RESEARCH ARTICLE

The association of HLA-G polymorphisms and the synergistic effect of sMICA and sHLA-G with chronic kidney disease and allograft acceptance

Vanessa Hauer, Matilde Risti, Bruna L. M. Miranda, José S. da Silva, Ana L. Cidral, Carolina M. Pozzi, Fabiana L. de C. Contieri, Ibrahim A. Sadissou, Eduardo A. Donadi, Danillo G. Augusto, Maria da G. Bicalho

1 Laboratório de Imunogenética e Histocompatibilidade, Departamento de Genética, Universidade Federal do Paraná, Curitiba, Paraná, Brazil, 2 Setor de Transplante Renal, Hospital Universitário Evangélico de Curitiba, Alameda Augusto Stellfeld, Curitiba, Brazil, 3 Laboratório de Biologia Molecular, Departamento de Clínica Médica, Faculdade de Medicina de Ribeirão Preto da Universidade de São Paulo, Ribeirão Preto, São Paulo, Brazil, 4 Laboratório de Genética Molecular Humana, Departamento de Genética, Universidade Federal do Paraná, Curitiba, Paraná, Brazil, 5 Departamento de Ciências Biológicas, Universidade Estadual de Santa Cruz, Ilhéus, Bahia, Brazil

*These authors contributed equally to this work.

Abstract

The HLA-G and MICA genes are stimulated under inflammatory conditions and code for soluble (sMICA and sHLA-G) or membrane-bound molecules that exhibit immunomodulatory properties. It is still unclear whether they would have a synergistic or antagonistic effect on the immunomodulation of the inflammatory response, such as in chronic kidney disease (CKD), contributing to a better prognosis after the kidney transplantation. In this study, we went from genetic to plasma analysis, first evaluating the polymorphism of MICA, NKG2D and HLA-G in a cohort from Southern Brazil, subdivided in a control group of individuals (n = 75), patients with CKD (n = 94), and kidney-transplant (KT) patients (n = 64). MICA, NKG2D and HLA-G genotyping was performed by polymerase chain reaction with specific oligonucleotide probes, Taqman and Sanger sequencing, respectively. Levels of soluble forms of MICA and HLA-G were measured in plasma with ELISA. Case-control analysis showed that the individuals with haplotype HLA-G*01:01/UTR-4 have a lower susceptibility to develop chronic kidney disease (OR = 0.480; p = 0.032). Concerning the group of kidney-transplant patients, the HLA-G genotypes +3010 GC (rs1710) and +3142 GC (rs1063320) were associated with higher risk for allograft rejection (OR = 5.357; p = 0.013 and OR = 5.357, p = 0.013, respectively). Nevertheless, the genotype +3010 GG (OR = 0.136; p = 0.041) was associated with kidney allograft acceptance, suggesting that it is a protection factor for rejection. In addition, the phenotypic analysis revealed higher levels of sHLA-G (p = 0.003) and sMICA (p < 0.001) in plasma were associated with the development of CKD. For patients who were already under chronic pathological stress and underwent a kidney transplant, a high sMICA (p = 0.001) in pre-transplant proved to favor immunomodulation and allograft acceptance. Even so, the association of genetic factors with differential levels of soluble
molecules were not evidenced, we displayed a synergistic effect of sMICA and sHLA-G in response to inflammation. This increase was observed in CKD patients, that when undergo transplantation, had this previous amount of immunoregulatory molecules as a positive factor for the allograft acceptance.

**Introduction**

Previous studies on the major histocompatibility complex (MHC) genomic region identified genes that are important for immune regulation [1–3]. Among these genes are HLA-G (human leukocyte antigen-G) and MICA (major histocompatibility complex class I chain-related gene A). The few studies that concomitantly evaluated those genes left questions to be clarified about their functions [4,5]. HLA-G and MICA are highlighted, because they are produced in inflammatory and pathological conditions [5,6], can be expressed on cell membranes and reach distant immunological targets when in the form of soluble isoforms (sHLA-G and sMICA) [7,8].

HLA-G is expressed in regulatory T-cells and endothelial cells [9]; its expression has also been observed in transplanted specimens and associated with better graft survival [10–14]. The immunomodulatory role of HLA-G is performed mainly through interaction with inhibitory receptors, such as the leukocyte Ig-like receptor family B member 1 (LILRB1) and member 2 (LILRB2) [15]. Soluble HLA-G induces regulatory mechanisms, such as apoptosis of CD8+ T and NK cells, inhibition of B-cell proliferation, differentiation, and Ig secretion [16]. The membrane-bound HLA-G1 and the secreted soluble HLA-G5 are the most widely investigated isoforms [7,8].

Considering the immunomodulatory role of HLA-G, its allelic variation may impact its function on immunoregulation [17]. For example, the G*01:04 allele group has been associated with increased expression levels of sHLA-G in kidney-transplant patients without acute rejection [18]. Still, the 3′-UTR HLA-G region (exon 8) has nine polymorphisms with cumulative effect towards differential levels of sHLA-G [14,19].

MICA is expressed in various cells, including the thymic medulla and the gastrointestinal epithelium [6,20–24]. The best-described interaction of MICA occurs with the natural killer group 2 member D ligand (NKG2D). This receptor is expressed in natural killer (NK) cells, γδ T-cells, and αβCD8+ T-cell membranes [25].

Functionally, the interaction of NKG2D with the membrane MICA can culminate in cytotoxic activity of NK cells, but this signal can be counteracted by HLA-G interaction with LILRB1 inhibitory signal [4]. However, MICA as a soluble isoform may induce internalization and degradation of NKG2D [20]. This can originate an arrest of CD4+ T cells cycle in the G0-G1 phases, as well as induce proliferation of immunosuppressive NKG2D+CD4+ T cells, able to act in a paracrine way, inducing growth arrest of other T cells, secretion of IL-10 and TGF-β, similarly to the T regulatory type 1 cell cytokine profiles [26].

The number of GCT repetitions at exon 5 generates five different MICA alleles (MICA-A4, A5, A5.1, A6, and A9). The A5.1 allele (dbSNP: rs41293539) contains five GCT repetitions and one insertion, which leads to a frameshift mutation and generates a truncated protein with a differential expression, cellular localization [27] and was related to a bad outcome in kidney transplantation [28]. The MICA-129 Val/Met polymorphism (dbSNP: rs1051792) at nucleotide 454 (G>A) leads to a substitution, Val129Met in the α2 domain of the MICA protein [25]. The MICA-129 Val/Met variant can affect NKG2D binding avidity, leading to an alteration in
the immune response mediated by NK cells, which also depends on the level of MICA expression [29,30]. The NKG2D gene has two haploblocks (set of haplotypes), hb1 and hb2. The first haploblock (hb1) can be discriminated in high and low natural cytotoxic activity (HNK1 and LNK1, respectively). HNK1 (described by the main dbSNP, rs1049174-G) is associated with higher activity of NK cells in the peripheral blood and a lower incidence of cancers originating from epithelial cells [25,31].

The immunomodulatory effects of MICA with NKG2D and HLA-G, together with the scarcity of data on the diversity of genes and their influence on the composition of these soluble molecules, stimulated the development of this study. We hypothesized, that genetic variation in these genes may be associated with differential levels of plasmatic sHLA-G and sMICA. These immunological alterations would be related to the susceptibility of developing chronic kidney disease and with the establishment of the immunological balance or rejection after renal transplantation. Here, we evaluated the impact of MICA, NKG2D and HLA-G genotypes as well as the sMICA and sHLA-G levels, comparing patients with chronic kidney disease with controls and patients who had undergone a kidney transplant and developed rejection with those who did not.

**Methods**

**Sample**

The variation of HLA-G, MICA and NKG2D was investigated in individuals from the state of Paraná, Southern Brazil. This sample was composed of 169 individuals, divided between a control group of individuals in homeostasis, without chronic kidney disease or other uncontrolled inflammatory disease (Ct, n = 75) and patients in the end stage of chronic kidney disease (CKD, n = 94). Of the CKD, a total of 64 patients (in the end stage of renal disease, with a glomerular filtration rate under 15 mL/min, calculated following the Chronic Kidney Disease Epidemiology Collaboration equation [32]) had undergone kidney transplantation along 2012 and 2013, and had developed rejection (KTR, n = 28), or had not (KTN, n = 36). The information regarding the development or not of kidney allograft-rejection episodes (such as acute cellular rejection, chronic allograft nephropathy, and/or chronic glomerulonephritis) was obtained up to 4 years after transplantation.

The characteristics of the population (Ct, CKD and KT patients) are described in Table 1. All participants gave their signed informed consent before blood collection. The study was approved by the Ethics Committee of the Federal University of Paraná, Brazil and the Evangelical University Hospital of Curitiba, Paraná, Brazil (protocol number: CAAE 53627315.0.0000.0102).

**Immunosuppression protocol from kidney-transplant patients**

For all KT patients, pre-transplant immunosuppression was based on intravenous corticosteroid administration (methylprednisolone, 500 mg, independently of weight). Patients with increased immunological risk for allograft rejection received induction therapy with anti-thymocyte globulin (ATG). Maintenance therapy was composed of three medications: a corticosteroid (prednisone), a calcineurin inhibitor (tacrolimus), and an antimetabolite agent (mycophenolate mofetil, which was replaced by mycophenolate sodium after discharge).

**HLA-G, MICA and NKG2D genotyping**

Blood samples were collected in tubes containing ethylene-diamine-tetra-acetic acid (EDTA), and DNA was extracted from buffy-coat samples, using a salting-out technique [33].
The sequencing of HLA-G gene was performed by Sanger method. We used the ABI Prism Big Dye Terminator v3.1 Cycle Sequencing kit (Applied Biosystems, CA, USA), and analyzed the data in ABI Prism SeqScape software version 2.7 (Applied Biosystems). Exons 2, 3 and 4 were amplified and sequenced using the following primers: i) exon 2, F2E: GGGTCGGGCGGGTGCTCAA, and R2E: TCCGTGGGCGATGGAGGT, ii) exon 3, F3E: CCCACGACCCTACCTGGAGA, R3E: CTCCTCTTGCTGGATGGCGGCTG, iii) exon 4, F4E: CCATGAGAGAATCGAAGTGCT and R4E: TGCTTTCCCTACAGACATGAT, as adapted from a previous study [34]. All

| Characteristics | Ct (N = 75) | CKD (n = 94) | p   | KTN (n = 36) | KTR (n = 28) | p   |
|-----------------|------------|--------------|-----|-------------|--------------|-----|
| Male            | 34.70%     | 59.60%       | 0.002 | 47.20%      | 75.00%       | 0.023 |
| Age             | 37.88 years (SD = 14.46) | 46.77 years (SD = 13.44) | <0.001 | 44.92 years (SD = 14.27) | 46.32 years (SD = 13.67) | 0.692 |

**Ethnic group**

|               |               |               |     |               |               |     |
|----------------|---------------|---------------|-----|---------------|---------------|-----|
| Mixed race     | 10.80%        | 9.60%         | 1.000 | 8.30%        | 17.90%        | 0.448 |
| Asian          | 1.40%         | 2.10%         | 1.000 | 2.80%        | 0.00%         | 1.000 |
| Black          | 0.00%         | 6.40%         | 0.034 | 2.80%        | 7.10%         | 0.577 |
| Mulatto        | 0.00%         | 2.10%         | 0.503 | 0.00%        | 7.10%         | 0.187 |
| White          | 87.80%        | 79.80%        | 0.211 | 86.10%       | 67.90%        | 0.127 |

**Diseases**

|               |               |               |     |               |               |     |
|----------------|---------------|---------------|-----|---------------|---------------|-----|
| Hypertension   | 16.00%        | 90.30%        | <0.001 | 54.20%      | 45.80%        | 0.375 |
| Diabetes       | 2.70%         | 23.70%        | <0.001 | 19.40%      | 14.30%        | 0.743 |
| Dyslipidemias  | 2.70%         | 16.10%        | 0.004 | 11.10%      | 7.10%         | 0.688 |
| Glomerulonephritis | 0.00%     | 37.60%        | <0.001 | 36.10%      | 46.80%        | 0.450 |
| Hypertensive nephropathy | 0.00% | 14.90% | <0.001 | 44.40% | 55.60% | 0.488 |

**Transplant patients**

|               | KTN (n = 36) | KTR (n = 28) | p   |
|----------------|--------------|--------------|-----|
| Male donors    | 41.70%       | 50.00%       | 0.615 |
| Living donors  | 41.70%       | 25.00%       | 0.193 |
| Donor’s age    | 40.39 years (SD = 12.29) | 48.18 years (SD = 12.07) | 0.014 |
| Previous transplants | 19.40% | 14.30% | 0.743 |
| Previous blood transfusion | 50.00% | 35.70% | 0.313 |
| Multiparous women | 42.11% | 35.70% | 0.378 |
| Detection of DSA | 5.60% | 21.40% | 0.064 |

**HLA mismatches**

|                |               |               |     |               |               |     |
|----------------|---------------|---------------|-----|---------------|---------------|-----|
| HLA-A—0/1/2   | 25.00%/41.70%/16.70% | 10.70%/42.90%/35.70% | 0.141 |
| HLA-B—0/1/2   | 22.20%/50.00%/11.10% | 00.00%/50.00%/39.30% | 0.003 |
| HLA-DRB1—0/1/2 | 25.00%/50.00%/8.30% | 21.40%/42.90%/25.00% | 0.227 |
| Unknown        | 16.70%        | 10.7%         | 0.720 |

|                |               |               |     |               |               |     |
|----------------|---------------|---------------|-----|---------------|---------------|-----|
| Time on dialysis | 38 months (SD = 33) | 55 months (SD = 51) | 0.100 |
| GFR            | 7.58 mL/min (SD = 3.44) | 7.11 mL/min (SD = 4.86) | 0.662 |
| Cold ischemia time | 15h:31min (SD = 6h:46min) | 14h:44min (SD = 8h:21min) | 0.754 |

Ct: Control group. CKD: Patients with chronic kidney disease. KTN: Kidney-transplant patients with no rejection. KTR: Kidney-transplant patients who developed episodes of rejection. SD: standard deviation. DSA: donor-specific antibody. GFR: Glomerular filtration rate in pre-transplant, calculated following CKD-EPI (Chronic Kidney Disease Epidemiology Collaboration) equation.

1Analyses performed with adjustment for sex and age.

2Ethnicity information was obtained by self-assessment.

3Calculated from the total number of females in the group.

4Calculated only for deceased donors. The cold ischemia time for all living donors was a maximum of 1 hour.

https://doi.org/10.1371/journal.pone.0212750.t001
sequences were analyzed based on the official alleles listed in the International Immunogenetics Information System (IMGT) [35].

Amplification of the HLA-G 3'UTR was performed using the following primers: HLA-G8F: TGTGAAACAGCTGGCCCTGTT [36] and HGUT.R1: GTCTTCCATTTA TTTTGTCTCT [33]. PCR was carried out in a final volume of 25 μL, containing one unit of Hot Start Go Taq DNA-polymerase (Promega, Madison, WI, USA), 1× of Colorless GoTaq Flexi Buffer (Promega), 0.2 mM of each dNTP, 1.5 mM MgCl₂, 0.4 pmol of each primer, and 25 ng of genomic DNA. The initial denaturation was carried out at 94°C for 5 min, followed by 32 cycles of 95°C for 45 s, 57°C for 45 s and 72°C for 1 min, with a final extension step at 72°C for 7 min. This protocol, adapted from Castelli [37], was used for all samples.

Enzymatic purification of amplicons was performed and exon 8 was sequenced using the reverse primer to prevent sequence overlap in heterozygous genotypes due to the 14-bp indel (+2960, dbSNP: rs371194629). Including the 14-bp Ins/del, a total of 17 sites described in Table 2 were analyzed and individually annotated in the amplicon.

The reverse sequence-specific oligonucleotide (rSSO) typing was used as the first step in MICA genotyping, according to the manufacturer’s recommendations, as outlined in the LABType MICA kit (One Lambda Inc., Canoga Park, CA, USA). This reaction was analyzed through a flow fluorimeter (LABScan 100, Austin, TX, USA), which employs LUMINEX technology. Analyses were performed with HLA Fusion Software, and MICA genotypes were discriminated according to MICA-129 Val/Met polymorphism as to the presence or absence (termed wild type in relation to MICA A5.1 or Wt in this study) of MICA A5.1 allelic variation at exon 5.

The NKG2D gene was genotyped using TaqMan allelic discrimination methodology and KLRC4-KLRK1 TaqMan SNP Genotyping Assays (product number: C_9345347_10, Applied Biosystems). The SNP-based genotyping was specific to detect a G>C (dbSNP: rs1049174) replacement and discriminate the allelic haplotypes HNK1 and LNK1.

Table 2. The 17 sites of the HLA-G 3'UTR region analyzed in this study.

| rs          | 14-bp Del | 14-bp Ins | +2960 |
|-------------|-----------|-----------|-------|
| rs371194629 |           |           | +2960 |
| rs56747015  | C         | T         | +3001 |
| rs1707      | C         | T         | +3003 |
| rs1710      | G         | C         | +3010 |
| rs17179101  | C         | A         | +3027 |
| rs146339774 | G         | C         | +3032 |
| rs17179108  | C         | T         | +3035 |
| No SNPIdNo SNPId¹ | A | T | +3044 |
| No SNPIdNo SNPId¹ | C | T | +3052 |
| rs180827037 | G         | T         | +3092 |
| No SNPIdNo SNPId¹ | C | G | +3107 |
| No SNPIdNo SNPId¹ | G | A | +3111 |
| rs138249160 | T         | C         | +3121 |
| rs1063320   | C         | G         | +3142 |
| rs9380142   | A         | G         | +3187 |
| rs1610696   | C         | G         | +3196 |
| rs1233331   | G         | A         | +3227 |

¹Castelli et al. (2017). Del: +2960 or 14-bp deletion and Ins: +2960 or 14-bp insertion.
ELISA for sHLA-G and sMICA

Soluble HLA-G concentrations were evaluated by a specific sandwich ELISA in plasma, using MEM-G/9 [38], anti-human β2-microglobulin as capture and detection antibodies, respectively [39]. This reaction recognizes sHLA-G1 and HLA-G5 isoforms. Similarly, sMICA was measured using a commercial kit according to the manufacturer’s protocol (DuoSet MICA ELISA, R & D Systems, Minneapolis, MN, USA). Plasma samples and calibrators were added to each well (100 μL) in duplicate to perform sMICA and sHLA-G measurements. The final concentration was determined from the optical density compared to standard curves (SD). The lower limit of detection for sHLA-G was 6.25 ng/mL and for sMICA was 31.25 pg/mL.

Data analysis

Genotypes obtained for HLA-G (3’-UTR and coding sequence regions), MICA (MICA-129 Val/Met and MICA A5.1/Wt) and NKG2D (HNK1 or LNK1) were analyzed for different genetic parameters.

The ELB algorithm was used to infer haplotypes for HLA-G and MICA for each individual, performing two runs, considering each gene individually. HLA-G 3’-UTR haplotypes were denominated according to the most recent UTR nomenclature, described by Castelli et al. [19]. The best phase (with reliability above 95%) obtained for HLA-G and MICA genes separately and individually for each individual was then used to perform analyses including NKG2D genotypes. The frequencies of each variation were computed as absolute and relative frequency. Also, the Hardy-Weinberg equilibrium was tested by the exact test of Guo and Thompson [40], for the control group.

Linkage disequilibrium (LD) parameters (p or LOD, D’ and r²) were also estimated and calculated for intragenic variations of MICA and HLA-G in each group (Ct, CKD, KTN and KTR patients) and considering all individuals. All parameters were estimated using ARLEQUIN 3.5 software [41], and LD was also analyzed using Haploview software [42].

Fisher’s Exact Test (FET) was performed to infer associations among the analyzed genes (alleles and defined haplotypes) and two pathological conditions (first, the risk for chronic kidney disease and second, the risk for kidney allograft rejection). Logistic regression was chosen to confirm predictions observed in FET. The case-control analysis was first performed without adjustment for specific variables (e.g. patients’ sex frequencies, donor’s age and frequencies of HLA-B mismatches); after that, adjustments were made to confirm the observed associations.

Furthermore, the amounts of sHLA-G and sMICA obtained for plasma samples from Ct (n = 75) were compared with CKD patients (n = 56). Pre-transplant plasma samples from KTR and KTN patients (n = 26) were also compared. These analyses included a Mann-Whitney U test and median evaluations. A Kruskal-Wallis test was performed to analyze differences of expression among allelic variations. All association analyses were performed using the software IBM SPSS version 25.

Results

HLA-G and MICA haplotypes definition for association analysis and description of new HLA-G haplotypes

Three haplotype inferences were performed between: (Fig 1a) MICA-129 Val/Met with MICA A5.1/Wt; (Fig 1b) the 17 variations found in 3’-UTR of the HLA-G gene, after identification according to UTR nomenclature [19]; and (Fig 1c) HLA-G UTRs with alleles.

Four new haplotypes for HLA-G 3’-UTR were described, termed UTR-47NC, UTR-48NC, UTR-50NC and UTR-51NC (Fig 1d). Also, the following five previously described variations
were found to be monomorphic, but with no SNP id (identity): +3044 (A), +3052 (C), +3092 (G), +3107 (C) and +3111 (G). The most frequent HLA-G haplotypes inferred for HLA-G alleles with UTRs were compared to published data and did not differ in their frequencies (Fig 1e).

These analyses were also done for HLA-G and MICA; however, 51.76% of the inferred haplotypes had less than 95% reliability and therefore they could not be included in other analyses. Seven individuals did not achieve reliability in haplotype UTR inference and were excluded.

The other inferences originated haplotypes with 95% reliability or higher. The haplotypes formed by inferences were 21 for HLA-G UTR, 31 for HLA-G coding DNA sequence plus UTR, and 3 for MICA. This data was used to perform case-control and LD analysis. Still, the polymorphic variations observed for the 3'-UTR of the HLA-G gene, the HLA-G alleles, MICA variations, and NKG2D haploblocks were analyzed for Hardy-Weinberg Equilibrium in the control group, and all loci were in equilibrium (see S1 Table).

**Association of HLA-G genotypes in CKD and KT patients**

Case-control association analyses were performed through a contingency table to first detect significant relationships of alleles nucleotide variations, genotypes and/or previously defined haplotypes. All description of alleles and genotypes frequencies with their association results are respectively described in S2 and S3 Tables. Then, we applied the logistic regression analysis to endorse the primary association.
The HLA-G haplotype HLA-G*01:01 UTR-4 (p = 0.035) (Table 3) and the MICA-129 Val/Met genotype (p = 0.043, OR = 0.519, 95% CI = 0.280–0.962) genotype were found to be protective factors for the control group versus CKD patients, according to FET. After adjustments for age and sex, the described association with the MICA-129 Val/Met genotype was no longer significant (p = 0.131), the association with the HLA-G haplotype was maintained (p = 0.032, OR = 0.480, 95% CI = 0.199–0.961), although in the logistic regression analysis was found that neither association was significant.

For the KTR patients, the risk associations (Table 3) were found for genotypes +3187 AG (p = 0.037), +3142 CG (p < 0.001) and +3010 CG (p < 0.001). A protective association was found for +3010 GG (p = 0.041). After adjustments for the patients’ sex frequencies, donor’s age and HLA-B mismatches, the risk association of +3010 CG and +3142 CG was maintained (both with p = 0.013, OR = 5.357, 95% CI = 1.417–20.261), as well as a trend toward +3010 GG as a protective factor (p = 0.041, OR = 0.136, 95% CI = 0.016–1.178).

In the logistic regression (Table 4), including all loci, the genotype +3010 CG was confirmed as a risk factor and also +3010 GG as a protective factor for kidney allograft rejection (p = 0.004 for the categorical variable HLA-G +3010 included in predicted model). The model had a prediction power of 71.88% as is described in Table 5. This association was maintained even after adjustment (p = 0.038). Information concerning the observed and predicted frequencies for rejection by logistic regression can be seen in S4 Table. For the KTR patients was not detected association at the allelic level.

### LD evaluation of HLA-G and MICA genes among case-control groups

LD analyses were performed with the purpose of clarifying whether or not associations between polymorphisms would differ between the case-control groups (Ct and CKD, KTN and KTR). These associations could be determined by unique events arising from pathological states (CKD and KTR patients), contributing to an understanding of their origins. Therefore, this analysis was first performed separately for each group and compared. After the analysis was performed including all individuals.

For each group and for all sample, the estimated linkage disequilibrium did not diverge, nor did it diverge with respect to the variations in cis. All polymorphic sites found in 3’-UTR of the HLA-G gene showed a strong LD between them, except for the last polymorphism at position +3227.
Of all the analyses performed for each group (Ct, CKD, KTN and KTR), the highest LD was found in the HLA-G gene between positions +3010 and +3142, +3142 and 14-bp Ins/Del, and MICA-129 Val/Met and MICA A5/Wt, as summarized in Table 6 and extended in S5 Table.

Table 4. Logistic regression analysis of kidney-transplant patients for rejection.

| Predictor                  | β   | SE β | Wald x2 | p    | eβ (OR) | 95% CI for eβ |
|----------------------------|-----|------|---------|------|---------|--------------|
| HLA-G +3010 (Categorical variable) | -   | -    | -       | -    | -       | -            |
| HLA-G +3010 CG             | 1.664 | 0.630 | 6.969   | 0.008 | 5.278   | 1.535 to 18.148 |
| HLA-G +3010 GG             | -0.470 | 0.806 | 0.340   | 0.560 | 0.625   | 0.129 to 3.035 |
| Constant                   | -0.916 | 0.483 | 3.598   | 0.058 | 0.400   | -            |

Overall model evaluation
- -2 Log likelihood: 87.720 vs. 75.304
- Wald test: 0.995 vs. 3.598
- Coefficient constant: 0.251 vs. -0.916

Goodness-of-fit test
- Hosmer & Lemeshow: 1.000
- Cox and Snell R: 0.176
- Nagelkerke R2: 0.236

Binary Logistic Regression (method: forward stepwise conditional). Analysis performed with genotypes for KTN (coded as "0"; n = 36) versus KTR (coded as "1"; n = 28). Predicted logit (equation) of rejection = (-0.916) + (1.664) * (HLA-G +3010 CG) + (-0.470) * (HLA-G +3010 GG). The contrast of categorical variables and their references were respectively the indicator and the last subcategory according to codes described in available file "Data set.xlsx". HLA-G +3010 categorical variable had as reference HLA-G +3010 CC genotype. KTN: Kidney-transplant patients with no rejection. KTR: Kidney-transplant patients who developed episodes of rejection. OR: Odds Ratio. CI: Confidence Interval.

Table 5. The observed and predicted frequencies for rejection by logistic regression with cutoff of 0.50.

| Predicted | Observed |
|-----------|----------|
|           | No | Yes | % Correct |
| No        | 27 | 9  | 75.00     |
| Yes       | 9  | 19 | 67.86     |
| Overall % correct | 71.88 |

Sensitivity = 19/ (9+19) % = 67.86%. Specificity = 27/ (27+9) % = 75.00%. False positive = 9/ (9+27) % = 25.00%.

Table 6. The most significant results for HLA-G 3'-UTR and MICA linkage disequilibrium for all groups (Ct, CKD, KTN and KTR).

| Locus 1 | Locus 2 | D'   | LOD  | r²    | Variants in cis          |
|---------|---------|------|------|-------|--------------------------|
| +3010 G>C | +3142 C>G | 1.00 | 7.90 to 39.27 | 0.87 to 1.00 | +3010 C and +3142 G, +3010 G and +3142 C |
| +3142 G>C | +3142 G | 0.91 to 1.00 | 5.06 to 18.86 | 0.42 to 0.62 | 14-bp Ins and +3142 G, 14-bp Del and +3142 C |
| MICA-129 Val/Met | MICA A5/Wt | 1.00 | 2.22 to 6.71 | 0.19 to 0.27 | MICA-129 Met and MICA Wt, MICA-129 Val and MICA A5 |

The significant linkage disequilibrium had an LOD > 3.000 and p < 0.05. LOD: is the log of the likelihood odds ratio. Ct: Control group. CKD: Patients with chronic kidney disease. KTN: Kidney-transplant patients with no rejection. KTR: Kidney-transplant patients who developed episodes of rejection. Wt: wild type, which does not show MICA A5 variation. Del: +2960 or 14-bp deletion and Ins: +2960 or 14-bp insertion.
The LDs inferred for HLA-G alleles and UTRs were also strong, especially among the HLA-G*01:03 with UTR-5 or UTR-13, and the HLA-G*01:04 with UTR-3 (for all LD results see S5 Table). These haplotypes are also found in other populations [19].

**Soluble HLA-G and MICA phenotypes association**

The quantitative analyses of sHLA-G and sMICA were performed to evaluate differences between two conditions, comparing patients with chronic kidney disease (CKD) with controls (Ct) and patients who had undergone a kidney transplant and developed rejection (KTR) with those who did not (KTN) (Fig 2a and 2c).

![Image](https://doi.org/10.1371/journal.pone.0212750.g002)

**Fig 2. Case-control analysis of sMICA and sHLA-G production.** Analyses of sMICA and sHLA-G were performed for control (Ct) versus chronic kidney disease (CKD) (a-b), and after kidney transplant (KT) in patients with (KTR) and without (KTN) episodes of rejection (c-d). The production of regulatory molecules, sHLA-G and sMICA, is stimulated in a pathological condition, such as in patients with chronic kidney disease (CKD), but generally not in individuals in homeostasis, as observed in Ct. Once the CKD is established, the regulatory molecules in the pre-transplant can act in the post-transplant period as enhancers of immunoregulation, leading to allograft acceptance (e). Mann-Whitney test\(^1\), Chi-square test\(^2\). SD: Standard deviation.

https://doi.org/10.1371/journal.pone.0212750.g002
The first evaluation showed significantly higher levels of both molecules in the CKD group compared to the Ct group (sMICA median for CKD: 119.39 pg/mL and Ct: 63.54 ng/mL, \( p < 0.001 \); sHLA-G median for CKD: 31.62 pg/mL and Ct: 17.70 ng/mL, \( p = 0.003 \)). Also, the percentage of individuals for which neither molecule had no detection (under the limit of detection of 6.25 ng/mL for sHLA-G and 31.25 pg/mL for sMICA) was higher for the Ct compared to the CKD (Fig 2a and 2b).

Comparison between the KTN and KTR groups showed a difference for sMICA (median for KTN: 136.61 pg/mL and for KTR: 80.57 pg/mL, \( p = 0.001 \)) but no difference for sHLA-G (median for KTN: 42.63 ng/mL and for KTR: 32.19 ng/mL, \( p = 0.836 \)). The percentage of individuals with no detectable sMICA was higher in the KTR group (Fig 2c and 2d).

MICA haplotypes were analyzed in association with sMICA quantification, and no differences were detected (\( p = 0.327 \)). sHLA-G and genetic variations were evaluated for the most frequently detected alleles (HLA-G*01:01, HLA-G*01:04 and HLA-G*01:03) and UTRs (UTR-1, -2, -3, -4 and -5), but no differences were observed (respectively, \( p = 0.448 \) and \( p = 0.585 \), more information see S7 Table and S1 Fig).

**Discussion**

Case-control analysis included HLA-G, MICA and NKG2D genotype data and quantification of soluble molecules (sHLA-G and sMICA) involved in immunomodulation of inflammatory response. Regarding the association tests performed for chronic kidney disease patients, the HLA-G*01:01/UTR-4 was found to be a protective factor for the development of chronic kidney disease (FET analysis). In particular, UTR-4 has previously been related to higher stability of HLA-G transcripts and immunoregulation in successful pregnancies [43]. This suggest the HLA-G*01:01/UTR-4 could be a protection against inflammatory conditions and pathological states, not just found in women with secondary recurrent miscarriage, but also in CKD patients.

For the kidney-transplant patients, the +3010 CG genotype was associated with kidney allograft rejection. This SNP has the strongest LD with the +3142 position. Also, the 14-bp Ins/Del had high LD with +3142 and +3010 SNPs. The variations found in cis were related to their effect when inherited together, as previously described for HLA-G [44–49]. Thus, the composition possibly associated with the risk of kidney allograft rejection by the decrease of HLA-G expression would be +3010 C, +3142 G and 14-bp Ins, whereas +3010 G with +3142 C, and +3142 C with 14-bp Del would be associated with an increase.

The presence of +3010 C would act as a negative factor for the allograft outcome (even in the heterozygote), as it is linked to the +3142 G variation that promotes greater affinity of HLA-G transcripts to microRNAs, leading to a decreased HLA-G expression [50]. This effect would be more evident in the presence of 14-bp Ins, from which unstable transcripts originate [44]. The analyses were not able to detect a direct relation regarding the +3010 CC and rejection, but twice as many patients who developed episodes of rejection have this genotype compared to the frequencies of +3010 GG patients, which was associated as a protection factor. The association of +3142 with 14-bp Ins/Del has been the subject of many studies aiming to identify risk factors [45,46,49,51]. However, we suggest that +3010 is a marker to predict the simultaneous presence of +3142 G with 14-bp Ins.

Concerning the sMICA and sHLA-G production, those molecules functionally preserve the expression of a similar inflammatory condition [11,52]. Our analysis showed that both molecules are more highly expressed in patients in a stressful condition, such as chronic kidney disease (CKD), compared to the control. Also as expected, in the Ct group the non-detection of sMICA and sHLA-G was more common than in the CKD group, which reflects the
stimulation of immunological changes and production of regulatory molecules (sHLA-G and sMICA) under a pathological condition.

In relation to the kidney-transplant patients and soluble molecules, there was evidence of higher production of sMICA in the KTN compared to the KTR group, who developed rejection. Therefore, once the chronic disease is established in the patient, the production of regulatory soluble molecules in the pre-transplant period acts as an immunoregulatory effect contributing to the acceptance of the allograft (Fig 2e). Thus, the pre-transplant status of sMICA can potentially influence the successful outcome of the kidney transplantation. Particularly, the evaluation of MICA expression need to take into account its isoform (membrane bound or soluble) related to its function (activation or inhibition of NK cells), thus MICA mRNA levels [5] may be a confounding manner to evaluate this biomarker.

Regarding to the difference in the sMICA and sHLA-G production, no allelic differentiation was observed. This demonstrates that the joint analysis of other forms, such as those bound to the membrane, is necessary to perform this type of analysis.

Finally, considering the Brazilian population is one of the most admixed in the world [53], new genetic variations can be found throughout the country, as well as similarities. In this study, the MICA, NKG2D and HLA-G genes evaluation also comprised part of this diversity. For the HLA-G gene, a total of 21 UTRs were observed, four of them new. The HLA-G haplotype frequencies did not differ from data reported by Castelli et al. (2014) [54]. For MICA, 3 haplotypes were detected and the MICA-129 Met with MICA A5.1 was not present. Of the HLA-G haplotypes, the combinations of HLA-G*01:03/UTR-5 or /UTR-13 with HLA-G*01:04/UTR-3 were most frequent, which is in accordance with the main complete extended haplotypes previously described [19].

**Conclusion**

The increased production of immunoregulatory molecules, such as sMICA and sHLA-G, was associated with chronic kidney disease. Still, a higher sMICA production in the pre-transplant period was found to be associated with a better kidney allograft outcome.

Association of genetic factors with different quantifications of sMICA and sHLA-G already reported in other works [14,17,18,27–30], were not observed here. However, the +3010 SNP (rs1710) in HLA-G gene, showed potential as a molecular marker for the characterization of +3142 SNP and 14-bp Ins/Del variation, all related to changes in HLA-G production and to kidney allograft rejection. Thus, this finding indicate that phenotype and genotype association analysis should include quantification of immunoregulatory molecules in the cell membrane-bound and soluble isoforms.

Moreover, our study described new HLA-G UTR haplotypes, which reinforces the importance of conducting further studies that aim to better characterize the diversity of the Brazilian population.

**Supporting information**

S1 Table. Analyses of Hardy-Weinberg equilibrium performed for HLA-G alleles and polymorphisms in 3’-UTR region, MICA and NKG2D variations in the control group (n = 75). Positions +3001, +3032, +3044, +3052, +3092, +3107, +3111 and +3121 are monomorphic sites in the control group. 1HLA-G haplotypes: inferred composition through the ELB algorithm of HLA-G alleles with determined UTRs. Wt: wild type, which does not show MICA A5.1 variation. SD: Standard deviation. (PDF)
S2 Table. Observed allele frequencies and Fischer’s Exact Test results for the HLA-G, MICA and NKG2D genes. Ct: Control group. CKD: Patients with chronic kidney disease. KTN: Kidney-transplant patients with no rejection. KTR: Kidney-transplant patients who developed episodes of rejection. Wt: wild type, which does not show MICA A5.1 variation. Del: +2960 or 14-bp deletion and Ins: +2960 or 14-bp insertion. Zero or not sufficient frequency to perform the calculation.

S3 Table. Observed genotype frequencies and Fischer’s Exact Test results for HLA-G, MICA and NKG2D genes. Ct: Control group. CKD: Patients with chronic kidney disease. KTN: Kidney-transplant patients with no rejection. KTR: Kidney-transplant patients who developed episodes of rejection. Wt: wild type, which does not show MICA A5.1 variation. Del: +2960 or 14-bp deletion and Ins: +2960 or 14-bp insertion.

S4 Table. Predicted probability of rejection for kidney transplant patients.

S5 Table. The linkage disequilibrium results for HLA-G 3’-UTR and MICA in each group (Ct, CKD, KTN and KTR). Analysis performed in Haploview. The linkage disequilibrium (LD) considered significant had an LOD > 3.000. LOD: is the log of the likelihood odds ratio. 95% CI relative to D’: confidence interval. Ct: Control group. CKD: Patients with chronic kidney disease. KTN: Kidney-transplant patients with no rejection. KTR: Kidney-transplant patients who developed episodes of rejection. Wt: wild type, which does not show MICA A5.1 variation. Del: +2960 or 14-bp deletion and Ins: +2960 or 14-bp insertion.

S6 Table. The linkage disequilibrium results for HLA-G 3’-UTR and MICA in each group (Ct, CKD, KTN and KTR). The inferences of HLA-G haplotype through ELB algorithm of HLA-G alleles with determined UTRs. Analysis performed in Arlequin. The significant linkage disequilibrium had a p < 0.05. Ct: Control group. CKD: Patients with chronic kidney disease. KTN: Kidney-transplant patients with no rejection. KTR: Kidney-transplant patients who developed episodes of rejection.

S7 Table. Association analysis performed between soluble molecules and genetic factors. Q1: First Quartile. Q3: Third Quartile.

S8 Table. Data set.xlsx. (a) Data_set_CKD_Ct_n_169. (b) Code_book_Data_set_CKD_Ct. (c) Data_set_KTR_KTN_n_64. (d) Code_book_Data_set_KTR_KTN.

S1 Fig. Representation of median, first and third quartile from soluble molecules associated with genetic factors. Soluble MICA (sMICA) and MICA haplotypes (MICA-129 Val/Met and MICA A5.1/Wt) association (p = 0.327) (a). Soluble HLA-G and HLA-G alleles association (p = 0.448) (b). Soluble HLA-G and most frequent HLA-G UTRs association (p = 0.585) (c).

Acknowledgments

We are most grateful to the participants for generously agreeing to provide samples for this study. We also thank the Hospital Universitário Evangélico de Curitiba, especially
LIGH and the staff of the Clinical Immunology Division (Ribeirão Preto), for their technical support.

Author Contributions

Conceptualization: Vanessa Hauer, Matilde Risti, Danillo G. Augusto, Maria da G. Bicalho.

Data curation: Vanessa Hauer, Matilde Risti, Bruna L. M. Miranda, Ana L. Cidral, Ibrahim A. Sadissou, Eduardo A. Donadi, Maria da G. Bicalho.

Formal analysis: Vanessa Hauer, Matilde Risti, José S. da Silva, Ana L. Cidral.

Funding acquisition: Vanessa Hauer, Maria da G. Bicalho.

Investigation: Vanessa Hauer, Matilde Risti, José S. da Silva, Ana L. Cidral, Carolina M. Pozzi, Fabiana L. de C. Contieri, Eduardo A. Donadi, Danillo G. Augusto.

Methodology: Vanessa Hauer, Matilde Risti, Bruna L. M. Miranda, José S. da Silva, Ana L. Cidral, Ibrahim A. Sadissou, Eduardo A. Donadi.

Project administration: Vanessa Hauer, Maria da G. Bicalho.

Resources: Vanessa Hauer, Matilde Risti, Carolina M. Pozzi, Fabiana L. de C. Contieri, Eduardo A. Donadi, Maria da G. Bicalho.

Software: Vanessa Hauer, José S. da Silva, Ana L. Cidral, Danillo G. Augusto.

Supervision: Vanessa Hauer, Eduardo A. Donadi, Danillo G. Augusto, Maria da G. Bicalho.

Validation: Vanessa Hauer, Matilde Risti, José S. da Silva.

Visualization: Vanessa Hauer.

Writing – original draft: Vanessa Hauer, Matilde Risti.

Writing – review & editing: Vanessa Hauer, Bruna L. M. Miranda, Ana L. Cidral, Carolina M. Pozzi, Fabiana L. de C. Contieri, Ibrahim A. Sadissou, Eduardo A. Donadi, Danillo G. Augusto, Maria da G. Bicalho.

References

1. Mosaad YM. Clinical Role of Human Leukocyte Antigen in Health and Disease. Scand J Immunol. 2015; 82: 293–306. https://doi.org/10.1111/sji.12329 PMID: 26099424

2. Dendrou CA, Petersen J, Rossjohn J, Fugger L. HLA variation and disease. Nat Rev Immunol. Nature Publishing Group; 2018; 18: 325–339. https://doi.org/10.1038/nri.2017.143 PMID: 29292391

3. Barkal AA, Weiskopf K, Kao KS, Gordon SR, Rosenthal B, Yiu YY, et al. Engagement of MHC class I by the inhibitory receptor LILRB1 suppresses macrophages and is a target of cancer immunotherapy. Nat Immunol. 2018; 19: 76–84. https://doi.org/10.1038/s41590-017-0004-z PMID: 29180808

4. Menier C, Riteau B, Carosella ED, Rouas-Freiss N. MICA triggering signal for NK cell tumor lysis is counteracted by HLA-G1-mediated inhibitory signal. Int J cancer. 2002; 100: 63–70. https://doi.org/10.1002/ijc.10460 PMID: 12115588

5. Racca AL, Veaute CM, Bailat AS, Gaite L, Arriola M, Hajos SE, et al. Expression of HLA-G and MICA mRNA in renal allograft. Transpl Immunol. Elsevier B.V.; 2009; 21: 10–12. https://doi.org/10.1016/j.trim.2009.01.002 PMID: 19193353

6. Groh V, Bahram S, Bauer S, Herman A, Beauchamp M, Spies T. Cell stress-regulated human major histocompatibility complex class I gene expressed in gastrointestinal epithelium. Proc Natl Acad Sci U S A. 1996; 93: 12445–50. https://doi.org/10.1073/pnas.93.22.12445 PMID: 8901601

7. LeMacoult J, Le Discorde M, Rouas-Freiss N, Moreau P, Menier C, McCluskey J, et al. Biology and functions of human leukocyte antigen-G in health and sickness*. Tissue Antigens. 2003; 62: 273–284. https://doi.org/10.1034/j.1399-0039.2003.00143.x PMID: 12974794
8. Carosella ED, Moreau P, Lemaout J, Rouas-Freiss N. HLA-G: from biology to clinical benefits. Trends Immunol. 2008; 29: 125–32. https://doi.org/10.1016/j.it.2007.11.005 PMID: 18249584

9. Sheshgiri R, Rao V, Turniat LC, Xiao R, Prodgier JL, Badiwala M, et al. Progesterone induces human leukocyte antigen-g expression in vascular endothelial and smooth muscle cells. Circulation. 2008; 118: S58–64. https://doi.org/10.1161/CIRCULATIONAHA.107.757781 PMID: 18824770

10. Créput C, Durbach A, Menier C, Guettler C, Samuel D, Daussel J, et al. Human leukocyte antigen-G (HLA-G) expression in biliary epithelial cells is associated with allograft acceptance in liver-kidney transplantation. J Hepatol. 2003; 39: 587–94. https://doi.org/10.1016/S0168-8278(03)00354-4 PMID: 12971970

11. Qiu J, Terasaki PI, Miller J, Mizutani K, Cai J, Carosella ED. Soluble HLA-G expression and renal graft acceptance. Am J Transplant. 2006; 6: 2152–6. https://doi.org/10.1111/j.1600-6143.2006.01417.x PMID: 16780545

12. Brugière O, Thabut G, Pretolani M, Krawice-Radanne I, Dill C, Herbreteau A, et al. Immunohistochemical study of HLA-G expression in lung transplant recipients. Am J Transplant. 2009; 9: 1427–38. https://doi.org/10.1111/j.1600-6143.2009.02650.x PMID: 19656137

13. Almasood A, Sheshgiri R, Joseph JM, Rao V, Kamali M, Tumiati L, et al. Human leukocyte antigen-G is upregulated in heart failure patients: a potential novel biomarker. Hum Immunol. Elsevier Inc.; 2011; 72: 1064–7. https://doi.org/10.1016/j.humimm.2011.08.016 PMID: 21925559

14. Donadi EA, Castelli EC, Arnaiz-Villena A, Roger M, Rey D, Moreau P. Implications of the polymorphism of HLA-G on its function, regulation, evolution and disease association. Cell Mol Life Sci. 2011; 68: 369–95. https://doi.org/10.1007/s00018-010-0580-7 PMID: 21107637

15. Carosella ED, Rouas-freiss N, Roux DT, Moreau P, Lemaout J. HLA-G: An Immune Checkpoint Molecule [Internet]. 1st ed. Advances in Immunology. Elsevier Inc.; 2015. https://doi.org/10.1016/bs.ai.2015.04.001 PMID: 26073983

16. Naji A, Menier C, Morandi F, Aguque S, Maki G, Ferretti E, et al. Binding of HLA-G to ITIM-Bearing Ig-like Transcript 2 Receptor Suppresses B Cell Responses. J Immunol. 2014; 192: 1536–1546. https://doi.org/10.4049/jimmunol.1300438 PMID: 24453251

17. Dias FC, Castelli EC, Collares CVA, Moreau P, Donadi EA. The Role of HLA-G Molecule and HLA-G Gene Polymorphisms in Tumors, Viral Hepatitis, and Parasitic Diseases. Front Immunol. 2015; 6: 9. https://doi.org/10.3389/fimmu.2015.00009 PMID: 25699038

18. Misra MK, Pandey SK, Kapoor R, Sharma RK, Kapoor R, Prakash S, et al. HLA-G gene expression influenced at allelic level in association with end stage renal disease and acute allograft rejection. Hum Immunol. American Society for Histocompatibility and Immunogenetics; 2014; 75: 833–839. https://doi.org/10.1016/j.humimm.2014.06.005 PMID: 24929145

19. Castelli EC, Gerasimou P, Paz MA, Ramalho J, Porto IOP, Lima THA, et al. HLA-G variability and haplotypes detected by massively parallel sequencing procedures in the geographically distinct population samples of Brazil and Cyprus. Mol Immunol. Elsevier Ltd; 2017; 83: 115–126. https://doi.org/10.1016/j.molimm.2017.01.020 PMID: 28136696

20. Bauer S, Groh V, Wu J, Steinle A, Phillips JH, Lanier LL, et al. Activation of NK cells and T cells by NKG2D, a receptor for stress-inducible MICA. Science. 1999; 285: 727–9. https://doi.org/10.1126/science.285.5428.727 PMID: 10426993

21. Katsuyama Y, Ota M, Ando H, Saito S, Mizuki N, Kera J, et al. Sequencing based typing for genetic polymorphisms in exons, 2, 3 and 4 of the MICA gene. Tissue Antigens. 1999; 54: 178–84. Available: http://www.ncbi.nlm.nih.gov/pubmed/10488745

22. Romphruk A V, Naruse TK, Romphruk A, Kawata H, Puapairoj C, Kulski JK, et al. Diversity of MICA (PERB11.1) and HLA haplotypes in Northeastern Thais. Tissue Antigens. 2001; 58: 83–9. Available: http://www.ncbi.nlm.nih.gov/pubmed/11696220

23. Steinle A, Li P, Morris DL, Groh V, Lanier LL, Strong RK, et al. Interactions of human NKG2D with its ligands MICA, MICB, and homologs of the mouse RAE-1 protein family. Immunogenetics. 2001; 53: 279–87. https://doi.org/10.1007/s002510010032S PMID: 11491531

24. Muro M, Lopez-Hernandez R, Llorente S, Miguel Bolarin J, Martinez P, Boix F, et al. MICA Molecules in Disease and Transplantation, a Double-Edged Sword? Curr Immunol Rev. 2012; 8: 307–325.

25. Risti M, Bicalho M da G. MICA and NKG2D: Is There an Impact on Kidney Transplant Outcome? Front Immunol. 2017; 8: 179. https://doi.org/10.3389/fimmu.2017.00179 PMID: 28289413

26. Groh V, Smythe K, Dai Z, Spies T. Fas ligand-mediated paracrine T cell regulation by the receptor NKG2D in tumor immunity. Nat Immunol. 2006; 7: 755–762. https://doi.org/10.1038/ni.1350 PMID: 16732291

27. Tonnerre P, Gérard N, Chatelais M, Poli C, Allard S, Cury S, et al. MICA variant promotes allosensitization after kidney transplantation. J Am Soc Nephrol. 2013; 24: 954–66. https://doi.org/10.1681/ASN.2012080814 PMID: 23539759
28. Tonnerre P, Gérard N, Gavlovsky P-J, Mazairey S, Hourmant M, Cheneau M-L, et al. MICA Mutant A5.1 Influences BK Polyomavirus Reactivation and Associated Nephropathy After Kidney Transplantation. J Infect Dis. 2016; 214: 807–16. https://doi.org/10.1093/infdis/jiw168 PMID: 27130430

29. Raache R, Belanteuk K, Amroun H, Benyahia A, Heniche A, Azzouz M, et al. Association of major histocompatibility complex class 1 chain-related gene A dimorphism with type 1 diabetes and latent autoimmune diabetes in adults in the Algerian population. Clin Vaccine Immunol. 2012; 19: 557–561. https://doi.org/10.1128/CVI.05473-11 PMID: 22323559

30. Isemhagen A, Schilling D, Monecke S, Shah P, Elsner L, Walter L, et al. The MICA-129Met/Val dimorphism affects plasma membrane expression and shedding of the NKG2D ligand MICA. Immunogenetics. 2016; 68: 109–23. https://doi.org/10.1007/s00251-015-0884-8 PMID: 26585323

31. Espinoza JL, Takami A, Onizuka M, Sao H, Akiyama H, Miyamura K, et al. NKG2D gene polymorphism has a significant impact on transplant outcomes after HLA-fully-matched unrelated bone marrow transplantation for standard risk hematologic malignancies. Haematologica. 2009; 94: 1427–34. https://doi.org/10.3324/haematol.2009.008318 PMID: 19794085

32. Levey AS, Stevens LA, Schmid CH, Zhang YL, Castro AF, Feldman HI, et al. A new equation to estimate glomerular filtration rate. Ann Intern Med. 2009; 150: 604–12. https://doi.org/10.7326/0003-4819-150-9-200905050-00006 PMID: 19414839

33. Lahiri DK, Nurnberger JI. A rapid non-enzymatic method for the preparation of HMW DNA from blood for RFLP studies. Nucleic Acids Res. 1991; 19: 5444. https://doi.org/10.1093/nar/19.19.5444 PMID: 1681511

34. Nardi F da S, Slowik R, Wowk PF, da Silva JS, Gelmini GF, Michelon TF, et al. Analysis of HLA-G Polymorphisms in Couples with Implantation Failure. Am J Reprod Immunol. 2012; 68: 505–514. https://doi.org/10.1111/ajr.12001 PMID: 23009094

35. Robinson J, Halliwell JA, Hayhurst JD, Flice P, Parham P, Marsh SGE. The IPD and IMGT/HLA database: allele variant databases. Nucleic Acids Res. 2015; 43: D423–31. https://doi.org/10.1093/nar/gku1161 PMID: 25414341

36. Bermingham J, Jenkins D, McCarthy T, O’Brien M. Genetic analysis of insulin-like growth factor II and HLA-G in pre-eclampsia. Biochem Soc Trans. 2000; 28: 215–9. Available: http://www.ncbi.nlm.nih.gov/pubmed/10816131 PMID: 10816131

37. Castelli EC, Mendes-Junior CT, Deghaide NHS, de Albuquerque RS, Muniz YCN, Simões RT, et al. The genetic structure of 3’ untranslated region of the HLA-G gene: polymorphisms and haplotypes. Genes Immun. 2010; 11: 134–41. https://doi.org/10.1038/gene.2009.74 PMID: 19798077

38. Rebmann V, LeMaoult J, Rouas-Freiss N, Carosella ED, Grosse-Wilde H. Report of the Wet Workshop for quantification of soluble HLA-G in Essen, 2004. Hum Immunol. 2005; 66: 853–863. https://doi.org/10.1016/j.humimm.2005.05.005 PMID: 16216668

39. Menier C, Saez B, Horejsi V, Martinozzi S, Krawic-Radanne I, Bruel S, et al. Characterization of monoclonal antibodies recognizing HLA-G or HLA-E: New tools to analyze the expression of nonclassical HLA class I molecules. Hum Immunol. 2003; 64: 315–326. https://doi.org/10.1016/S0198-8859(02)00821-2 PMID: 12590976

40. Guo SW, Thompson EA. Performing the exact test of Hardy-Weinberg proportion for multiple alleles. Biometrics. 1992; 48: 361–72.

41. Excoffier L, Laval G, Schneider S. Arlequin (version 3.0): an integrated software package for population genetics data analysis. Evol Bioinform Online. 2007; 1: 47–50. Available: http://www.ncbi.nlm.nih.gov/pubmed/19329582 PMID: 19329582

42. Barrett JC, Fry B, Maller J, Daly MJ. Haploview: analysis and visualization of LD and haplotype maps. Bioinformatics. 2005; 21: 263–5. https://doi.org/10.1093/bioinformatics/bth457 PMID: 15297300

43. Meuleman T, Drabbel EP, Mattar SB, Bompeixe EP, Silva J dos S, Pirri A, et al. Association of HLA-G alleles and 3’UTR 14 bp haplotypes with recurrent miscarriage in Brazilian couples. Hum Immunol. 2011; 72: 479–85. https://doi.org/10.1016/j.humimm.2011.02.011 PMID: 21954454

44. Cilião Alves DC, De Oliveira Crispim JC, Castelli EC, Mendes-Junior CT, Deghaide NHS, Barros Silva GE, et al. Human leukocyte antigen-G 3’ untranslated region polymorphisms are associated with better kidney allograft acceptance. Hum Immunol. 2012; 73: 52–59. https://doi.org/10.1016/j.humimm.2011.10.007 PMID: 22044721
47. Martelli-Palomino G, Pancotto JA, Muniz YC, Mendes-Junior CT, Castelli EC, Massaro JD, et al. Polymorphic Sites at the 3’ Untranslated Region of the HLA-G Gene Are Associated with Differential hla-g Soluble Levels in the Brazilian and French Population. PLoS One. 2013; 8: 1–10. https://doi.org/10.1371/journal.pone.0071742 PMID: 24204558

48. Amodio G, Canti V, Maggio L, Rosa S, Castiglioni MT, Rovere-Querini P, et al. Association of genetic variants in the 3’UTR of HLA-G with Recurrent Pregnancy Loss. Hum Immunol. The Authors; 2016; 77: 886–891. https://doi.org/10.1016/j.humimm.2016.06.020 PMID: 27370685

49. Rajasekaran A, Shivakumar V, Kalmady S V., Narayanaswamy JC, Subbana M, Venugopal D, et al. The impact of HLA-G 3’ UTR variants and sHLA-G on risk and clinical correlates of schizophrenia. Hum Immunol. American Society for Histocompatibility and Immunogenetics; 2016; 77: 1166–1171. https://doi.org/10.1016/j.humimm.2016.08.013 PMID: 27567986

50. Veit TD, Chies JAB. Tolerance versus immune response—microRNAs as important elements in the regulation of the HLA-G gene expression. Transpl Immunol. Elsevier B.V.; 2009; 20: 229–31. https://doi.org/10.1016/j.trim.2008.11.001 PMID: 19038339

51. Zambra FMB, Biolchi V, de Cerqueira CCS, Brum IS, Castelli EC, Chies JAB. Immunogenetics of prostate cancer and benign hyperplasia—the potential use of an HLA-G variant as a tag SNP for prostate cancer risk. HLA. 2016; 87: 79–88. https://doi.org/10.1111/tan.12741 PMID: 26889902

52. Li J-J, Pan K, Gu M-F, Chen M-S, Zhao J-J, Wang H, et al. Prognostic value of soluble MICA levels in the serum of patients with advanced hepatocellular carcinoma. Chin J Cancer. 2013; 32: 141–8. https://doi.org/10.5732/cjc.012.10025 PMID: 22704489

53. Passos-Bueno MR, Bertola D, Horovitz DDG, de Faria Ferraz VE, Brito LA. Genetics and genomics in Brazil: a promising future. Mol Genet Genomic Med. 2014; 2: 280–91. https://doi.org/10.1002/mgg3.95 PMID: 25077170

54. Castelli EC, Ramalho J, Porto IOP, Lima THA, Felício LP, Sabbagh A, et al. Insights into HLA-G Genetics Provided by Worldwide Haplotype Diversity. Front Immunol. 2014; 5: 476. https://doi.org/10.3389/fimmu.2014.00476 PMID: 25339953