

TO THE EDITOR:

Validation of genetic associations with acute GVHD and nonrelapse mortality in DISCOVeRY-BMT

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In their recent report in *Blood Advances*, Martínez-Laperche et al¹ proposed a predictive model for the risk of acute and chronic graft-versus-host disease (GVHD) based on the selection of clinical variables and 25 single-nucleotide polymorphisms (SNPs), spanning different cytokines previously found relevant in the biology of GVHD. The study included 509 patients and their sibling donors from the Spanish Group for Hematopoietic Stem Cell Transplantation (GETH). We attempted to validate their SNP associations with acute GVHD (aGVHD) and nonrelapse mortality (NRM) in DISCOVeRY-BMT, a 2-cohort study of almost 3000 8/8 HLA-matched unrelated donor-recipient paired samples from individuals of European American ancestry.²⁻⁵

The analyses of the GETH data were performed using univariate logistic and least absolute shrinkage and selection operator (LASSO) models.⁶ Univariate logistic regression models for grade 2 to 4 aGVHD, grade 3 to 4 aGVHD, and NRM were constructed via regression of clinical and genetic variables. LASSO models were constructed for all outcomes using genetic and clinical variables from a training set of 85% of patients, with the remaining 15% comprising the test set. Risk scores were calculated using the LASSO outcomes, and recipients were stratified into high- and low-risk groups based on the proportion of events in the total data. To validate the clinical and genetic associations seen with grade 2 to 4 aGVHD, grade 3 to 4 aGVHD, and NRM, we constructed univariate clinical logistic regression models identical to those of Martínez-Laperche et al¹ and LASSO models for grade 2 to 4 aGVHD, grade 3 to 4 aGVHD, and NRM. Twenty-four of the 25 SNPs tested were available for univariate and LASSO analyses in DISCOVeRY-BMT; rs9267487 was used as a surrogate for rs361525 in *TNF*, because the SNPs are in linkage disequilibrium at $r^2 = 0.82$.⁷

Clinical characteristics of both GETH and DISCOVeRY-BMT data are listed in Table 1. In GETH data, univariate clinical models showed total-body irradiation and female donor/male recipient increased the odds of aGVHD and NRM, respectively ($P < .05$). In DISCOVeRY-BMT, total-body irradiation increased the odds of grade 2 to 4 aGVHD (odds ratio [OR], 1.3; 95% confidence interval [CI], 1.12-1.51; $P = 5.9 \times 10^{-4}$) and grade 3 to 4 aGVHD (OR, 1.22; 95% CI, 1.01-1.47; $P = .04$), whereas female donor/male recipient was not associated with NRM at $P < .05$.

We selected 6 SNPs at $P < .05$, with 95% CIs that did not include 1, and tested them for association with aGVHD or NRM in DISCOVeRY-BMT. We assumed the same transmission models (dominant, additive, recessive, or codominant) for donors and recipients presented in the report by Martínez-Laperche et al¹ and adjusted for significant clinical variables but not multiple comparisons. Donor GG/AG vs AA at rs3819024 was associated with NRM in DISCOVeRY-BMT cohort 1 at $P < .05$ (OR, 0.79; 95% CI, 0.64-0.99; $P = .04$) but not cohort 2 (OR, 0.89; 95% CI, 0.61-1.29; $P = .53$), although the effect direction was also risk reducing, as in the GETH data (Figure 1). The dominant models for rs16944, rs1143627, and rs2275913 trended toward a risk reduction in both cohorts in DISCOVeRY-BMT, with an OR of < 1 , whereas these genetic models showed an OR of > 1 in the GETH cohort (Figure 1).

We assessed effect sizes detectable in DISCOVeRY-BMT and GETH for the 25 SNPs tested ranging in MAF from 0.4% to 45%. After correcting for testing in 22 (of 25) independent SNPs

Table 1. Characteristics of European Americans in DISCOVeRY-BMT GETH cohorts

Category	Cohort 1* (n = 2110)	Cohort 2 (n = 777)	GETH (n = 359)
Median age (range), y†	45.3 (0.6-74.5)	50 (0-74)	45 (0-68)
Recipient sex			
Male	1191 (56.4)	429 (55.2)	225 (63)
Female	919 (43.6)	348 (44.8)	134 (37)
Donor sex‡			
Male	1396 (68)	554 (72.6)	201 (56)
Female	656 (32)	209 (27.4)	158 (44)
Female donor/male recipient†			
Yes	332 (15.7)	101 (13.0)	91 (25)
No	1778 (84.3)	676 (87.0)	268 (75)
Disease‡			
AML	1282 (60.8)	488 (62.8)	116 (32)
ALL	483 (22.9)	94 (12.1)	49 (13.5)
MDS§	345 (16.4)	195 (25.1)	34 (9.5)
Other	0 (0)	0 (0)	160 (44)
Stem cell source‡			
Peripheral blood	1365 (64.7)	567 (73)	250 (69.6)
Bone marrow	745 (35.3)	210 (27)	109 (30.4)
Conditioning intensity†			
Myeloablative	1540 (73)	551 (71)	253 (70)
Reduced intensity	570 (27)	225 (29)	106 (30)
Conditioning regimen			
TBI	973 (46.1)	280 (36)	94 (26)
No TBI	1137 (53.9)	497 (64)	265 (74)
Outcome			
Grade 2-4 aGVHD	973 (46.1)	358 (46.1)	115 (32)
Grade 3-4 aGVHD	389 (18.4)	168 (21.6)	50 (14)
NRM	405 (19.2)	141 (18.1)	86 (24)

Data are n (%) unless otherwise noted.

ALL, acute lymphoblastic leukemia; AML, acute myeloid leukemia; MDS, myelodysplastic syndrome; TBI, total-body irradiation.

*To be comparable to multivariate analysis in literature, for analysis of grade 2-4 aGVHD and grade 3-4 aGVHD, patients who died without grade 2-4 aGVHD or grade 3-4 aGVHD, respectively, within the 100 d were excluded. After exclusion, difference in number of patients between grade 2-4 aGVHD cohort and grade 3-4 aGVHD cohort was <0.6% in cohorts 1 and 2, so we did not include separate patient characteristics for grade 3-4 aGVHD cohort in table.

†Tested in clinical models by Martinez-Laperche et al.¹

‡There were 2052 donors in cohort 1 and 763 donors in cohort 2 for analysis of NRM, 1906 donors in cohort 1 and 698 donors in cohort 2 for analysis of grade 2-4 aGVHD II-IV, and 1894 donors in cohort 1 and 692 in cohort 2 for grade 3-4 aGVHD.

§Includes MDS patients in DISCOVeRY-BMT and both MDS and myelofibrosis patients in GETH.

||Other includes non-Hodgkin lymphoma, Hodgkin disease, multiple myeloma, chronic myeloid leukemia, and aplastic anemia.

($\alpha = 0.05/22 = 0.0023$), DISCOVeRY-BMT was powered ($\beta = 0.80$) to detect hazard ratios from 4.0 to 1.2 (NRM and aGVHD 3-4) and 2.4 to 1.1 (aGVHD 2-4) for MAFs from 0.4% to 45%, respectively.³ We were therefore powered to validate the effect sizes detected in the Martinez-Laperche et al¹ report.³ In contrast, the GETH cohort was powered ($\beta = 0.80$) to detect medium effect sizes (OR, 1.6) for a MAF of 45% in grade 2 to 4 aGVHD; however, for variants at the bottom end of the MAF range, the detectable OR was ~ 40 , with further reduced power for NRM and grade 3 to 4 aGVHD outcomes. Compounding this sample size challenge was the fact that GETH SNP data were analyzed for all outcomes using logistic regression. For aGVHD, these events occur early on, and this model may be reasonable. However, failure to analyze

overall survival or NRM with survival models can result in information loss, and the use of survival models is preferable.^{5,8-11}

The well-described LASSO and prediction modeling performed in the GETH cohorts provides us the opportunity to consider important issues in association testing and prediction modeling of transplantation outcomes using genetic variables.^{6,12} Because of problems with interpretations of the proposed LASSO models in the report, we did not attempt validation of the exact model in DISCOVeRY-BMT. Specifically, the same SNPs were included under multiple transmission assumptions for all LASSO models. For example, in the LASSO model for grade 3 to 4 aGVHD, 7 of 11 SNPs were included in 2 different transmission models, and 1 SNP was included in

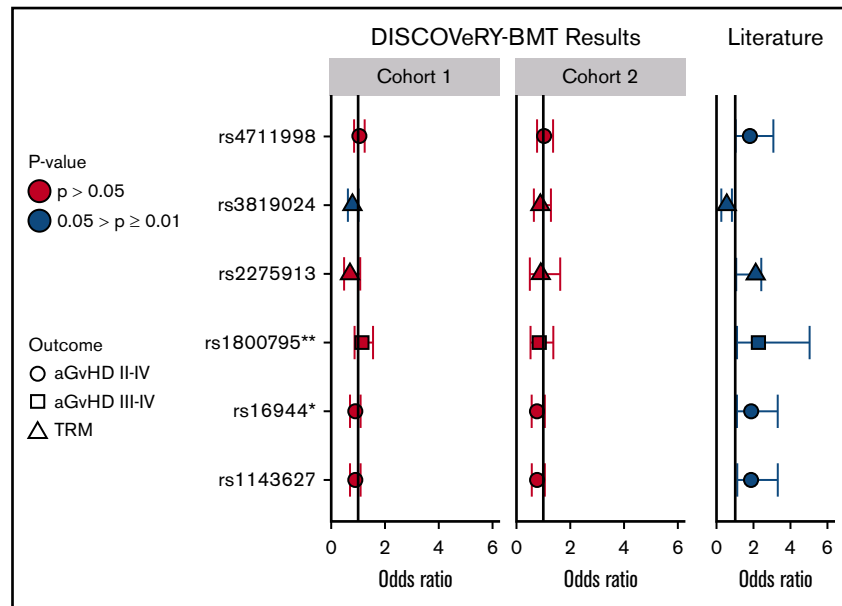


Figure 1. Validation in DISCOVeRY-BMT of significant SNP associations in Martínez-Laperche et al report. This figure shows the ORs, 95% CIs, and *P* values as reported by Martínez-Laperche et al¹ (literature) and validation of these associations in DISCOVeRY-BMT cohort 1 and cohort 2. Martínez-Laperche et al reported 6 SNPs with *P* < .05 and 95% CIs that did not include OR = 1, which are shown along the y-axis. The results are grouped by those shown in the Martínez-Laperche et al report (literature) and DISCOVeRY-BMT (cohort 1 and cohort 2), as indicated in the gray boxes displayed along the top of the figure. Each circle, square, and triangle represents an OR from regression analysis, with the shapes corresponding to grade 2 to 4 aGVHD, grade 3 to 4 aGVHD, or NRM, respectively. The colors represent *P* values of the analyses: .05 > *P* ≥ .01 (blue) and *P* > .05 (red). rs2275913 and rs3819024 are donor variants; the remaining 4 SNPs are recipient associations. *SNPs with a 5% and **SNPs with a 10% difference in minor allele frequency (MAF) between DISCOVeRY-BMT and GETH, respectively; all other MAFs were comparable between the groups. For example, (recipient) SNP rs4711998 was associated with increased risk of grade 2 to 4 aGVHD in GETH (blue and to the right of OR = 1) but resides on OR = 1 and *P* > .05 in DISCOVeRY BMT.

3 transmission models (ie, rs8193036 and rs2430561 [recipients] and rs2275913 [donors]) were included as recessive [2 copies of the minor allele impact risk] and additive [each additional copy of the minor allele impact risk]. The LASSO models for all outcomes had various combinations of additive, dominant, recessive, and codominant models for the same SNP. In human population genetic studies, it is not biologically reasonable that the same SNPs are acting in multiple contradictory ways to change the risk of transplantation outcomes. Although it is common practice for genetic association and prediction studies to use additive genetic models,^{13,14} other appropriate analytic approaches have been developed to assess modes of transmission.^{15,16}

To consider the 25 SNPs in aggregate, we constructed 3 separate LASSO models for grade 2 to 4 aGVHD, grade 3 to 4 aGVHD, and NRM under an additive genetic model. All 25 recipient and donor SNPs for cohorts 1 and 2 were possible variables (this approach most closely represents Table 4 in the Martínez-Laperche et al¹ report); each cohort was divided into a training (85%) and test (15%) set. To select the variables with the smallest prediction errors, we built logistic LASSO regression using fivefold cross validation to find the best penalty parameter λ , and we repeated this cross validation 50 times to give a robust estimate of λ by taking the median. The final LASSO model selected the SNPs with the best λ . LASSO models for 2 outcomes selected 1 SNP; however, the coefficients were almost 0 ($<5 \times 10^{-15}$). Thus, for unrelated donor-recipient pairs, we concluded the 25 SNPs were not associated with either aGVHD or NRM and therefore did not pursue additional predictions. In addition, the predictive models specified in the report

cannot be generalized to other transplantation patient cohorts or applied in a prospective setting, because high- and low-risk cut points were determined by the proportion of events in the GETH cohort. The successful stratification of risks for aGVHD and NRM outcomes in 1 cohort may not necessarily be carried over to another cohort when prediction is needed.¹⁷

Our inability to validate the univariate associations or find SNPs predictive of either aGVHD or NRM may have been driven by the differences in transplantation type (related sibling vs unrelated donor), distribution of disease, genomic ancestry, and/or event rates between DISCOVeRY-BMT and GETH. It is important to consider that these single SNPs were initially identified as important in small expression studies or were selected because they reside in gene promoter regions. However, recent large-scale functional studies of SNPs in thousands of samples can now be leveraged.¹⁸⁻²⁰ For example, rs3819024, although selected as an *IL17A* donor variant, is correlated with the expression of *PAQR8*, not *IL17A*, and only in whole blood, although tested in >70 tissues.¹⁸⁻²⁰ Therefore, collectively, these variants may not be informative for the genes of interest. Irrespective of this, we must consider that the DISCOVeRY-BMT results show these SNP findings are not generalizable to other transplantation populations, and to start building successful prediction models of transplantation outcomes, we need larger homogeneous studies across multiple patient populations.

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