	0.96	0.75-1.22	.720
Female-female	393	0.91	0.77-1.08	.270	393	0.92	0.78-1.08	.300	392	0.67	0.51-0.88	.004
Female-male	664	0.99	0.86-1.14	.870	664	0.98	0.86-1.13	.810	663	0.91	0.74-1.12	.390
Year of transplantation												
1997-2003	455	1.00			455	1.00			455	1.00		
2004-2010	2191	0.76	0.64-0.91	.002	2191	0.79	0.67-0.93	.005	2189	0.61	0.49-0.76	<.001
Donor-recipient CMV status*												
Negative-negative	520	1.00			520	1.00			520	1.00		
Negative-positive	183	1.40	1.08-1.81	.099	183	1.34	1.05-1.71	.017	183	1.23	0.84-1.79	.290
Positive-negative	479	1.20	0.98-1.47	.081	479	1.16	0.96-1.40	.140	478	1.27	0.95-1.70	.100
Positive-positive	539	1.13	0.93-1.38	.210	539	1.11	0.93-1.34	.260	539	1.13	0.85-1.51	.400
Patient KIR-ligand status												
C1C1 or C1C2	2244	1.00			2244	1.00			2242	1.00		
C2C2	402	1.26	1.09-1.47	.027	402	1.30	1.13-1.50	.002	402	1.14	0.91-1.42	.240
Donor origin												
National	2091	1.00			2091	1.00			2090	1.00		
International	555	1.23	1.07-1.41	.004	555	1.23	1.08-1.41	.002	554	1.29	1.06-1.56	.010

Correlation of clinical predictors with OS, DFS, and TRM. Significance level set to P < .01. Significant results are bold.

*Performed as subanalysis with separate model.

Our study included patients with non-Hodgkin lymphoma and multiple myeloma. These disorders have substantially contributed to the transplant activities in Germany in recent years. Most of the transplantations in our study have been performed between 2006 and 2010 and draw a more recent clinical picture than other published large studies, which thus far include patients transplanted before 2007.¹¹⁻¹³ On the other hand, inclusion of more recent transplants in our study resulted in shorter median follow-up compared with other studies. Nevertheless, because most adverse events occur within the first 2 years following HSCT, the observations of our study can be considered accurate.

Improvements in therapy have made conditioning regimens safer, especially for older patients. As a consequence, the proportion of older patients undergoing transplantation is rising, and this might influence outcome results. In addition, the immunogenetic background of a study population could influence results, as HLA differences in genetically diverse populations may confer different risks than in genetically more homogenous cohorts.²⁵

Limitations of our study are that analysis of graft versus host disease was not possible and data about ATG treatment and CMV status were partly missing.

Despite the mentioned methodological differences, we could confirm patterns that were described in previous reports: HLA-C antigen mismatches are strongly predictive for adverse outcome, whereas the impact of HLA-C allele mismatch remains uncertain.^{11-13,39} Nevertheless, when directly compared with HLA-A and HLA-B mismatches, HRs are lower for HLA-C mismatches, especially regarding OS. This observation is consistent across different studies and can be considered robust in view of the high numbers of HLA-C antigen mismatches analyzed. It might indicate manifestation of beneficial KIR receptor-ligand incompatibilities in a subset of patients with HLA-C mismatches. We saw a strong effect of DRB1 allele mismatching (Table 3), especially regarding its impact on TRM. The low number of HLA-DRB1 mismatches in our analysis, however, limits the interpretation of these results.

Our data support HLA-A/B/C/DRB1 typing and matching as minimum requirement for donor search. In general, allele and antigen mismatches should be treated as equivalent, however, keeping in mind uncertainty of the impact of HLA-C-allele mismatches. A 10/10 matched situation is desirable; 9/10 or even 8/10 matched donors may be acceptable, in particular if a single HLA-DQB1 allele mismatch is present, when no better matched unrelated donors are available.⁴⁰ In cases with more mismatches, a careful risk/benefit evaluation is warranted and alternatives, such as cord blood or haploidentical transplantation, should be considered, especially if rapid transplantation is required.^{37,41,42} If >1 mismatch is to be taken into account, it is advisable to avoid a combined HLA class I/II incompatibility.

Furthermore, in cases where initially no perfectly matched donor can be found, the clinical risk for disease progression must be weighed against the probability of finding a better-matched donor Figure 2. OS of patients with 10/10 matched donors, grouped by transplantations performed with national donors (solid line) or international donors (dashed line), logrank P = .047. Of the 213 patients with international donors, 211 were of white and 2 were of Asian ethnicity.



in a prolonged search. Depending on the clinical situation of the patient, in many cases, a mismatched transplantation performed in early disease stage may be a better choice for the patient than a transplant in advanced disease, even if this would eventually be performed with a better-matched donor.

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