RESEARCH ARTICLE

The relationship between donor-recipient genetic distance and long-term kidney transplant outcome [version 1; peer review: 3 approved]

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Abstract

Background: We set out to quantify shared genetic ancestry between unrelated kidney donor-recipient pairs and test it as a predictor of time to graft failure.

Methods: In a homogenous, unrelated, European cohort of deceased-donor kidney transplant pairs (n pairs = 1,808), we calculated, using common genetic variation, shared ancestry at the genic (n loci=40,053) and genomic level. We conducted a sub-analysis focused on transmembrane protein coding genes (n transcripts=8,637) and attempted replication of a previously published nonsynonymous transmembrane mismatch score. Measures of shared genetic ancestry were tested in a survival model against time to death-censored graft failure.

Results: Shared ancestry calculated across the human leukocyte antigen (HLA) significantly associated with graft survival in individuals who had a high serological mismatch (n pairs = 186) with those who did not have any HLA mismatches indicating that shared ancestry calculated specific loci can capture known associations with genes impacting graft outcome. None of the other measures of shared ancestry at a genic level, genome-wide scale, transmembrane subset or nonsynonymous transmembrane mismatch score analysis were significant predictors of time to graft failure.

Conclusions: In a large unrelated, deceased-donor European ancestry renal transplant cohort, shared donor-recipient genetic ancestry, calculated using common genetic variation, has limited value in...
predicting transplant outcome both on a genomic scale and at a genic level (other than at the HLA loci).

**Keywords**
shared genetic ancestry, identity-by-state, identity-by-descent, transplant, kidney transplant, graft failure, HLA

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List of Abbreviations

eGFR, estimated glomerular filtration rate; HLA, human leukocyte antigen; IBD, identity by descent; IBS, identity by state; SNP, single nucleotide polymorphism; UKIRT, United Kingdom and Ireland Renal Transplant Consortium; GWAS, Genome-wide association study; PCA, principal components analysis.

Introduction

Organ allocation is a major issue in the field of transplantation. It is vital that organs are matched in a way that facilitates the best possible outcome for the recipient and for the transplanted organ, so that the organ goes to the person for whom it will last the longest. For many decades, renal transplants have been allocated largely on the basis of preferential HLA scoring, with reports dating back as far as 1965 showing the impact of HLA matching on graft outcome. A perfect HLA “match” does not guarantee a successful outcome and in individuals with zero HLA mismatches, acute rejection and early graft failure are still present. This indicates that other, potentially genetic, factors influence graft outcome and the success of the transplant.

There are large differences in graft survival between ethnicities, with African Americans having the poorest graft survival rates. However, it is unclear to what extent these differences are attributable to genetic, socioeconomic, and/or behavioural factors. A recent study of donor and recipient kidney transplant pairs found that ethnically matched transplants had significantly better graft survival, when compared to non-ethnically matched pairings. However this significance disappeared when adjustment was made for clinical variables including HLA mismatch, sensitization, sex, age and cold ischaemic time. Thus, the impact of shared ancestry between the kidney donor and recipient on graft outcome remains unclear.

Recent studies have developed distance scores based on exome sequencing data which are reported to predict graft outcome. For example, one study created an allogenomic mismatch score using non-synonymous variants from transmembrane protein genes. In a cohort of 53 donor-recipient kidney transplant pairs found that ethnically matched transplants had significantly better graft survival, when compared to non-ethnically matched pairings. However this significance disappeared when adjustment was made for clinical variables including HLA mismatch, sensitization, sex, age and cold ischaemic time. Thus, the impact of shared ancestry between the kidney donor and recipient on graft outcome remains unclear.

In this study, we set out to investigate the impact of shared genetic ancestry between donor-recipient transplant pairs, on graft outcome. We quantified shared genetic ancestry at the genomic level, genic level and across a specific subset of transmembrane protein encoding genes and tested these shared ancestry measures as a predictor of time to graft failure in a population of unrelated, European, deceased-donor kidney transplant pairs.

Methods

Cohort

Kidney donor and recipient pairs (n=2,094) were recruited via the United Kingdom and Ireland Renal Transplant Consortium (UKIRT). All transplants were deceased donor grafts and paired on the basis of preferential HLA matching and recipient waiting time. Transplants in the UKIRT cohort took place between 1981–2007. Only data on the first transplant was considered. All individuals were GWAS genotyped as part of the UKIRT and the Wellcome Trust Case Control Consortium 3 (see 11 for further details).

The following inclusion/exclusion criteria were applied: participants must have 1) been of European ancestry (determined via principal components analysis (PCA) – see Extended data, Supplementary methods for details) 2) have donated/received a renal transplant and 3) be unrelated to level of 3rd degree relative (PIHAT<0.1 calculated using PLINK v1.9).

Clinical predictors of graft survival were tested using STATA v13 in a Cox proportional hazards model and significant variables were included in the analysis as covariates. Acute rejection status was available for 1,099 individuals in our cohort. See Extended data, Supplementary methods for details on imputation. Details on HLA typing were published elsewhere.

Ethics approval and consent to participate

Ethics approval was granted by the Hammersmith and Queen Charlotte’s & Chelsea Research Ethics Committee REC No 08/H0707/1.

Genome-wide shared ancestry

We quantified shared genetic ancestry between donors and recipients at a genome-wide scale using three methods.

Method 1: Identity by descent (IBD). IBD here refers to the amount of shared genetic material between two individuals that is inherited from a recent common ancestor. We quantified the proportion of the genome that is IBD between the donor-recipient pairs using the PIHAT function in PLINK. The data was pruned for linkage disequilibrium using a 1000-kb window size, 100-kb step size and r² threshold of 0.2.

Method 2: Identity by state (IBS). IBS quantifies the number of shared alleles between two individuals. Alleles that are IBS can be inherited from a recent common ancestor or shared
by chance. We quantified the proportion of shared alleles across the genome between each donor and recipient pair using the DST measure in PLINK\textsuperscript{13,14}. Data was pruned for linkage disequilibrium following the same criteria as outlined above for the IBD method.

**Method 3: Chromopainter.** We created a mosaic of an individual’s genome from the haplotypes of all the other individuals in the dataset using Chromopainter\textsuperscript{10}. We calculated the matrix of donated haplotype lengths and associated haplotype-donors, for each individual in our dataset (i.e. the “chromolength” output from Chromopainter). The correlation between the chromolength chunks (donated from each individual in our dataset) of the donor and recipient in a given donor/recipient pair was calculated using the “corr” function in Microsoft Excel (with a perfectly matched mosaic genome having a score of 1). The correlation was then used as the measure of shared genetic ancestry.

These three measures of shared ancestry were tested against time to graft failure using a Cox Proportional Hazards Model using the R survival package (version 2.41–3) and the cox.ph function\textsuperscript{15,16}. We analysed two different survival analysis models; in the ‘full’ model we included significant clinical predictors as covariates, in the ‘baseline’ model we excluded clinical covariates.

**Per genic region analyses**

We examined whether shared ancestry between donor and recipient pairs at any specific gene, calculated using the IBD PIHAT and IBS DST methods, was associated with graft survival. Hg19 (human genome assembly 19) coordinates of protein-coding loci were obtained using the UCSC Genome Browser Table Browser via the “Genes and Predictions” group on the USCS gene track\textsuperscript{17}. The coordinates with a 50-kb buffer region added were used to extract the imputed genotype information for each individual in each donor recipient pair using PLINK, creating a file per genic locus. The files were pruned for LD in PLINK (“--indep-pairwise” command) using 1000-kb window size, 100-kb step-size and r\textsuperscript{2} threshold of 0.2. The PIHAT and DST scores were calculated for each of the thinned genic region files using PLINK’s --genome command. The PIHAT and DST scores for each gene were then extracted for each donor recipient pair.

PIHAT and DST scores for each gene was tested against time to graft failure, adjusting for significant clinical covariates, using R’s survival package\textsuperscript{18}. We corrected for multiple testing via the Bonferroni method. The total number of loci tested was 40,053 but greater than 50% of these were transcripts coded from overlapping regions and so we only considered independent loci (n= 16,765) when correcting for multiple testing.

As sanity check for the genic approach, we assessed whether genic shared ancestry measures could detect the known effect of the HLA on transplant outcome. As the UKIRTC dataset is composed of donor recipient pairs who were paired by preferential HLA matching and thus skewed towards pairs with fewer mismatches, we focused on the subset of transplant pairs with 0, 5 or 6 serological HLA mismatches. Time to death censored graft failure was tested against the total serological HLA mismatch measure, and against the IBD and IBS measures, in a Cox proportional hazards model using R. As this is a replication of the HLA mismatch measures, we did not apply Bonferroni correction and used a standard significance threshold of p=0.05.

**Shared ancestry across transmembrane protein coding genes**

As a subgroup analysis, we focused on the scores for transmembrane protein coding genes specifically, calculated in donor-recipient pairs using the IBD PIHAT and IBS DST measures (see methods section 5.2). The list of transmembrane protein coding was obtained using the same methods described previously\textsuperscript{9}. PIHAT and DST scores were extracted for these loci from the per-genic region results (see methods section 5.3). The PIHAT score was summed across all of the transmembrane genes to create an overall score. This transmembrane PIHAT score was then normalized, so the mean was 0 and standard deviation was 1. The same process was repeated for DST. The normalized transmembrane DST and PIHAT scores were then tested against time to graft failure in a Cox proportional hazards model using the survival package in R. Models were adjusted for significant clinical predictors of time to graft failure (see Extended data, Supplementary Table 1\textsuperscript{15}).

We also analysed the same set of transmembrane genes applying a univariate approach, where the PIHAT/DST calculated for each gene was considered separately and tested against time to graft failure as per the previous paragraph. To correct for multiple testing within this sub-group analysis, each gene was corrected for as opposed to each transcript (as some genes had multiple overlapping transcripts). In total, we corrected for 4,873 tests.

Finally, we carried out an exact replication analysis of the nonsynonymous transmembrane mismatch score which was recently reported to predict time to failure\textsuperscript{6}. Following the protocol as described by Reindl-Schwaighofer et al.\textsuperscript{9}, we imputed the HLA eplet mismatches using HLA matchmaker. Next, using VEP we annotated the imputed genotypes. We used the DR-mismatch program\textsuperscript{2} to calculate the nonsynonymous transmembrane mismatch score. We also calculated the overall genotype mismatch score and the overall nonsynonymous mismatch score (i.e. not limited to transmembrane protein encoding genes). Matching the parameters of the analysis presented in 9, we censored graft survival to 10 years post-transplant and excluded individuals with less than 3 months of follow-up post-transplant (see Extended data, Supplementary Table 5\textsuperscript{2} for cohort description). We carried out a Cox proportional hazards model against time to death censored graft failure of the interquartile range of the nonsynonymous transmembrane mismatch score including genome-wide
mismatch score, genome-wide nonsynonymous mismatch score, HLA serological mismatch, HLA eplet mismatch, donor age, donor sex, transplant centre and IBD (using PIHAT score as measure). We also tested the nonsynonymous transmembrane mismatch score in a univariate model which was not adjusted for additional covariates.

Results
Graft failure data was available for 1,808 donor-recipient pairs in our cohort with a total of 399 failure events (death censored). The average time to death censored graft failure was 6.04 years (range, 37–9,021 days) and average follow-up in those without a failure event was 8.87 years (range, 89–9,038 days). In order to determine the influence of clinical covariates on time to graft failure in the UKIRTC dataset, we applied a Cox proportional hazards model (see Extended data, Supplementary Table 1[2]). Donor age and acute rejection at 1 year emerged from the Cox model as significant predictors of graft failure and therefore were included as covariates in our shared ancestry survival analysis.

Genome-wide shared ancestry
We set out to test the effect of the three different genome-wide shared ancestry measures (IBS, IBD and Chromopainter) on time to death censored graft failure using a Cox proportional hazard model.

These shared ancestry measures were successfully calculated on 1,815 donor-recipient pairs (see Extended data, Supplementary Figure 1[2] for distributions of Chromopainter, IBD and IBS measures). Of these pairs, 927 had no detectable IBD relationship (i.e. PIHAT = 0) with 888 pairs having a PIHAT greater than 0. Due to the large number of pairs with PIHAT = 0, we carried out subsequent analyses on two groups; 1) those pairs who shared an IBD of less than or equal to 3rd degree relative, excluding those with no detected IBD and 2) pairs with an IBD of less than or equal to 3rd degree relative, including those with no detected IBD (see Extended data, Supplementary Table 2[2]).

These shared ancestry measures were then tested against time to graft failure in a Cox proportional hazards model. None of the genome-wide shared ancestry measures were significant in our full (with clinical covariates) or baseline models (without clinical covariates, see Table 1).

Shared ancestry per genic region results
Given the lack of correlation between graft outcome and donor-recipient genome-wide-level shared ancestry measures, we next tested if the shared ancestry between a donor and recipient at any one particular gene could predict graft outcome.

The HLA is a known predictor of graft outcome. To determine if our shared ancestry measures could reproduce the known effect of the HLA on graft outcome, we tested the correlation between IBD and IBS scores calculated across the HLA region, and the clinically reported combined serological HLA mismatch score (at HLA-A, B and DR). Both IBD and IBS were highly significantly correlated with total serological mismatch scores (P = 3.57x10^{-22}/r^2 = 0.48 and 5.74x10^{-22}/0.47, respectively (see Figure 1 and Figure 2).

As donor recipient pairs in the UKIRTC were selected on the basis of preferential HLA matching, the data was skewed towards having a low number of HLA mismatches. In this context, we selected donor-recipient pairs who had either zero-, or 5/6 HLA mismatches for regression analyses testing HLA distance with graft outcome. In this subset of donor-recipient pairs, we found that the total serologically typed HLA mismatch was associated time to graft failure (p=0.009, N=186, N events = 40, HR = 1.22, SE = 0.08).

| Table 1. Genome-wide shared ancestry measures in time to graft failure Cox proportional Hazards model. Full model was adjusted for donor age and acute rejection (within 12 months post-transplant). Baseline model was not adjusted for additional covariates. |
|-------------------|----------------|----------------|----------------|----------------|----------------|----------------|
| Model             | HR             | lower          | upper          | P              | N              | N events       |
| Chromopainter corr| full           | 5              | 0.28           | 89.58          | 0.27           | 1099           | 179           |
| PIHAT             | full           | 1.50E-08       | 2.40E-20       | 8693.76        | 0.19           | 1099           | 179           |
| PIHAT - Zero =NA  | full           | 1.40E-13       | 2.70E-32       | 750000         | 0.18           | 512            | 82            |
| DST               | full           | 1.20E-16       | 2.30E-62       | 5.90E+29       | 0.49           | 1099           | 179           |
| Chromopainter corr| baseline       | 4.64           | 0.78           | 27.52          | 0.09           | 1808           | 399           |
| PIHAT             | baseline       | 0.02           | 5.90E-10       | 599466         | 0.65           | 1808           | 399           |
| PIHAT - Zero =NA  | baseline       | 2.00E-08       | 4.30E-20       | 9457           | 0.2            | 885            | 202           |
| DST               | baseline       | 0.009          | 4.20E-35       | 1.70E+30       | 0.9            | 1808           | 399           |

HR, hazard ratio; 95% CI, upper and lower 95% confidence intervals; P, uncorrected p-value. PIHAT - Zero =NA, any donor recipient pair with a PIHAT score of 0 was removed from this analysis; Chromopainter corr, chromopainter correlation measure; N, number of donor-recipient pairs included in the analysis (i.e. had full clinical covariate data and PIHAT/DST scores available).
Figure 1. Correlation between total HLA mismatches and PIHAT across HLA ($p = 3.57 \times 10^{-223}$ and $r^2$ of 0.48). Total mismatches were calculated across HLA-A, B and DR using typing by serology. PIHAT (measure of IBD) was calculated for HLA-A, B and DR individual and combined by summing the PIHAT scores for each locus.

Figure 2. Correlation between total HLA mismatches and DST across HLA ($p = 5.74 \times 10^{-222}$ and $r^2 = 0.47$). Total mismatches were calculated across HLA-A, B and DR using typing by serology. DST (measure of IBS) was calculated for HLA-A, B and DR individual and combined by summing the DST scores for each locus.
Having reproduced the known predictive ability of HLA serological scores, we next compared, in the same group of individuals, the IBD and IBS HLA scores, to time to graft failure in a Cox proportional Hazards model. IBD and IBS scores were both significantly associated with time to graft failure with more shared IBD, or a greater amount of IBS associated with a longer time to graft failure (see Table 2). When we conditioned on serological HLA matches, the IBD/IBS signals were no longer significant, suggesting that the IBD/IBS scores are capturing similar variability as the serological typing.

The concordance statistic measures the proportion of pairs of individuals in which the individual with the higher-risk predictor had a failure event before the individual with the lower-risk predictor. We tested this statistic for the serological HLA score and the IBD/IBS scores (serological mismatch concordance = 0.56, standard error (SE) = 0.03; IBS across HLA concordance = 0.56, SE = 0.05; IBD across HLA concordance = 0.57, SE = 0.05). This indicates that IBS/IBD per gene measures can capture known associations with genes impacting graft outcome.

We then went on to test shared ancestry in individual genes across the entire genome. We compared the IBS and IBD measures calculated from each coding region in the genome (that contained at least one SNP after quality control measures were applied) to the afore-mentioned outcome measures. No gene reached genome-wide significance, for either the IBD or IBS measures (see Extended data, Supplementary tables 9–12).

The HLA-A (IBD p = 0.017 and IBS p = 0.045) and DR (IBD p = 0.009 and IBS p = 0.01) loci came out in the top 5% of the distribution for the time to graft failure analysis, but did not survive correction for multiple testing (see Supplementary table 3).

We also examined the per gene IBD and IBS scores in a baseline model which excluded clinical covariates. No single gene survived correction for multiple testing (n = 1,804, n events = 375; see Extended data, Supplementary tables 11–12).

Shared ancestry across transmembrane protein encoding genes results
Given previously published analysis showing a significant association between donor and recipient allele mismatches across genes encoding transmembrane proteins and graft outcome, we tested the association between the amount of shared genetic ancestry calculated across these transmembrane proteins and time to death censored graft survival. Neither the normalized transmembrane IBD nor IBS score were significant predictors of time to graft failure even before correction for multiple testing in our baseline (IBD p = 0.50, IBS p=0.35, n = 1,804, n events = 395) or full models (n= 1030, n events = 160, see Extended data, Supplementary tables 13–17).

Transmembrane genes were also analysed in a univariate model, where the IBD/IBS measure for each transmembrane gene was tested against time to graft failure. None of the transmembrane genes reached significance after correcting for multiple testing (corrected for 4783 tests, see Extended data, Supplementary tables 13–17).

Finally, we went on to attempt to replicate the findings of Reindl-Schwaighofer et al., who demonstrated a significant correlation between a transmembrane nonsynonymous donor-recipient mismatch score and time to ten-year and death censored graft failure. We did not find a significant correlation between in allogenomic mismatch score and time to graft failure in the univariate or multivariate (adjusted for additional covariates - see Methods section) models (see Extended data, Supplementary tables 7 and 8 for results).

**Discussion**
Our findings show that shared donor-recipient genetic ancestry, beyond 3rd degree relative, has limited value in predicting transplant outcome both on a genomic scale and at a genic level in deceased donor European ancestry renal transplant pairs. Our findings also suggest that IBD and IBS-based distance methods can be used to capture serological mismatch scores, and show comparative predictive ability.

Our analysis of shared ancestry at the genome-wide level indicates that, in a European-descent homogenous population, the level of shared ancestry between unrelated (defined here as <3rd degree relative) deceased donor and recipient pairs is not a clinically relevant predictor of graft survival. This is an important finding as it suggests that when choosing between deceased donors and recipients and where all of North-West European ancestral background, the level of shared ancestry (beyond 3rd degree) between donor and recipient is irrelevant.

| Table 2. Cox proportional hazards results: DST and PIHAT HLA measures vs time to graft failure. |
|---------------------------------|----------|----------|----------|-------|-------|----------|
| HR | Lower CI | Upper CI | P | N | N events |
| PIHAT HLA A, B, DR | 0.65 | 0.43 | 0.98 | 0.04 | 186 | 40 |
| DST HLA A, B, DR | 0.29 | 0.1 | 0.89 | 0.03 | 186 | 40 |

Outcome, time to graft failure; PIHAT HLA A, B, DR, PIHAT/DST between the given donor and recipient calculated across HLA A, B and DR loci; HR, hazards ratio; CI, 95% confidence interval; p, p-value; N, number of donor-recipient pairs included in the analysis; N events, number of failure events.
This study was carried out on a homogenous North West European ancestry population and therefore should not be generalized to admixed, or populations of different ancestry. Further work is required to determine if shared ancestry between donors and recipients impacts graft survival when the levels of shared ancestry has greater tendency for fluctuation within a given population, in particular admixed populations (for example, in African American communities).

We did not find any single gene that significantly associated with time to graft failure. This result indicates that, beyond the HLA, single gene shared-ancestry effects do not have a clinically relevant impact on graft survival. Notably, the IBD and IBS measures calculated at HLA-A and DR were found in the top 5% of genes most significantly associated with time to graft failure, but did not reach genome-wide significance. However, we found an effect of shared ancestry calculated across the HLA when we compared highly mismatched individuals for HLA serological typing, to those who were perfectly matched. We also showed a highly significant correlation between the total number of HLA serological mismatches and the amount of shared genetic ancestry calculated across the same HLA loci (A, B and DR). This demonstrates that these shared ancestry measures can pick up clinically relevant genetic differences between donors and recipients, and indeed have potential as a proxy for serological testing. This is in line with previous studies that showed known associations with the HLA region can be picked up using IBD measures\cite{9,10}. It is probable the HLA genes would have reached genome-wide significance in our genic analysis, if we had larger numbers of donor: recipient pairs and/or a cohort of transplant donors and recipients which had not been paired on the basis of preferential HLA matching.

Neither IBD nor IBS measures calculated across transmembrane protein coding loci were predictive of graft survival in our analysis. A previous exome mismatch study\cite{4} and genome-wide association study (GWAS) based analysis\cite{8} found that a score created via the quantification of specifically non-synonymous allelic mismatches at transmembrane and secreted proteins significantly predicted graft outcome whereas our shared ancestry measures considered all variation (i.e. including synonymous variation). We attempted to replicate the findings of the GWAS-based transmembrane nonsynonymous mismatch score\cite{9}; we did not see a significant correlation between the score and time to death censored graft survival with or without adjustment for additional covariates. However, we did see the same direction of effect as the original study. Potentially, differences between the cohort in this study and the original study may explain the lack of significance in our analysis. Our cohort was an older era and so was exposed to different immunosuppressants compared to that in the original study which was a more contemporary cohort. Further analysis with additional patient cohorts is required to determine whether this mismatch score will replicate in cohorts beyond that in the original report. The previous exome study\cite{4} captured both common and rare variation, whereas our study considered common variation only. It is also possible and indeed likely that advancements in immunosuppressants are suppressing the effect of shared ancestry.

**Conclusions**

In a large deceased-donor European ancestry renal transplant cohort, the amount of shared genetic ancestry between a donor and recipient had limited value in predicting transplant outcome both on the genome scale and at a genic level. However, the distance measures described in this study could describe a similar amount of the variability that is captured through serological HLA typing, and so have potential as an alternative to serological testing. Although we did not detect an effect of shared ancestry on graft outcome, studies capturing rare variation as well as studies examining admixed populations may uncover an important role of shared genetic ancestry on graft outcomes.

**Data availability**

**Underlying data**

The data used and/or analysed in the presented study was GWAS genotype chip array data and clinical data collected as part of the UKIRTC. Due to the sensitivity of the clinical patient data and genotype data used in this study, we are unable to share the data publicly. Access can be requested by researchers at accredited institutions. Access to data can be requested by emailing Peter J. Conlon (peterconlon@beaumont.ie) or Gianpiero L. Cavalleri (gcavalleri@rcsi.ie).

**Extended data**

Harvard Dataverse: Supplementary Information: The relationship between donor-recipient genetic distance and long-term kidney transplant outcome. https://doi.org/10.7910/DVN/ZXAXTN\cite{12}.

File ‘Supplementary Information - shared ancestry manuscript’ (DOCX) contains the following extended data:

- Affiliations for UKIRTC consortium members
- Supplementary methods
- Supplementary Tables 1–8
- Supplementary Figure 1

Supplementary Tables 9–16 are available as individual DOCX files as part of the repository project.

**Authors’ contributions**

CPS, GLC and PJC designed the analyses. CPS carried out the data analyses and wrote the manuscript. GLC and PJC supervised the data analysis and manuscript writing. GML coordinated the data collection and genotyping of the UKIRTC dataset. The UKIRTC members carried out the patient recruitment, clinical data collection and provided DNA samples/genotype information for patients. All authors reviewed and provided feedback for the manuscript.

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UK and Ireland Renal Transplant Consortium investigators include:


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Stapleton and colleagues use data on Deceased donor transplants from the UK/IR transplant registry to show that genome wide IBS/IBD do not independently associate with graft failure. They also do not find prior association with Austrian cohort data regarding total non Synonymous mismatches in genes predicted in a transmembrane or secreted location.

These are important data to report from this homogenous north west European cohort. These findings are different from Reindl-Schwaighofer et al, and Zhang et al, 2020 (not referenced - should be appropriately referenced)1.

- The homogeneity of the patient cohort used here could be depicted using a PCA plot based on the 1000 genome data. This would show difference between this and Zhang et al. report which was a clearly admixed cohort.
- An IBD distribution plot will also help visualize this.
- Details of the genotyping method used i.e. SNP chip etc. are missing. Were SNP chip versions different across data from this large /multicenter cohort?
- Was imputation performed? Details of pipeline used should be provided?
- Why were higher levels of relatedness excluded? Including them could still bring out a role for IBD that is greater than HLA mismatches?
- Is HLA serological mismatch used here? Both prior reports used PCR based mismatches. While PCR based data is also referenced later - it is unclear. Could have led to differences in findings.
In the correlation with HLA mismatch levels and within group comparisons the 0 vs 5/6 is shown. The role of IBS/IBD in other levels of mismatches should be tested. In these intermediate levels a greater role may be identified.

Overall valuable data to report.

References

PubMed Abstract | Publisher Full Text

Is the work clearly and accurately presented and does it cite the current literature? No

Is the study design appropriate and is the work technically sound? Yes

Are sufficient details of methods and analysis provided to allow replication by others? Yes

If applicable, is the statistical analysis and its interpretation appropriate? Yes

Are all the source data underlying the results available to ensure full reproducibility? Yes

Are the conclusions drawn adequately supported by the results? Yes

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: Kidney transplantation, long term graft damage, Donor recipient genetics.

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Reviewer Report 16 February 2021

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Krzysztof Kiryluk
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This is an interesting study that tests the role of genetic similarity between kidney donors and recipients in predicting long-term allograft outcomes. The study has notable strengths, including a large number of genotyped donor-recipient pairs used in the analysis and a comprehensive set of genetic similarity metrics tested as potential predictors of allograft survival. The methods are well described and the statistical models appear appropriate. The results were largely negative, leading to the conclusion that the degree of DR ancestry sharing (with the exception of the HLA region) was not significantly associated with time to allograft failure. The authors appropriately discuss the fact that these findings were based on a homogenous European cohort, and might not be generalizable to more diverse or admixed populations. Overall, the paper is informative and the negative results do provide a meaningful contribution.

I do have a few minor suggestions that could improve the manuscript:

1. Given that this study produced largely negative results, it would be important to provide a power analysis (estimate the probability of type II error) and discuss the range of detectable effect sizes given the cohort size, the available follow up time, and the observed event rates.

2. The main hypothesis of the study is that genetic donor-recipient dissimilarity would increase the risk of allograft failure by increasing the risk of rejection. It would therefore be informative to also include the analysis of rejection (any type), since rejection represents a more proximal outcome, and the study may have a better power to test for this association.

3. I am not entirely clear why the full model presented in Table 1 is adjusted for rejection events, since presumably rejection would mediate the effects on allograft survival. This model should probably be adjusted for HLA mismatch instead of rejection (to test for the effects of genetic mismatch that is independent of HLA, which I think would be a more meaningful hypothesis).

4. It would also be informative to see the following in the supplement: (1) histograms for the distributions of the key DR ancestry similarity metrics tested as predictors in the study, (2) PCA plot for the DR cohort in relationship to major continental ancestries, (3) QQ-plots for gene-based similarity tests. Please add HGNC-approved gene symbols to the Supplemental Tables. I would also consider replacing the term “genic regions” with “gene regions”.

Is the work clearly and accurately presented and does it cite the current literature?
Yes

Is the study design appropriate and is the work technically sound?
Yes

Are sufficient details of methods and analysis provided to allow replication by others?
Yes

If applicable, is the statistical analysis and its interpretation appropriate?
Yes

Are all the source data underlying the results available to ensure full reproducibility?
Are the conclusions drawn adequately supported by the results? Yes

**Competing Interests:** No competing interests were disclosed.

**Reviewer Expertise:** Human genetics.

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

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Dr. Stapleton and colleagues have produced a well-structured, well-written and refreshingly honest overview of genetic analyses undertaken to assess the relationship of shared genetic ancestry and time to graft failure in kidney donor recipients.

Sufficient background information relevant to the study has been provided. The methods are thoroughly described and potential limitations of the analyses are made very clear. The results are depicted clearly and discussed in a very balanced manner.

Is the work clearly and accurately presented and does it cite the current literature? Yes

Is the study design appropriate and is the work technically sound? Yes

Are sufficient details of methods and analysis provided to allow replication by others? Yes

If applicable, is the statistical analysis and its interpretation appropriate? I cannot comment. A qualified statistician is required.

Are all the source data underlying the results available to ensure full reproducibility? Partly

Are the conclusions drawn adequately supported by the results?
Yes

**Competing Interests:** No competing interests were disclosed.

**Reviewer Expertise:** Clinical genetics

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.