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Long- and short-term outcomes in renal allografts with deceased donors: A large recipient and donor genome-wide association study

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Improvements in immunosuppression have modified short-term survival of deceased-donor allografts, but not their rate of long-term failure. Mismatches between donor and recipient HLA play an important role in the acute and chronic allogeneic immune response against the graft. Perfect matching at clinically relevant HLA loci does not obviate the need for immunosuppression, suggesting that additional genetic variation plays a critical role in both short- and long-term graft outcomes. By combining patient data and samples from supranational cohorts across the United Kingdom and European Union, we performed the first large-scale genome-wide association study analyzing both donor and recipient DNA in 2094 complete renal transplant-pairs with replication in 5866 complete pairs. We studied deceased-donor grafts allocated on the basis of preferential HLA matching, which provided some control for HLA genetic effects. No strong donor or recipient genetic effects contributing to long- or short-term allograft survival were found outside the HLA region. We discuss the implications for future research and clinical application.

**KEYWORDS**

basic (laboratory) research/science, genomics, graft survival, kidney transplantation/nephrology, rejection, translational research/science

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**INTRODUCTION**

Kidney transplantation is a highly successful treatment for end-stage renal failure, with significant benefits for recipients both in survival and quality of life. Early outcomes have steadily improved over the last 10 years,\(^1\) with risk-adjusted and death-censored, 1-year renal graft survival rates of 94% and 97% for deceased and living donor transplants, respectively.\(^2\) However, both late allograft loss and increased mortality among transplant recipients remain key challenges for the transplant community. There are a wide number of factors that are known to influence long-term transplant outcome, including donor factors such as age and comorbidity, recipient factors such as comorbidity and response to immunosuppression, as well as allograft ischemic time, the degree of HLA mismatch, and the development of donor-specific antibodies.\(^3\)\(^5\) However, a comprehensive understanding of the pathophysiology
of graft failure has remained elusive, with the observed variation in patient outcomes still inadequately explained by our current understanding of risk factors. An improved understanding of the determinants of transplantation outcome would allow the development of truly personalized approaches to the management of transplant recipients.

The importance of genetic factors in transplantation has been clear since the inception of the technique, with the first successful kidney transplant having been performed between identical twins in 1954. Renal transplantation between identical twins continues to show excellent long-term outcomes,\(^6,7\) and HLA matching has a large impact on graft survival even in the modern era of immunosuppression.\(^8\)

HLA genes are highly polymorphic, and demonstrate the importance of genetic variation in donor-recipient pairing that impacts on long-term outcomes. However, over recent decades, our ability to assay human genetic variation beyond the HLA region has increased considerably.

A number of studies have been published over recent years exploring the association between genotypes of interest and renal transplant outcomes.\(^9,10\) A large proportion of these studies have concentrated on immune-related genes, based on the hypothesis that the risk of acute rejection or late allograft loss may be modulated by genetic variation in the immune response. As summarized in Table S1, associations have been described between various transplant phenotypes and single nucleotide polymorphisms (SNPs) in a number of genes including those encoding tumor necrosis factor-\(\alpha\), interleukins-1, -6, and -10, and interferon-\(\gamma\). Of note, many of these studies have reported inconsistent findings. For example, analysis of DNA from donor-recipient pairs in the Collaborative Transplant Study failed to replicate an earlier finding that particular combinations of C3 genotypes in the donor and recipient were associated with reduced graft survival.\(^11,12\) While some of this discrepancy might be explained by methodological or populational differences between these studies, it is difficult to draw firm conclusions about the role of these genetic variations.\(^13\)

More recently, attention has also focused on non-immune-related genetic risk variants. Donor genetic variation in CAV1 (caveolin-1),\(^14\) APOL1 (apolipoprotein-L1),\(^15,16\) or ABCB1 (ATP-binding cassette, sub-family-B, member-1, expressed in the kidney) genes\(^17,18\) has been reported to be associated with increased risk of allograft failure or poorer recipient survival. Recipient genetic variation effects on graft and patient survival have only been reported in 1 cohort for CAV1.\(^14\) In addition to effects of donor genetic variation, it might be expected that recipient genotypes in other pharmacometabolic pathways might also impact on transplant outcomes such as increased risk of acute rejection.\(^19\)

In general, candidate gene studies in renal transplantation have so far failed to provide consistent and reproducible results. Some of the reasons for this may include small sample sizes, variations in genotyping methodology and strategy, and, perhaps most importantly, a lack of consistency in clinical phenotyping.\(^20\) Genome-wide association studies (GWAS) have contributed greatly to an increased understanding of complex common conditions such as inflammatory bowel disease, hypertension, type 2 diabetes, and schizophrenia.\(^21\) A small number of GWAS have been reported in the field of renal transplantation, describing SNPs associated with cardiovascular adverse events in recipients taking calcineurin inhibitor immunosuppression,\(^22\) 2 SNPs associated with serum creatinine levels at 5 years posttransplant,\(^23\) and a number of SNPs associated with the development of new-onset diabetes after transplantation.\(^24\) Recently, a GWAS using pooled DNA of recipient-only origin found variation in 2 new loci associated with acute rejection in both univariate and multivariate analysis.\(^25\) However, these studies were underpowered for discovery of genetic variants with small effect sizes.

The Wellcome Trust Case Control Consortium (www.wtccc.org.uk/ccc3) has led the deployment of GWAS in a wide range of conditions. As part of WTCCC-3, all renal transplant centers in the United Kingdom and Ireland formed the United Kingdom and Ireland Renal Transplant Consortium (UKIRTC; www.ukirtc.org). Collaborative initiatives such as these are essential for the collection of adequate sample numbers, for the sharing of expertise, standardization of techniques, and building consensus on accurate phenotyping of clinical data. Through this consortium, 3936 samples comprising 2094 complete donor-recipient pairs were tested in the GWAS discovery phase, and an additional 5866 complete donor-recipient pairs in the replication phase, making this the largest GWAS conducted to date in renal transplantation outcomes.

### 2 | METHODS

#### 2.1 Discovery study participants

The large multicenter United Kingdom and Ireland Renal Transplant Consortium (www.UKIRTC.org), coordinated by King’s College London in partnership with the WTCCC-3 and the National Health Service Blood and Transplant database (NHS-BT), sourced all available good-quality stored DNA samples and pre-existing GWAS data from both recipients and their corresponding donors from all renal transplantation centers in the United Kingdom and the Republic of Ireland (listed in Table S2). The study was approved by the Hammersmith and Queen Charlotte’s & Chelsea Research Ethics Committee REC No 08/H0707/1, on October 14, 2009. Third-party anonymized clinical data were provided by NHS-BT UK. All samples and anonymized data for the replication cohort were provided by the University of Heidelberg.

Following genotyping and GWAS quality control (see below), there were 2689 unique recipients, 2204 unique donors, and 2094 complete donor-recipient transplantation pairs available for analysis. Figure 1 describes the study design and analysis steps. Table 1 provides additional information for the complete donor-recipient pairs (discrepancies between Figure 1 and Table 1 are because some recipients received more than 1 graft, some donors donated 2 kidneys to different recipients, and not all recipients had matching donor GWAS data and vice versa). The samples and data referred to transplants that took place between December 1981 and December 2007.
Inclusion criteria for the study were as follows: (1) deceased donor kidney transplants only; (2) recipient is an adult (>16 years old); (3) reported European ancestry for recipients; and (4) graft survival time >3 months. Donor-recipient allocation followed NHS-BT standard UK-protocols during the study period.

2.2 | Replication phase participants

A cohort of 5866 complete donor-recipient pairs, with similar ethnicity to that of the discovery study, were obtained from the Collaborative Transplant Study DNA Biobank held at the University of Heidelberg, Germany (Table 1).

Clinical variables, datasets, and analysis are described in Supplementary Methods.

2.3 | Discovery phase genotyping and analysis

A whole genome amplification step was undertaken (Source BioScience, Nottingham) for samples containing 5-40 μL of DNA (n = 990 samples). Samples were assayed via Illumina 670 Quad Custom GWAS chips, and subjected to standard postgenotyping quality control procedures before being imputed to the 1000-genomes reference dataset. Imputation of HLA alleles from SNP genotype data was undertaken using HLA*IMP software and compared to serologically typed alleles. A series of GWAS analyses were performed to investigate different genetic models and the renal transplant outcomes of interest. Graft survival genome-wide analyses were performed using Cox proportional hazards modeling for (1) donor SNP genotype main effects; (2) recipient SNP genotype main effects; (3) donor-recipient SNP genotype interaction effects (1df and 3df tests); and (4) CNV-tag-SNPs genotype mismatch effects (2 different models). Acute rejection genome-wide analyses were performed using logistic regression for models (1)-(3). An end-stage renal failure genome-wide analysis was performed using logistic regression for model (2), and to take advantage of the opportunity, an intracranial hemorrhage genome-wide analysis was performed using logistic regression for model (1).

2.4 | Replication-phase genotyping and analysis

Replication DNA samples were received at King’s College London and sent to the Wellcome Trust Sanger Institute for replication analysis. A
replication panel of 139 SNPs (post-quality control) was tested based on a combination of low P value (<10^-6) from the discovery phase, plus good support of association signals from SNPs in local linkage disequilibrium (LD), or on prior candidature from previous association studies. SNPs were tested according to the same model as motivated their inclusion in the replication panel (for example, if nominated based on a low P value for acute rejection in recipients, then that was also the test of interest in the replication analysis). Meta-analysis of discovery and replication results was carried out using inverse variance meta-analysis.27

For further details see Figure 1 and the Methods section in the Supplementary Material.

2.5 | Role of the funding source

The funding sources did not participate in the study design, collection, analysis or interpretation of the data, nor did they have a role in writing the report or the decision to submit for publication.

3 | RESULTS

3.1 | SNP association analysis

Despite the large size of our study (Table 1 and Table S2), none of the phenotypes and genetic models tested in the
discovery phase produced any LD-supported single-SNP results of genome-wide significance ($P \leq 5 \times 10^{-8}$). We also performed a partitioned heritability analysis via stratified LD score regression, which failed to reveal any significant enrichment of heritability in genomic regions marking tissue- or cell-type-specific activity (Figure 2).

We pursued a second "replication" phase in the hope that the most significant SNPs would be enriched for true association signals. One
hundred thirty-nine SNPs (post-quality control) were tested, based either on a low P value from the discovery phase plus LD support or on prior candidature from previous association studies. In general, the distribution of replication P values for all the tests did not depart appreciably from that expected under a global null hypothesis (Table S3). There was some enrichment for low P values in the recipient genotype main effect tests for acute rejection in the 12 months following transplantation. However, the P values from meta-analysis (across both discovery and replication phases) did not reach genome-wide significance ($P \leq 5 \times 10^{-8}$). The single SNP with strongest evidence for association was rs2289887 (Figure S1 and Table S4), which had consistent effects in both cohorts but a meta-analysis P value of only 0.00011, indicating that further studies are needed to establish the validity of this signal.

We collated all previously published association signals for early graft rejection and long-term allograft survival and none of these replicated in our study (Table S1).

### 3.2 | HLA mismatch analysis

We took advantage of recorded serological information to check for known associations with donor-recipient mismatch levels and to compare recorded information with imputed mismatches based on SNP genotype information (Figure 3). As expected for deceased donor transplants, the number of non-zero mismatch transplants in our data was low, reducing our power to detect associations. Nevertheless, we confirmed significant associations with HLA-A ($P = 0.022$) and HLA-DRB ($P = 0.00049$) mismatches using the recorded data.

On the other hand, the imputed mismatch results did not reproduce the expected associations. We attribute this to 2 factors: (1) the true mismatch = 2 frequency was low, due to donor selection; (2) as with any statistical noise, the imputation inaccuracy tended to act with disproportionate effect in the extremes of the distribution (here, the mismatch = 2 group), resulting in higher relative errors in that group. Thus, although the overall imputation accuracy was high (Tables S5 and S6), the rate of false positives was disproportionately high in the mismatch = 2 categories, swamping our ability to detect the true association signals (Table S7).

### 4 | DISCUSSION

In this article, we report the results of the first large-scale GWAS in renal transplantation. Despite our considerable sample size, we did not replicate any proposed findings from previous candidate gene studies nor did we discover any convincing new variants in our own analyses. There are a number of plausible reasons that may explain this.

Firstly, although this was a study involving thousands of individuals, by GWAS standards it was at the lower end of the range of sample sizes that have been employed for other human traits. A decade of GWAS across multiple complex traits has shown that single effect sizes for any 1 causal variant are typically low, and thus for some traits even bigger sample sizes than ours are needed to discover them. For example, the first robustly associated locus for schizophrenia was found in a discovery GWAS cohort of 3322 cases and 3587 controls. The number of reliably associated signals for schizophrenia has now grown

![FIGURE 3 Kaplan-Meier plots of graft survival by number of mismatches by serological typing (A, B, and C), imputed 2-digit resolution (D, E, and F), and imputed 4-digit resolution (G, H, and I). P values were obtained from likelihood ratio tests on Cox proportional hazards models](image-url)
to 108, providing new biological insights into the disease, thanks to a meta-analysis that involved 40 000 cases and 113 000 controls.32

Secondly, part of our study sought to look for recipient-donor genetic interactions. Interaction effects require even larger sample sizes to be reliably discovered. With a few notable exceptions, GWAS studies on other traits have been unsuccessful in discovering reliable interaction effects.

Thirdly, and perhaps most importantly, the transplantation outcomes we considered were relatively crude measures obtained retrospectively from national registry data, collected over many years for reasons other than for acting as endpoints in a genome-wide association study. Our outcomes were therefore heterogeneous, subject to missingness, and subject to environmental factors that likely weakened the genetic signals. Our data, for example, arising from different treatment protocols over time and space. Genetic data can also be subject to nonrandom missingness, the batch effect structure would need to be the same in both the discovery and replication cohorts for a signal to be replicated.

Batch effects are also a concern. Both the discovery and replication data were obtained from multiple different collection points in different countries over many years, and thus our phenotypes may be subject to batch effects, for example, arising from different treatment protocols over time and space. Genetic data can also be subject to batch effects, for example, arising from differences in sample collection protocols and unknown differences in population structure. We applied both covariate selection and genetically derived principal component axes to try to mitigate such effects. We also note that, as with nonrandom missingness, the batch effect structure would need to be the same in both the discovery and replication cohorts in order for false-positive association signals to be replicated. Nevertheless, together these extraneous factors may well have increased the noise in our data, and so reduced (and made less detectible) the genetic effects in our data.

HLA imputation from chip-based genetic data might in time be of value as an adjunct to serological typing. However, our investigations found that the additional noise introduced by imputation uncertainty prevented the imputed data from picking up the well-known mismatch signals at HLA-A and HLA-DRB. This suggests that the accuracy of HLA imputation will need to be improved before it can be used reliably as an alternative to serotyping.

In contrast, our serological data successfully detected the HLA-A and HLA-DRB mismatch signals, despite the reduced power to do so resulting from HLA-matching of deceased donor allografts. This indicates that, despite the various shortcomings of our study, strong HLA signals were detectable in our data. The implication therefore, at least as far as our primary graft survival endpoint is concerned, is that signals outside the HLA region are weaker than those already established in the HLA region. We therefore anticipate that genetic effects outside of the HLA region are more likely to be of value in elucidating biological pathways than in direct clinical prediction.

We explored this last point further via formal power calculations. These indicated that we were well powered to detect any main effect graft survival association signals involving causal SNPs with allelic hazard ratios in the range 1.4-1.9 (log-additive risk model, alpha = 5 × 10⁻⁸, power = 0.8, minor allele frequency>0.05), and to detect main-effect acute rejection association signals with allelic odds ratios in the range 1.7-2.9. We emphasize that these effect sizes are applicable to the traits investigated in this study, but that larger effects might be found in future studies under more precise phenotyping.

In summary, while our study was able to replicate known mismatch signals in the HLA region, we failed to find convincing association signals outside of the HLA region. Both phenotype heterogeneity and sample size may have contributed to this result. Looking ahead, we note that the general lessons from GWAS applied to multiple human traits over more than a decade have brought home 3 clear messages.29,30 The first is that all complex traits contain a genetic component, and harbor a large number of causal variants throughout the genome. The second is that larger GWAS studies, often obtained via meta-analysis with previous studies, inevitably succeed in discovering some portion of these causal variants. The third is that when a large enough portion of these variants is discovered, new insights into the biology of the trait are gained. With these points in mind, we look forward to an international consortium (iGeneTrain34), which has been formed to share and meta-analyze genetic and phenotypic data from most major transplant cohorts worldwide. We also look forward to efforts to collect more detailed phenotypes of relevance to transplant failure, which should provide greater genetic resolution. The data provided by our study form a foundation for ongoing efforts seeking to uncover the biology and improve the prospects for renal transplantation outcomes.

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AUTHOR CONTRIBUTIONS


DISCLOSURE

The authors of this article have conflicts of interest to disclose as described by the American Journal of Transplantation. M.E.W. is an employee of Genomics plc, a company providing genomic analysis services to the pharmaceutical and health care sectors. M.H.F. and R.B. are employees of UCB Celltech, a pharmaceutical company. Their involvement in the conduct of this research was solely in their capacity as academics at King’s College London.

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SUPPORTING INFORMATION

Additional Supporting Information may be found online in the supporting information tab for this article.