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Development and Validation of a Multiplex Non-HLA Antibody Assay for the Screening of Kidney Transplant Recipients

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The best treatment for patients with end-stage renal disease is kidney transplantation. Although graft survival rates have improved in the last decades, patients still may lose their grafts partly due to the detrimental effects of donor-specific antibodies (DSA) against human leukocyte antigens (HLA) and to a lesser extent also by antibodies directed against non-HLA antigens expressed on the donor endothelium. Assays to detect anti-HLA antibodies are already in use for many years and have been proven useful for transplant risk stratification. Currently, there is a need for assays to additionally detect multiple non-HLA antibodies simultaneously in order to study their clinical relevance in solid organ transplantation. This study describes the development, technical details and validation of a high-throughput multiplex assay for the detection of antibodies against 14 non-HLA antigens expressed on the donor endothelium.
INTRODUCTION

Kidney transplantation is the preferred treatment for most patients with end-stage renal disease. Pretransplant donor-specific anti-human leukocyte antigen (HLA) antibodies (DSA) have been shown to be a major risk factor in kidney graft loss (1–3). In recent years non-HLA antibodies are increasingly being recognized as a cause for antibody-mediated graft dysfunction leading to graft loss, due to reports of antibody-mediated rejection or C4d deposition in the absence of circulating DSA (4–6). A landmark study (7) in 2005 indicated a role for non-HLA immunity by showing a relation between graft loss in HLA-identical sibling kidney transplants and percentage pan-reactive antibodies (PRA) against HLA antigens. As graft loss could not be attributed to DSA, PRA served as an indicator of increased immunity against the graft, including non-HLA antigens (7). In the past years, several reports have been published describing AMR in the absence of circulating DSA (8–17). These reports have raised interest in the identification of immunogenic non-HLA molecules and a number of target antigens were identified. At present, there is a need for the development of coherent screening assays for the detection of multiple non-HLA antibodies simultaneously in order to study their clinical relevance in solid organ transplantation (18).

Based on a literature search we have selected fourteen non-HLA targets deemed relevant for kidney transplantation (see Table 1 for a detailed description). Antibodies against angiotensin-II type I receptor (AT1R) and endothelin type A receptor (ETAR) expressed on the endothelium have already been associated with kidney graft loss using a commercially available ELISA (23, 25). The other twelve antibodies, found in patients with kidney disease or after kidney transplant, have not yet been associated with graft loss. In this study, we describe the development, technical details, and validation of a multiplex assay on a Luminex platform that can be used to detect non-HLA antibodies in kidney transplant recipients. In the PROCARE study (2) the clinical relevance of these non-HLA antibodies will be measured in 4770 sera collected prior to kidney transplantation, and the results will be reported separately. Using this high-throughput assay, the non-HLA antibody status can be determined in large cohorts and potentially be used in pre- and post-transplant risk stratification.

MATERIALS AND METHODS

In-house Protein Production

For the production of the HaloTag-proteins, we generated a universal vector containing the HaloTag® open reading frame [from the HaloTag® CMV/dl Flexi® Vector (Promega, Madison, Wisconsin)] cloned in a pcDNA3.1 derived plasmid (Thermo Fisher Scientific, Waltham, Massachusetts) with multiple cloning site altered and added woodchuck posttranscriptional regulatory element (WPRE) before the poly A tail in order to stabilize the mRNA and enhance protein production (30). For PLA2R without a HaloTag we used the pcDNA3.1-WPRE vector only. We ordered the different inserts each starting with a BglII restriction site, then a Kozak sequence and a standard signal peptide, the codon optimized DNA sequence coding for the protein of interest, followed by 6-Histidines for purification purposes and ending with an XhoI restriction site from the Invitrogen GeneArt Gene synthesis (Thermo Fisher Scientific (Figure 3)). The BglII and XhoI restriction sites we transferred the synthesized inserts from the suppliers vector to our HaloTag expression vector (Figure 3). The amino acid sequence used for the in-house produced proteins are provided in Supplementary Table S1. For all constructs the complete open reading frames were sequence verified by Sanger sequencing. Subsequently, 1 x 10⁸ HEK293-F cells were co-transfected with a 100 µg DNA mix (containing per non-HLA an optimized ratio of the pAdVantage™ vector (Promega) and the vector containing the sequence of the protein of interest) and 130 µg 293-Fectin in FreeStyle™ 293 Expression Medium (both Thermo Fisher Scientific). As the universal vector contained a signal peptide, the proteins were secreted into the culture supematant, which was harvested at day 4. An AKTA start protein purification system (GE Lifesciences, Chicago, Illinois) fitted with a HisTrap™ HP column (GE Lifesciences) was used to purify proteins from the cell supernatants. The fractions containing the protein of interest were pooled. To remove imidazole and other contaminants smaller than 10 kDa, the pooled fractions were dialyzed to PBS in a Slide-A-Lyzer™ Dialysis Cassette 10 kDa (Thermo Fisher Scientific). The proteins were detected by Western blot using various antibodies. See Supplementary Table S2 for a list of all antibodies used in the Western blot and their dilutions.

**Abbreviations:** AMR, antibody-mediated rejection; APMAP, Adipocyte plasma membrane-associated protein; ARHGDIB, Rho GDP-dissociation inhibitor 2; ARHGGEF6, Rho guanine nucleotide exchange factor 6; AT1R, Angiotensin receptor (ETAR) expressed on the endothelium have already been associated with kidney graft loss using a commercially available ELISA (23, 25). The other twelve antibodies, found in patients with kidney disease or after kidney transplant, have not yet been associated with graft loss. In this study, we describe the development, technical details, and validation of a multiplex assay on a Luminex platform that can be used to detect non-HLA antibodies in kidney transplant recipients. In the PROCARE study (2) the clinical relevance of these non-HLA antibodies will be measured in 4770 sera collected prior to kidney transplantation, and the results will be reported separately. Using this high-throughput assay, the non-HLA antibody status can be determined in large cohorts and potentially be used in pre- and post-transplant risk stratification.

**MATERIALS AND METHODS**

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TABLE 1 | Fourteen non-HLA proteins selected based on a literature search in patients with kidney disease or after kidney transplant.

<table>
<thead>
<tr>
<th>Protein</th>
<th>Other name</th>
<th>UniProtKB</th>
<th>Localization</th>
<th>Expression</th>
<th>References</th>
<th>Patient population</th>
</tr>
</thead>
<tbody>
<tr>
<td>Agrin</td>
<td></td>
<td>O00468</td>
<td>Secreted, extracellular matrix</td>
<td>Wide expression including glomerular basement membrane</td>
<td>(19)</td>
<td>Patients with transplant glomerulopathy</td>
</tr>
<tr>
<td>APMAP (Adipocyte plasma membrane-associated protein)</td>
<td>C20orf3</td>
<td>Q9HDC9</td>
<td>Single pass membrane protein</td>
<td>Ubiquitously expressed</td>
<td>(20)</td>
<td>Patients awaiting kidney retransplant</td>
</tr>
<tr>
<td>ARHGDIIB (Rho GDP-dissociation inhibitor 2)</td>
<td>RhoGDII</td>
<td>P52566</td>
<td>Intracellular: cytoplasm</td>
<td>Wide expression including renal pelvis and glomerulus</td>
<td>(21)</td>
<td>Chronic hemodialysis patients</td>
</tr>
<tr>
<td>ARHGEF6 (Rho guanine nucleotide exchange factor 6)</td>
<td>Q15052</td>
<td></td>
<td>Intracellular: cytoplasm</td>
<td>Ubiquitously expressed</td>
<td>(22)</td>
<td>Pediatric kidney transplant recipients</td>
</tr>
<tr>
<td>AT1R (Angiotensin type I receptor)</td>
<td>P30556</td>
<td></td>
<td>Transmembrane protein</td>
<td>Adipose and soft tissues</td>
<td>(5, 23)</td>
<td>Kidney transplant recipients</td>
</tr>
<tr>
<td>Endorepellin (C-terminal fragment of perlecian)</td>
<td>P98160</td>
<td></td>
<td>Secreted, extracellular matrix</td>
<td>Ubiquitously expressed</td>
<td>(24)</td>
<td>Kidney transplant recipients with acute vascular rejection</td>
</tr>
<tr>
<td>ETAR (Endothelin A receptor)</td>
<td>EDNRA</td>
<td>P25101</td>
<td>Transmembrane protein</td>
<td>Ubiquitously expressed</td>
<td>(25)</td>
<td>Kidney transplant recipients</td>
</tr>
<tr>
<td>Lamin B1</td>
<td></td>
<td>P20700</td>
<td>Intracellular: Nuclear membrane</td>
<td>Ubiquitously expressed</td>
<td>(21)</td>
<td>Chronic hemodialysis patients</td>
</tr>
<tr>
<td>LPLUNC1 (BPI fold-containing family B member 1)</td>
<td>BPIFB1/C20orf114</td>
<td>Q8TDL5</td>
<td>Secreted, extracellular matrix</td>
<td>Respiratory epithelia, stomach, small intestine and salivary gland</td>
<td>(20)</td>
<td>Patients awaiting kidney retransplant</td>
</tr>
<tr>
<td>PECR (Peroxisomal trans-2-enoyl-CoA reductase)</td>
<td>Q9BY49</td>
<td></td>
<td>Intracellular: cytoplasm, peroxisome</td>
<td>Ubiquitously expressed</td>
<td>(26)</td>
<td>Patients with transplant glomerulopathy</td>
</tr>
<tr>
<td>PLA2R (Phospholipase A2 receptor)</td>
<td>Q13018</td>
<td></td>
<td>Membrane, secreted</td>
<td>Renal glomeruli</td>
<td>(27)</td>
<td>Patients with membranous nephropathy</td>
</tr>
<tr>
<td>PRKCC2 (Protein kinase C zeta type)</td>
<td>Q05513</td>
<td></td>
<td>Intracellular: cytoplasm</td>
<td>Ubiquitously expressed</td>
<td>(28)</td>
<td>Pediatric kidney transplant recipients</td>
</tr>
<tr>
<td>Tubb4B (Tubulin beta-4B)</td>
<td>P68371</td>
<td></td>
<td>Intracellular: cytoplasm</td>
<td>Ubiquitously expressed</td>
<td>(21)</td>
<td>Chronic hemodialysis patients</td>
</tr>
<tr>
<td>Vimentin</td>
<td></td>
<td>P08670</td>
<td>Intracellular: cytoplasm</td>
<td>Ubiquitously expressed</td>
<td>(21, 29)</td>
<td>Chronic hemodialysis patients and renal transplant recipients with IFTA (interstitial fibrosis and tubular atrophy)</td>
</tr>
</tbody>
</table>

**Treatment With PNGase F**

All purchased and in-house produced proteins with an N-Acetylg glucosamine (N-GlcNAC) according to UniProt (Table 2) were treated with PNGase F (New England Biolabs) according to manufacturer’s protocol. The PNGase F treated and untreated proteins were detected by Western blot using various antibodies (Supplementary Table S2).

**Direct Coupling of Proteins to Microspheres**

First, using a Vivaspin® 500 (Sartorius, Göttingen, Germany) with pore size 10 kDa, 10 of the 15 purchased proteins which were dissolved in glycerol or Tris-HCl were transferred to PBS. Covalent coupling of the sixteen (proteins Table 2 upper part) different carboxylated MagPlex® Microspheres (Luminex Corp, Austin, Texas) each with different emitting fluorescence pattern was performed by following the procedures recommended by Luminex (31). In short, 6.25 × 10⁶ microspheres (500 µl of stock) suspension was resuspended in microtiter tubes containing 0.1 M sodium phosphate buffer (pH 6.1) to a final volume of 200 µl. Fresh solutions of N-hydroxy-sulfosuccinimide (Sulfo-NHS) and 1-ethyl-3-(3-dimethylaminopropyl)-carbodiimide hydrochloride (Pierce), both at 50 mg/ml, were prepared in phosphate buffer, and 25 µl of each solution was sequentially added to stabilize the reaction and activate the microspheres. This suspension was incubated for 5 min at room temperature and then resuspended in 625 µl protein solution (1.5 µg protein per 1 × 10⁶ microspheres). The mixture was incubated overnight in the dark with continuous shaking. The microspheres were then incubated with 625 µl PBS-0.05% Tween-20 for 4 h. After aspiration, the microspheres were blocked with PBS-0.1% BSA-0.1% sodium azide to a final concentration of 2,500 beads/µl. The microspheres were counted on Beckman Coulter Counter and stored in the dark at 4°C. Coupling efficiency was tested using anti-protein antibodies (Figure 1 and Supplementary Table S2).

**Coupling of HaloTag Proteins to Microspheres**

First, microspheres were coupled with HaloTag Amine (O4) Ligand (Promega). 6.25 × 10⁶ microspheres were suspended in...
TABLE 2 | Overview of all purchased and in-house produced proteins.

<table>
<thead>
<tr>
<th>Coupling</th>
<th>Protein</th>
<th>Source</th>
<th>Company</th>
<th>Size including Tags (kDa)</th>
<th>N-GlcNAc</th>
<th>Glycosylation confirmed by PNGase F treatment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Direct</td>
<td>IgG</td>
<td>Human serum</td>
<td>Sigma Aldrich</td>
<td>150</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>Agrin</td>
<td>CHO cell line</td>
<td>R&amp;D systems</td>
<td>100</td>
<td>No</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>APMAP</td>
<td>Wheat Germ (in vitro)</td>
<td>Abnova</td>
<td>73</td>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td></td>
<td>ARHGDIB</td>
<td>Wheat Germ (in vitro)</td>
<td>Abnova</td>
<td>48</td>
<td>No</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>ARHGEF6</td>
<td>Wheat Germ (in vitro)</td>
<td>Abnova</td>
<td>114</td>
<td>No</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>AT1R</td>
<td>Wheat Germ (in vitro) with proprietary liposome technology</td>
<td>Abnova</td>
<td>41</td>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td></td>
<td>Endorepellin</td>
<td>Mouse myeloma cell line</td>
<td>R&amp;D systems</td>
<td>90</td>
<td>Yes</td>
<td>Yes</td>
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<tr>
<td></td>
<td>ETAR</td>
<td>Wheat Germ (in vitro)</td>
<td>Abnova</td>
<td>75</td>
<td>Yes</td>
<td>No</td>
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<td></td>
<td>LMNB1</td>
<td>Wheat Germ (in vitro)</td>
<td>Abnova</td>
<td>93</td>
<td>No</td>
<td>–</td>
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<tr>
<td></td>
<td>LPLUNC1</td>
<td>HEK293 Cells</td>
<td>Sino Biological Inc.</td>
<td>53</td>
<td>Yes</td>
<td>Yes</td>
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<tr>
<td></td>
<td>PECR</td>
<td>E coli</td>
<td>Abcam</td>
<td>35</td>
<td>No</td>
<td>–</td>
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<tr>
<td></td>
<td>PLA2R</td>
<td>HEK293 Cells</td>
<td>in-house production</td>
<td>160</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td></td>
<td>PPKCZ</td>
<td>Wheat Germ (in vitro)</td>
<td>Abnova</td>
<td>94</td>
<td>No</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>Transferrin</td>
<td>Serum of non-immunized animals</td>
<td>Jackson ImmunoResearch</td>
<td>80</td>
<td>No</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>TUBB4B</td>
<td>Wheat Germ (in vitro)</td>
<td>Abnova</td>
<td>75</td>
<td>No</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>Vimentin</td>
<td>E coli</td>
<td>R&amp;D systems</td>
<td>55</td>
<td>No</td>
<td>–</td>
</tr>
<tr>
<td>HaloTag</td>
<td>Agrin_HaloTag</td>
<td>HEK293 cells</td>
<td>in-house production with HaloTag</td>
<td>138</td>
<td>No</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>PECR_HaloTag</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>PLA2R_HaloTag</td>
<td>HEK293 cells</td>
<td>in-house production with HaloTag</td>
<td>113</td>
<td>Yes</td>
<td>Yes</td>
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<tr>
<td></td>
<td>PPKCZ_HaloTag</td>
<td>HEK293 cells</td>
<td>in-house production with HaloTag</td>
<td>52</td>
<td>Yes</td>
<td>Yes</td>
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<tr>
<td></td>
<td>Transferrin_HaloTag</td>
<td>HEK293 cells</td>
<td>in-house production with HaloTag</td>
<td>105</td>
<td>No</td>
<td>–</td>
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<tr>
<td></td>
<td>TUBB4B_HaloTag</td>
<td>HEK293 cells</td>
<td>in-house production with HaloTag</td>
<td>91</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td></td>
<td>Vimentin_HaloTag</td>
<td>HEK293 cells</td>
<td>in-house production with HaloTag</td>
<td>96</td>
<td>No</td>
<td>–</td>
</tr>
</tbody>
</table>

NA, not available, N-GlcNAC, N-Acetylglucosamine; see Table 1 for the full protein names and additional information.

A solution of 100 mmol/l MES (pH 6.0) containing 5 mg/ml 1-ethyl-3-(3-dimethylaminopropyl)-carbodiimide hydrochloride (Pierce) and 0.2 mg HaloTag Amine Ligand. After 2 h incubation in the dark, the microspheres were washed and resuspended in 100 mmol/l MES (pH 4.5) and stored at 4°C for at least 16 h before using for protein coupling. Next, after aspiration the microspheres coupled with HaloTag amine ligand were washed with PBS-0.05% Tween-20. This suspension was incubated
with 1 ml HaloTag-protein fraction (protein concentration ranging from 25 to 100 µg/ml) for 1 h in the dark with continuous shaking. After washing with PBS-0.05% Tween-20, the microspheres were aspirated and blocked with PBS-0.1% BSA-0.1% sodium azide to a final concentration of 2,500 beads/µl. The microspheres were counted on Beckman Coulter Counter and stored in the dark at 4°C. Coupling efficiency was tested using anti-protein antibodies (Figure 1 and Supplementary Table S2).

**Multiplex Luminex Assay**

A mix of 31 different microspheres was made consisting of the 15 directly-coupled proteins, 13 in-house produced HaloTag-coupled proteins, the IgG-coupled microsphere as a positive control, the HaloTag amine-coupled microsphere, and an empty microsphere. 1,500 microspheres per well were used of each specificity diluted in PBS–0.1% BSA (wash buffer). First, 48 µl diluted microsphere mix was aliquoted into a 96-well Bio-Plex Pro Flat bottom plate (Bio-Rad, Hercules, California) and then 2 µl serum was added (serum dilution 1:25). For assay validation, instead of serum anti-protein antibodies were added. After overnight incubation in the dark at room temperature with continuous shaking, samples were washed using a Bio-Plex Pro Wash station (Bio-Rad). Next, 50 µl of 1:50 R-phycocerythrin-conjugated goat-anti human antibody (Jackson ImmunoResearch, Cambridgeshire, UK) diluted in wash buffer was added per well. After 30 min incubation at room temperature with continuous shaking, 50 µl wash buffer was added and samples were measured on a Luminex 200 flow analyzer (Luminex Corp). See Supplementary Table S3 for a list of all antibodies used in the Luminex assay and the used dilution. Readout is provided as median fluorescence intensity (MFI).

**Commercial Autoantibody Luminex Assay**

For comparison, LABScreen Autoantibody assay group 1 (Lot 1: 32 targets) and group 2 (Lot 1: 1 target) from One Lambda (Canoga Park, CA) was used to determine the non-HLA antibody profile in patient sera according to manufacturer’s instructions. One Lambda donated reagents but was not involved in either the conduct of the study or the preparation of the manuscript. In brief, 10 µl of Autoantibody microsphere mix was incubated with 40 µl of serum in a 96 well V-bottom plate for 30 min in the dark at room temperature with continuous shaking. After washing, PE-conjugated anti-human IgG was added and samples were again incubated for 30 min in the dark at room temperature with continuous shaking. Finally, samples were washed again, PBS was added and samples were measured on a Luminex 200 flow analyzer. Readout is provided as median fluorescence intensity (MFI) and adjusted for background by subtracting the MFI of the negative control microsphere from the MFI of the specific microsphere. This adjusted value is referred to as Baseline.

**Serum Samples**

The use of sera and experimental protocols was approved by the Research Ethics Committee for Biobanks and the Medical Ethics Committee of the University Medical Center Utrecht and was performed in accordance with the FEDERA Code of Conduct. Sera were passed through a 96-well 1.2 µm MultiScreen filter plate (Millipore, Billerica, MA) to clear debris before used in the Luminex assay. No other serum pretreatment was performed.

Eighteen patient sera from a national inter-laboratory comparison of the qualitative anti-PLA2R indirect immunofluorescence test (IIFT) were used for validation of our PLA2R Luminex assay. The national consensus conclusions were used to compare to MFI levels we measured. Sera of 87 controls (healthy persons fit to work or deceased donors approved for lung donation) were used to assess non-HLA antibodies in persons without end-stage renal disease. From a national consortium study (2), we selected eight sera from patients with end-stage renal disease with high MFI levels against a variety of non-HLA specificities to test their specificity in inhibition experiments using each produced non-HLA antigen added in fluid phase. From the same consortium study, fourteen sera were selected containing antibodies against 8 non-HLA proteins both present in our assay and the commercially available assay in order to compare results of our assay to the commercially available Autoantibody assay.

**Statistical Analysis**

Statistical analyses were performed with SAS 9.4 (SAS Institute, Cary, NC, United States). Continuous data were analyzed with the Mann-Whitney U-test. The correlation between our assay and the commercially available Autoantibody assay was evaluated.
using Pearson’s correlation coefficient. \( P < 0.05 \) was considered as statistically significant.

RESULTS

Protein Production

We chose two different strategies to covalently couple the protein antigens to Luminex microspheres: (1) direct coupling via the amino group to the carboxyl group on the microsphere (Figure 2A) and (2) indirect (HaloTag) coupling via the HaloTag-recombinant protein to the HaloTag amine ligand bound to the microsphere (Figure 2B). Both coupling strategies result in covalent and stable binding of the proteins to the microspheres. For direct coupling we purchased all non-HLA proteins, except PLA2R (produced in-house), and for the coupling via the HaloTag we produced all non-HLA proteins in-house. Table 2 provides an overview of all proteins with their production method or source, size including Tags, whether N-Acetylglucosamine (N-GlcNAc) was present according to UniProt, and whether glycosylation could be confirmed by treatment with PNGase F. We were not able to produce AT1R and PECR in-house because contaminants such as the HaloTag protein (34 kDa) which does not contain histidine residues. As we produced relatively low amounts of proteins, we were not able to check for purity by Coomassie blue protein staining. The HaloTag coupling to the microspheres is another purification step, as HaloTag is usually used for protein purification (as advertised by Promega). Therefore, protein size and glycosylation status were studied by Western blot using appropriate antibodies (Figure 4) and compared to the predicted size (Table 2). The estimated size was comparable to the predicted size for all 28 proteins. Glycosylation could not be demonstrated for three of six purchased and directly-coupled proteins with expected N-GlcNAc (APMAP, AT1R, and ETAR), since the size of proteins was unchanged after treatment with PNGase F (Figure 4B). The Western blot of the purchased AT1R, purchased ETAR (both Figure 4B) and in-house produced ETAR_HaloTag (Figure 4D) showed multiple bands which might be caused by the presence of SDS-PAGE and DTT stable aggregates. For the three other proteins with N-GlcNAc used for direct coupling (Endorepellin, LPLUNC1 and PLA2R) glycosylation could be confirmed by PNGase F treatment. These proteins were produced in a mouse myeloma cell line or in HEK293 cells. Also for all five HaloTag proteins with N-GlcNAc we could confirm the glycosylation (Figure 4D).

Validation of the Multiplex Non-HLA Antibody Assay

After analysis of all proteins using Western blot as described above, the proteins were coupled to MagPlex microspheres. In order to optimize detection of non-HLA antibodies, we tested different incubation times, incubation temperatures and serum dilutions. Overnight incubation at room temperature using a serum dilution of 1:25 resulted in the highest specific MFI levels compared to minimal background signal using anti-PLA2R IIFT positive sera (data not shown). Similar results were observed for patient sera containing anti-Vimentin antibodies.

To assess whether the PLA2R microspheres were correctly coupled, we used patient sera with anti-PLA2R antibodies determined with the qualitative IIF test (Figure 5). The average MFI ± SD for the PLA2R directly-coupled microspheres was 548 ± 390 for the PLA2R IIFT negative sera and 4,548 ± 3,256 for the PLA2R IIFT positive sera (\( P = 0.0008 \)). For the PLA2R-HaloTag microspheres this was 1,593 ± 1,408 for the PLA2R IIFT negative sera and 7,058 ± 2,152 for the PLA2R IIFT positive sera (\( P = 0.0003 \); Figure 5A). Even though, there is a high variation between the patient sera, the PLA2R IIFT positive and negative sera could easily be distinguished based on the measured MFI of our assay. According to the qualitative IIF test, PLA2R positive sera could be subdivided into weak positive and positive. Using this stratification, we plotted the MFI values measured in our assay with the two different PLA2R microspheres (direct and HaloTag coupling). The MFI values of the positive sera were higher compared to the weak positive sera (\( P = 0.106 \) for PLA2R weak positive compared to PLA2R positive, and \( P = 0.048 \) for PLA2R_HaloTag weak positive compared to PLA2R_HaloTag positive). In our assay we defined that the MFI of the IgG microspheres should be at least 10,000 to assure correct secondary antibody binding, in line with the commercially available Luminex assay to determine HLA antibodies. The four negative control microspheres had overall low MFI values (empty microsphere average MFI ± SD: 360 ± 247; Transferrin: 692 ± 961; HaloTag amine ligand 844 ± 673; Transferrin_HaloTag 479 ± 604). 10 PLA2R IIF positive sera were used to determine assay variation on several days in our laboratory using the same lot. The mean coefficient of variation was 15.1% for the MFI of PLA2R and PLA2R_HaloTag microspheres (data not shown).

Human sera with antibodies against the other non-HLA targets are not commercially available nor present in external proficiency testing, as is the case for anti-PLA2R antibodies. To test whether coupling of the microspheres was successful, we used commercially available anti-non-HLA antibodies (Supplementary Table S3). As the principle of ELISA is comparable to our Luminex assay, we selected animal source anti-non-HLA antibodies that were validated for ELISA. We obtained validated antibodies from animal sources for 9 of the 15 non-HLA targets (Supplementary Table S3). So 6 of these specific antibodies from animal sources were not validated.
We refer to direct coupling when the amino group of the protein is directly coupled to the carboxylated microsphere. For the HaloTag coupling a HaloTag amine ligand serves as a connection between the carboxyl group and the HaloTag of the protein. Due to this indirect coupling, the protein of interest is freely accessible compared to the direct situation. Part of the figure is adapted from Promega.

In the general pcDNA3-WPRE vector starting with a CMV promoter and including the HaloTag sequence and the WPRE sequence, which enhances expression, we ligated an insert consisting of a BglII restriction site, then a signal peptide, the DNA sequence of the protein of interest, followed by 6-Histidines and ending with an XhoI restriction site.

for the use in an ELISA assay and could therefore result in lower MFI values. Using these antibodies we compared the MFI values for the directly- and HaloTag-coupled microspheres (Figure 1). For most of the microspheres we observed relatively high MFI values, confirming the coupling and recognition of the non-HLA proteins. Potential causes of relatively low MFI values for some microspheres could be due to the fact that the antibody is not suitable for ELISA/Luminex, that the epitopes these antibodies recognize are not accessible on the direct- or HaloTag-coupled microspheres, or that the coupling efficiency is low. To check whether the signals were specific, we preincubated 8 patient sera, with high MFI levels against a variety of non-HLA specificities, with the specific protein or with transferrin (concentrations ranging from 0 to 60 ng/μl). Inhibition in antibody binding (decrease in MFI) was observed if sera were pre-incubated with the specific protein, whereas no inhibition was found after preincubation with transferrin (data not shown).

Multiplex Assay With Healthy Control Sera
Next, we screened sera of 87 healthy controls with the multiplex non-HLA antibody assay. The control microspheres performed as expected: IgG-MFI was above 10,000 and the 4 negative control microspheres showed in general low MFI values (mean MFI < 500; Figure 6A). For the direct-coupled microspheres, 8 targets had a mean MFI below 1,000 (Agrin, APMA, ARHGDIB, ARHGEF6, ETAR, LMNB1, LPLUNC1, and PLA2R), 4 targets had a mean MFI between 1,000 and 1,500 (PECR, PRKCZ, TUBB4B, and Vimentin; Figure 6B). AT1R had a mean MFI ± SD of 3,489 ± 2,322. The results acquired with the direct-coupled Endorepellin are not reliable and were excluded from further analyses. For the direct-coupled Endorepellin, the majority of
microspheres had a relatively high MFI with a mean MFI ± SD of 11,769 ± 7,492. This Endorepellin was produced in a mouse cell line, and the high MFI-values appeared due to cross-reactivity of our secondary goat anti-human antibody to mouse (data not shown). For the HaloTag coupled microspheres, the mean MFI values for the healthy controls were much lower compared to the direct-coupled microspheres, except for ARHGDIIB and PLA2R (Figure 6C). Most mean MFI values for the HaloTag coupled microspheres were below 1,000, only the mean MFI of ARHGDIIB_HaloTag was above 1,000 (mean MFI ± SD: 1,255 ± 1,532).

Comparison With Commercially Available Autoantibody Assay
Recently, a commercial Autoantibody multiplex assay became available which included 8 of the 14 non-HLA targets we selected (Agrin, ARHGDIIB, Endorepellin (C-terminal part LG3), LMNB1, PECR, PLA2R, PRKCG, and Vimentin). 14 patient sera were selected based on a variety of MFI levels against the 8 targets in our assay. We analyzed these sera with our direct- and HaloTag-coupled non-HLA antibody assay and compared the baseline calculations to the commercially available autoantibody assay. The correlation ($r^2$) of the baseline calculations between our assay and the commercial assay varied between 0.0903 and 0.9628 (Table 3). Although it is reassuring to see that for some non-HLA targets there is a high correlation between our assay and the commercial assay, it is difficult to properly compare the assays because essential details from the commercial assay are unknown such as protein size, glycosylation, source, purity and coupling procedure to the microspheres.

DISCUSSION
The major objective of this study was to develop a high-throughput non-HLA antibody assay that can be used for the screening of kidney, heart and lung transplant recipients. At
Detection of anti-PLA2R antibodies in patient sera

(A) 18 patient sera (6 negative and 12 positive according to the qualitative anti-PLA2R indirect immunofluorescence test (IIFT)) were incubated with a microsphere mix consisting of empty microspheres and microspheres coupled with HaloTag amine ligand, Transferrin (direct-coupling), Transferrin_HaloTag (together four negative controls), coupled with IgG (positive control), and also two specific microspheres coupled with PLA2R (direct) or PLA2R_HaloTag. The MFI values of each independent patient serum are depicted including the mean and standard deviation.

(B) The 12 IIFT positive patient sera can be divided into weak positive (n = 5) and positive (n = 7) according to the immunofluorescence staining. Using this stratification, we plotted the MFI values measured in our assay with the two different PLA2R microspheres (direct- and HaloTag-coupling).

In this study we coupled non-HLA antigens directly to the microspheres as well as indirectly via a HaloTag linker. At present it is unknown which epitopes are predominantly recognized on each of the non-HLA antigens included, but as the coupling process may influence the epitopes recognized, our rationale to include both coupling procedures was to empirically find out which coupling procedure is best suited to detect clinically relevant non-HLA antibodies. To have all relevant epitopes exposed upon coupling of proteins to the microspheres, we anticipate that the HaloTag is superior to the direct-coupling, although the trade-off is that less molecules can be bound to individual microspheres resulting in less antibody binding leading to lower MFI levels.

In addition, whilst time-consuming, the HaloTag system is more flexible compared to direct coupling, allowing research groups to produce and develop their own assay with any protein of interest. Another major disadvantage of the direct-coupled assay is the dependency on commercially available proteins, (either full length proteins or peptides), and the cell lines in which they are produced can have an effect on posttranslational modifications such as glycosylation. For three purchased proteins we could not demonstrate glycosylation. This lack of glycosylation is probably explained by the \textit{in vitro} production in wheat germ (Abnova). As we don’t know the production details of the proteins used in the commercially available Autoantibody assay, the comparison between the assays is probably not completely valid.

To determine whether a serum sample is positive or negative for a non-HLA antibody, a clinically meaningful cut-off has to be defined. The commercial autoantibody multiplex assay
used for comparison in this study makes use of serum samples from non-transplanted individuals. Using the results of 100 individuals, the reference background value was calculated from the median of the MFI plus 2 times the SD defining ~3–5% of healthy as being positive for non-HLA antibodies. However, this is by definition not a clinically relevant cut-off as every healthy individual potentially has antibodies against autoimmune targets, and the definition used is unrelated to clinical key-parameters such as graft loss or antibody-mediated rejection. Patients with end-stage renal disease have massive cellular damage in their kidneys and occurrence of cell death or apoptosis is known to result in release of intracellular proteins which are normally not accessible by antibodies. This release likely results in a boost of autoantibody formation generated to clear these antigens from the circulation, thus including a healthy control cohort as reference for non-HLA antibody levels for patients with end-stage renal disease may not be optimal. In our opinion, the best way to determine the clinically relevant cut-off for kidney transplant recipient is to use a clinical end point such as rejection or graft failure (32). The high-throughput multiplex non-HLA assay described here can be used to assess defined patient groups to determine whether these 14 non-HLA antibodies are associated with graft rejection or graft loss. Recently, we determined the effect of pretransplant single antigen bead-defined donor-specific HLA antibodies in 4770 kidney transplant recipients (2). This well defined cohort will be
used to determine the clinical relevance of these pretransplant non-HLA antibodies and results will be published in a separate study.

AUTHOR CONTRIBUTIONS


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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fimmu.2018.03002/full#supplementary-material

REFERENCES


Conflict of Interest Statement: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.