We are IntechOpen, the world’s leading publisher of Open Access books
Built by scientists, for scientists

4,200 Open access books available
116,000 International authors and editors
125M Downloads

154 Countries delivered to
TOP 1% Our authors are among the most cited scientists
12.2% Contributors from top 500 universities

WEB OF SCIENCE™
Selection of our books indexed in the Book Citation Index in Web of Science™ Core Collection (BKCI)

Interested in publishing with us?
Contact book.department@intechopen.com

Numbers displayed above are based on latest data collected.
For more information visit www.intechopen.com
Donor-Specific Anti-HLA Antibodies in Organ Transplantation: Transition from Serum DSA to Intra-Graft DSA

Tsukasa Nakamura, Hidetaka Ushigome, Takayuki Shirouzu and Norio Yoshimura

Additional information is available at the end of the chapter

http://dx.doi.org/10.5772/intechopen.79846

Abstract

In the field of organ transplantation, donor-specific anti-HLA antibodies (DSA) have gained more popularity, as antibody-mediated rejection (AMR) has been recognized as an important factor to determine allograft survival. Thus, it is reasonable to believe that appropriate control of DSA is directly linked to well-managed immunosuppression, resulting in free from AMR. First, in order to prevent and manage AMR, it is of vital importance to be familiar with updated knowledge regarding crossmatch test and DSA detection methods, including intra-graft DSA. Second, it is also crucial to understand the standard criteria to diagnose AMR. Although pathological diagnosis and serum DSA (s-DSA) detection play the central role, the recent trend seems to be detection of intra-graft DSA (g-DSA). Third, regarding organ transplantation between sensitized pairs, the acceptable outcomes are obtained owing to recent preoperative desensitization protocols: depletion/modification of B cells, apheresis for antibodies, and inhibition of reaction between DSA and HLA. Finally, we would like to discuss the treatment of AMR. Further advances in diagnosis methods and emergences of effective treatments would be expected for acceptable control of AMR. In this chapter, we will review from the basics to recent topics in order to understand DSA and AMR.

Keywords: organ transplantation, antibody-mediated rejection, donor-specific anti-HLA antibodies, intra-graft donor-specific anti-HLA antibodies, immunocomplex capture fluorescence analysis
1. Introduction

Recent advances in immunosuppression permit for organ transplantation between sensitized recipients and donors with acceptable outcomes. However, it is true that the management of acute or, in particular, chronic antibody-mediated rejection (AMR) due to donor-specific anti-HLA antibodies (DSA) is still a crucial issue to improve long-term graft survival. Because chronic changes of AMR are irreversible, it is also true that an early accurate diagnosis of AMR is required to prevent severe consequences. Thus, it is reasonable to believe that DSA is a main research topic to improve the outcome of organ transplantation. In the 1960s, the introduction of azathioprine brought the beginning of contemporary organ transplantation [1]. Following this era, incompatibilities, anti-AB [2, 3] and anti-HLA antibodies [4], were recognized as a crucial barrier for organ transplantation. In the next half-century, the main attention was paid to cellular rejection: T-cell-mediated rejection [5]. This resulted in the development of calcineurin inhibitors [6] and several antibody drugs [7]: a depletion of lymphocytes that enabled the control of T-cell-mediated rejection. However, AMR remains an important issue that is still not addressed. Then, DSA have finally received strong attention in the twenty-first century because DSA are important factors to determine long-term graft survival. In this chapter, we will review DSA in organ transplantation and discuss effectiveness of a novel application of immunocomplex capture fluorescence analysis (ICFA), as well as crossmatch examinations, protocols of desensitization, and outcomes of crossmatch positive organ transplantation.

2. Assessment for donor-specific anti-HLA antibodies

To assess reactive DSA in recipient serum, a crossmatch test or measurement of DSA is performed routinely in clinical organ transplantation. There are several crossmatch methods: lymphocyte cytotoxic test (LCT), flow cytometry crossmatch (FCXM), and ICFA. In addition to crossmatch test, antibody detection methods such as flow PRA screening and single antigen bead assay (SAB) are utilized in clinical settings. Generally, it is important to understand the advantages and disadvantages of these methods and interpret appropriately.

2.1. Lymphocyte cytotoxic test

Antigen–antibody and complement-dependent reactions are observed in this traditional direct-crossmatch test. Donor lymphocytes are incubated in the recipient serum, followed by the addition of complements. Under the circumstances where there are DSA in the recipient serum, lymphocytes are necrotized by the complement-dependent cytotoxicity reaction. Then, adding eosin dye, the ratio of necrotized lymphocytes is counted by a phase-contrast microscopy (Figure 1). The disadvantages of LCT are relatively low sensitivity, difficulty in obtaining donor alive lymphocytes, and subjective judges. Furthermore, it should be noted that sometimes non-DSA reaction can be observed. Conversely, only this traditional method is capable of visualizing real reactions against donor cells, including non-DSA reactions [8]. Thus, it is true that careful attention should be paid when positive reaction in LCT is detected [9].
2.2. Flow cytometry crossmatch

FCXM is an examination that flow cytometry technology is applied. As characteristics of FCXM test, sensitivity is high enough to detect a slight amount of DSA. Therefore, it should be paid attention that non-HLA antibodies might be detected: false positive. Because of high-sensitivity examination, rigorous quality control is required. To apply FCXM, donor lymphocytes and recipient serum samples were utilized. Subsequently, antihuman IgG antibodies are added and positivity is determined [10]. In addition, to add CD3 or CD19, T cells and B cells can be separated and analyzed simultaneously (Figure 2).

2.3. Immunocomplex capture fluorescence analysis

WAKFlow HLA antibody class I and II ICFA (Wakunaga Pharmaceutical Co., Ltd., Osaka, Japan) is one of the crossmatch tests by using donor lymphocytes and recipient serum [11]. This technique can detect DSA immunocomplex with high specificity [12]. As a detection system, Luminex xMAP technology (Luminex Corporation, Austin, TX) is applied. Schematic presentation of ICFA is shown in Figure 3. First, HLA and DSA complexes are formed following the reaction between donor lymphocytes and recipient serum containing DSA. Second, lymphocytes are lysed and complexes remain in lysates. Third, these complexes are captured by anti-HLA monoclonal antibodies fixed on Luminex beads, and subsequently PE-conjugated human anti-IgG is added. Finally, Luminex system detects these PE-conjugated anti-human
IgG signals. As ICFA characteristics, the following features are noted: (1) the specificity of identifying HLA antibodies is high, and (2) class I and II antibodies can be identified separately. Furthermore, in terms of recent advances in ICFA, DR, DQ, and DP, DSA can be identified separately.

### 2.4. FlowPRA screening

To identify anti-HLA antibodies in serum, FlowPRA screening test is performed. First, a reaction is caused between anti-HLA antibodies and latex beads coated with HLA antigens. Then,
FITC-conjugated antihuman IgG is added. Subsequently, mean fluorescence intensity (MFI) and shift from negative control are calculated based on flow cytometry analyses (Figure 4). Each mixed class I and II HLA antigen is separately coated on latex beads, derived from about 30 types of panel cells [13]. Depending on the human race, there is a possibility that rare HLA

![FlowPRA (Screening)](image)

**Figure 4.** FlowPRA screening. (a) Schematic presentation of FlowPRA screening. (b) Examples of positive FlowPRA screening results about class I (upper) and class II (lower).

![Single Antigen bead assay](image)

**Figure 5.** Schematic presentation of single antigen bead assay (SAB).
antigens are not covered. These disadvantages should be recognized. Generally, if a positive reaction would be observed, the following SAB is applied to identify the specificity of anti-HLA antibodies.

2.5. Single antigen bead assay

To identify the specificity of anti-HLA antibodies, SAB is performed when crossmatch test and/or FlowPRA is positive. First, a reaction is observed between anti-HLA antibodies and Luminex beads coated with a single HLA antigen extracted from gene-modified cells. Following steps are similar to those of FCXM. Finally, these reactions, MFI of PE signals, were calculated by Luminex system (Luminex Corporation). According to HLA typing of donor, the presence of DSA is judged. It is often used in LABScreen single antigen HLA class I/II beads: LABScreen single antigen class I/LABScreen single antigen class II (One Lambda Inc., Canoga Park, CA) or WAKFlow HLA antibody class I HR and WAKFlow HLA antibody class II HR (Wakunaga Pharmaceutical Co., Ltd) (Figure 5). As an important point, because these single HLA antigen beads do not include all types of HLA antigens, we should be familiar with SAB kit to judge the existence of DSA appropriately.

3. Assessment for intra-graft donor-specific anti-HLA antibodies

Due to recent advances in examinations for DSA, the assessment for g-DSA has been paid attention. In fact, it is hard to understand that s-DSA damage allografts without localization in target organs. Thus, hereafter, the assessment of g-DSA would gain more popularity as diagnosis or prognosis factors. Although the presence of g-DSA is not included in AMR diagnosis criteria currently, g-DSA might be a key criterion for considering AMR in the near future. It would be better that clinicians and researchers are aware of this novel topic. Here, we will present representative two different methods. We will also delve into graft ICFA technique in this section.

3.1. Dissociation between HLA and DSA (acid elution method)

To obtain free DSA from allografts, dissociation HLA and DSA complexes are attempted. So far, g-DSA detection in the kidney [14–16], liver [17], and lung [18] were reported. This method requires the following steps: (1) Wash more than seven times to prevent from detection s-DSA incorrectly. (2) Dissociate these complexes by acid (buffer). (3) Detect dissociated DSA by SAB (Figure 6). As compared to graft ICFA, mentioned later, acid elution method has weak points regarding simplicity and remains doubtful points whether DSA denature or not during the acid elution step. However, SAB analysis following acid elution seems to be accepted widely and allows to identify specific DSA even where multiple candidate DSA exist. The common recognition of g-DSA assessment seems to be that g-DSA is an important factor to determine graft survival and more sensitive than s-DSA [14].

3.2. Graft immunocomplex capture fluorescence analysis (non-dissociation technique)

WAKFlow HLA antibody class I and II ICFA is an attractive tool to identify HLA/DSA complexes as mentioned above, by means of WAKFlow HLA antibody class I and II (Wakunaga
Pharmaceutical). It is not only for serum of recipients but also for allograft specimens. The first graft ICFA was introduced as an effective tool for detection of g-DSA in 2017 [19, 20]. Graft ICFA can be performed as previously described by Nakamura et al. Graft samples were obtained by means of a percutaneous needle biopsy. To standardize the results of graft ICFA, 2 mm specimen was used for each analysis. Regarding graft ICFA to identify HLA expression, samples were washed enough in PBS. As compared to acid elution method of DSA from grafts associated to g-DSA detection by SAB, graft ICFA does not allow to identify individual responsible HLA alleles, such as HLA-A24, etc. However, in the setting of real organ transplantation, generally HLA alleles are identified prior to transplantation. Thus, it does not seem to be difficult to narrow down candidate HLA. It is true, therefore, that current graft ICFA is clinically useful to diagnose AMR. Moreover, the combination of acid elution method and graft ICFA allows to obtain useful information regarding g-DSA. Further advances in graft ICFA would be expected.

(Ethics Committee approval was obtained from the internal research ethics committee of Kyoto Prefectural University of Medicine. The clinical trial registration number is UMIN000023787.)

3.3. Data interpretation

MFI of samples was calculated by the Luminex system. A ratio of sample MFI/blank beads MFI of all negative samples, including HLA matched recipients’ samples, + 2SD is demonstrated below 0.9 (data not shown) (please refer to [19, 20]). Then the ratio was determined, and ≥ 1.0 was considered as a positive result. Furthermore, to compensate baseline reaction, the following index was also calculated. Index = \((X_{\text{HLA}}- (N_{\text{HLA}}-N_{\text{BB}}) X_{\text{BB}}/N_{\text{BB}}) / X_{\text{BB}}\) \(X_{\text{HLA}}\) sample
MFI; $X_{\text{mbr}}$, sample blank beads MFI; $N_{\text{HLA}}$, the mean MFI of negative samples; and $N_{\text{mbr}}$, the mean blank beads MFI of negative samples. Given the results of negative samples, the index $\geq 1.5$ was considered as a positive result.

3.4. DSA-HLA complexes in the liver, heart, lungs, pancreas, and small intestine are also successfully detected by graft ICFA

To confirm whether graft ICFA can be applied to other organs besides the kidney [19, 20], we employed ten liver transplant recipients and a liver transplant recipient who underwent autopsy due to primary graft dysfunction. Samples of other possible organs such as the heart, lung, pancreas, and small intestine from this patient were pretreated according to the graft ICFA preparation method. Contaminated blood cells in samples were minimum, confirmed by histology. Luminex analyses detected PE signals from the positive control samples in all organs. Thus, it can be concluded that graft ICFA can be applied for all organ transplantation (Figure 7).

3.5. Sensitivity and specificity of graft ICFA to determine pathological AMR in renal transplantation

In order to prove a hypothesis—graft ICFA is useful to detect g-DSA and AMR—a total of 40 Japanese renal transplant recipients were included prospectively. They underwent graft biopsy and were examined by graft ICFA as previously described [19, 20]. According to the results of graft ICFA, these patients were divided into the g-DSA+ and g-DSA- groups to assess

![Figure 7. Intra-graft DSA (g-DSA) in all possible allografts can be analyzed by graft ICFA.](image)
the sensitivity and specificity of graft ICFA to predict pathological AMR. In the current study, this technique has demonstrated 100% sensitivity (12/12) and 92.9% specificity (26/28). Thus, it is reasonable to believe that positive graft ICFA results strongly suggest the onset of AMR.

4. Diagnosis of antibody-mediated rejection

4.1. Renal transplantation

With renal allograft dysfunction, it can be diagnosed as AMR, provided that C4d deposited in peritubular capillaries (ptc) and antibodies or complement deposition are confirmed in vascular fibrinoid necrosis, in addition to s-DSA detection. Recently, AMR can be diagnosed according to the Banff classification 2015 [21].

Acute AMR:
The following three criteria should be met when acute AMR is diagnosed:

1. Histological features of acute tissue injury, including at least one of the following
   • Microvascular inflammation g > 0 (excluding recurrent or de novo glomerulonephritis), ptc > 0, or v > 0 (intimal/transmural arteritis).
   • Acute thrombotic microangiopathy (without any other apparent reasons).
   • Acute tubular injury (without any other apparent reasons).

2. Histological features due to DSA and vascular endothelium reaction
   • Linear C4d deposition in ptc (C4d2 or C4d3 (frozen sections)/C4d > 0 (paraffin embedded sections))
   • Moderate microvascular inflammation g + ptc ≥ 2.
   • Detection of genetic transcript expression in biopsy specimens due to endothelial injury.

3. Detection of s-DSA

Chronic AMR:
The following three criteria have to be met:

1. Histological features of chronic tissue injury, including at least one of the following
   • Transplant glomerulopathy (cg > 0), in the case of no chronic thrombotic microangiopathy, includes histologic features detected by electron microscope (EM).
   • Severe ptc basement membrane multilayering by EM.
   • New onset of arterial intimal fibrosis (without any other apparent reasons).
2. Histological features due to DSA and vascular endothelium reaction

- Linear C4d deposition in PTC (C4d2 or C4d3 (frozen sections)/C4d > 0 (paraffin embedded sections))

- Moderate microvascular inflammation g + ptc ≥ 2.

- Detection of genetic transcript expression in biopsy specimens due to endothelial injury.

3. Detection of serum DSA

![Figure 8. Histopathological impacts of g-DSA presence. A. The Banff histologic scores are analyzed individually based on the g-DSA status: G-DSA- or g-DSA+. B. The Banff histologic scores based on the g-DSA status: G-DSA- or g-DSA+ without ABO-incompatible cases. C. These items are reanalyzed depending on g-DSA values: G-DSA-, g-DSA < 10, and g-DSA ≥ 10, including ABO-incompatible cases. ****p < 0.0001, ***p < 0.0005, **p < 0.005, and *p < 0.05.](image-url)
4.1.1. The presence of g-DSA (graft ICFA) is associated with microvascular lesions in renal transplantation

To confirm the consistency of graft ICFA results, the individual scores of the Banff classification were analyzed between g-DSA- and g-DSA+ renal transplant recipients (g-DSA+ 15, g-DSA- 25 recipients). As a result, individual g, cg, ptc, and ptc-bm scores were significantly higher in g-DSA+ patients. Interestingly, there was no apparent difference in the C4d staining score, primarily due to the presence of ABO-incompatible cases in both groups (32.0% and 33.3% in the g-DSA- and + groups, respectively) (Figure 8A). Thus, ABO-incompatible cases were removed from both groups. Then, the C4d result showed that 0.45 ± 0.17 and 1.73 ± 0.33 in the g-DSA- and g-DSA+ groups, respectively (p = 0.0184) (Figure 8B). Next, g-DSA+ patients were divided into low g-DSA group, g-DSA+ < 10, and high g-DSA group, g-DSA+ ≥ 10. Then, this result demonstrated that g and ptc deteriorated in g-DSA concentration manner. In contrast, only g-DSA+ ≥ 10 group showed significant higher scores in cg, mm, ptc-bm, and C4d (Figure 8C). These results might indicate that g-DSA causes microcirculation lesions and high g-DSA means chronic allograft damages. To correspond to a recent functional concept, g + ptc (microvascular inflammation), g + cg + ptc (microvascular lesions), and cg + mm (microvascular deterioration), we also analyzed these scores again. Expectedly, g + ptc and g + cg + ptc deteriorated stepwise according to g-DSA scores, but cg + mm referring chronic lesions is clearly higher only in the g-DSA ≥ 10 group (Figure 9). It is true, therefore, that g-DSA assessment by graft ICFA accurately supports the diagnosis of AMR.

4.2. Liver transplantation

Generally, it is often discussed that liver allografts tend to be resistant against AMR due to Kupffer cell DSA clearance, HLA expression in microvasculature, allografts size, and regenerative capacity of the liver. Diagnosis criteria are suggested in the Banff meeting [22].

Define acute AMR:

The following four criteria should be met to diagnose acute AMR:

1. Histological features of acute tissue injury, including at least one of the following Portal microvascular endothelial cell hypertrophy, eosinophilic and neutrophilic portal microvasculitis, portal edema, and ductular reaction; cholestasis is usually present but variable; edema and periportal hepatocyte necrosis, active lymphocytic, and/or necrotizing arteritis.

2. Positive s-DSA.

3. Diffuse microvascular C4d staining (C4d = 3).

4. Excluding other lesions possibly mimic AMR.

Suspicious for AMR (both criteria required).

Serum DSA + and positive histopathology score (h-score) (C4d + h-score ≥ 3).
Indeterminate for AMR:

1. C4d + h-score ≥ 2.
2. DSA and C4d immunohistochemistry not available and not apparent.
3. Coexisting pathophysiology might cause similar injury.

Liver allograft chronic active AMR is also considered as an important factor for determining long-term outcome [23, 24]. Chronic active AMR is diagnosed when the condition meets the all following criteria:

1. Histologic features consistent with chronic active AMR. (a) At least mild mononuclear portal and/or perivenular inflammation with interface and/or perivenular necroinflammatory activity. (b) At least portal/periportal, sinusoidal, and/or perivenular fibrosis.
2. Detection of serum DSA.
3. Focal C4d deposition in portal tract microvascular endothelia (> 10%).
4. Other lesions can be denied.

4.3. Heart transplantation

Following heart transplantation, the onset of AMR is generally estimated around 10–20% [25]. In the field of heart transplantation, AMR is also considered as an important prognosis factor. In fact, the consequences are severe, and the development of cardiac allograft vasculopathy (CAV) has a huge impact. As with other organ transplantations, timing of AMR onset is diverse: both acute AMR within 1 week after transplantation and chronic AMR in the remote period can be seen. Again, allograft biopsy also plays a crucial role to accurately diagnose AMR. Repeated allograft biopsies seem to be required for achieving good long-term outcomes. Interestingly, regarding de novo DSA synthesized after 2 months following heart transplantation, class II DSA are dominant [25].

Diagnosis criteria of heart allograft AMR were discussed in the international society for heart and lung transplantation working formulation consensus [26]. First of all, substrates were divided into two categories: histologic investigations (H) and immunopathologic studies (I). Histologic features of heart AMR are as follows: activated mononuclear cell infiltration, interstitial edema, hemorrhage, necrosis, and vascular thrombosis. On the other hand, C4d, CD68 (paraffin section), C4d, or C3d (frozen section) are considered mandatory panels of immunopathologic studies. Furthermore, CD3, CD20, C3d, endothelial cell CD31 or CD34, complement regulatory proteins (paraffin section), and fibrin and immunoglobulin G/M (frozen section) are regarded as secondary or optional panels. In total, the current report of AMR in heart is noted from Grade 0 to 3 as follows:

pAMR 0: negative for pathologic AMR (H0/I0)
pAMR 1(H+): histopathologic AMR (H+/I−)
pAMR 1 (I+): immunopathologic AMR (H−/I+)
pAMR 2: pathologic AMR (H+ I+)
pAMR 3: severe AMR (interstitial hemorrhage, capillary fragmentation, mixed inflammatory cell infiltration, endothelial cell pyknosis and/or karyorrhexis, and marked edema and I+)
pAMR 3 cases usually demonstrate severe hemodynamic dysfunction and poor consequences.

Scoring system of criteria for pathologic diagnosis of cardiac AMR is slightly different between immunohistochemistry and immunofluorescence analyses. These criteria are summarized in Table 1.

4.4. Lung transplantation

Regarding AMR following lung transplantation, the main reason is also DSA, although a role of non-HLA antibodies was also described [27]. Hachem et al. [28] have reported, in 2010, a prospective study on AMR after lung transplantation, which deepened our
understanding of clinical AMR managements. In 2016, a consensus statement has been proposed from the international society for heart and lung transplantation [29].

Lung allograft AMR can be divided into clinical and subclinical AMR, depending on whether there is allograft dysfunction or not. Next, as similar to other organs, these clinical and subclinical AMR are subcategorized as define, probable, or possible. To determine certainty of clinical AMR, the following five criteria are proposed: allograft dysfunction, other causes excluded, lung histology, lung biopsy C4d, and s-DSA. Define lung AMR is determined by positive for all five criteria. In case anyone of them is negative, it is considered as probable AMR, excluding allograft dysfunction (Table 2). There are histopathologic features of AMR, including the following—neutrophil margination, neutrophil capillaritis/arteritis without any signs of pneumonia, or other apparent reasons—though these features are not specific for lung AMR. In terms of lung histology, further advancement and organization would be required.

<table>
<thead>
<tr>
<th>Immunohistochemistry scoring system</th>
<th>Immunofluorescence scoring system</th>
</tr>
</thead>
<tbody>
<tr>
<td>Capillary C4d distribution</td>
<td>Capillary C4d/ C3d distribution</td>
</tr>
<tr>
<td>0 &lt; 10% Negative</td>
<td>0 &lt; 10% Negative</td>
</tr>
<tr>
<td>1 10–50% Focal</td>
<td>1 10–50% Focal</td>
</tr>
<tr>
<td>2 &gt; 50% multifocal/diffuse</td>
<td>2 &gt; 50% Multifocal/diffuse</td>
</tr>
<tr>
<td>Capillary C4d intensity</td>
<td>Capillary C4d/ C3d intensity</td>
</tr>
<tr>
<td>0 Negative/equivocal</td>
<td>0 Negative/equivocal</td>
</tr>
<tr>
<td>1 Faint positive</td>
<td>1 Faint positive = 0–1+</td>
</tr>
<tr>
<td>2 Strong positive</td>
<td>2 Strong positive = 2–3+</td>
</tr>
<tr>
<td>Intravascular CD68 distribution</td>
<td>HLA-DR distribution</td>
</tr>
<tr>
<td>0 &lt; 10% Negative</td>
<td>0 &lt; 10% Negative</td>
</tr>
<tr>
<td>1 10–50% Focal</td>
<td>1 10–50% Focal</td>
</tr>
<tr>
<td>2 &gt; 50% Multifocal/diffuse</td>
<td>2 &gt; 50% Multifocal/diffuse</td>
</tr>
<tr>
<td>HLA-DR intensity</td>
<td></td>
</tr>
<tr>
<td>0 Negative/equivocal</td>
<td></td>
</tr>
<tr>
<td>1 Faint positive = 0–1+</td>
<td></td>
</tr>
<tr>
<td>2 Strong positive = 2–3+</td>
<td></td>
</tr>
</tbody>
</table>

Table 1. Immunohistochemistry and immunofluorescence scoring system in heart AMR.

Lung allograft AMR can be divided into clinical and subclinical AMR, depending on whether there is allograft dysfunction or not. Next, as similar to other organs, these clinical and subclinical AMR are subcategorized as define, probable, or possible. To determine certainty of clinical AMR, the following five criteria are proposed: allograft dysfunction, other causes excluded, lung histology, lung biopsy C4d, and s-DSA. Define lung AMR is determined by positive for all five criteria. In case anyone of them is negative, it is considered as probable AMR, excluding allograft dysfunction (Table 2). There are histopathologic features of AMR, including the following—neutrophil margination, neutrophil capillaritis/arteritis without any signs of pneumonia, or other apparent reasons—though these features are not specific for lung AMR. In terms of lung histology, further advancement and organization would be required.

4.5. Small bowel transplantation

Due to recent increased recognition of AMR, AMR is also considered as a serious issue in the small bowel transplantation field. However, the define diagnosis criteria have not been established in the small bowel transplantation yet. There are few series of case reports in terms of intestine AMR. Although common understandings of AMR also seem to be C4d deposition, capillaritis, and s-DSA positivity [30], there is a report which did not find clinical evidence between C4d positivity and the onset of AMR [31]. The establishment of diagnostic criteria should be required to standardize and manage AMR in intestine transplantation. The future contribution of g-DSA assessment would be also expected in this field.
5. Preoperative desensitization

To perform organ transplantation between a sensitized recipient and donor pair, preoperative desensitization is required. Generally, crossmatch positive organ transplantation is mainly performed in kidney and liver transplantation. Desensitization can be divided into three main treatments: depletion/modification of B cells, apheresis for antibodies, and inhibition of reaction between DSA and HLA. The golden standard of B-cell depletion therapy is rituximab (anti-CD20 antibodies) administration [32–35]. There are a wide variety of rituximab administration protocols in terms of a dosage and schedule. In our institution, generally, rituximab (375 mg/m$^2$) is administered in 2 weeks prior to organ transplantation [20]. Subsequent B-cell count in the peripheral blood is measured by flow cytometry. In addition to rituximab, to deplete B cells, anti-CD52 antibodies (alemtuzumab) also can be used, because higher than 95% B cells express CD52 on the cell surface [36]. It has been reported that alemtuzumab-combined regimens are safe and effective for highly sensitized recipients [36, 37]. Other antibodies’ introduction would be expected regarding induction regimens.

To decrease DSA production and reaction, the importance of intravenous immune globulin (IVIG) has been reported. IVIG infusion significantly decreased the baseline flowPRA levels [38]. There is a variety of reports regarding the dosage and duration of IVIG (100 mg/kg/day–4 g/kg/day, etc.), partially due to the cost problems.

Regarding apheresis therapy, double filtration plasmapheresis (DFPP) or plasma exchange (PE) is generally performed. Usually, as an index of DSA titer, MFI has gained its popularity. Given the fact of g-DSA and s-DSA assessments, DSA with MFI < 2500 might not deposit and cause clinical damages to allografts. Thus, although it depends on each institution, acceptable DSA MFI prior to surgery might be estimated around 2000 [20] in renal transplantation. However, on the other hand, intensive posttransplant desensitization also can result in

<table>
<thead>
<tr>
<th>Allograft dysfunction</th>
<th>Other causes excluded</th>
<th>Lung histology compatible with AMR</th>
<th>C4d</th>
<th>s-DSA</th>
</tr>
</thead>
<tbody>
<tr>
<td>Define</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Probable*</td>
<td>+</td>
<td>+</td>
<td>–</td>
<td>+</td>
</tr>
<tr>
<td>+</td>
<td>–</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>+</td>
<td>+</td>
<td>–</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>+</td>
<td>+</td>
<td>+</td>
<td>–</td>
<td>+</td>
</tr>
<tr>
<td>+</td>
<td>–</td>
<td>–</td>
<td>+</td>
<td>–</td>
</tr>
<tr>
<td>+</td>
<td>–</td>
<td>+</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>+</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
</tbody>
</table>

There is growing evidence that C4d negative AMR exists. Thus, the second line * cases are considered as an independent group.

Table 2. Lung AMR diagnosis criteria.

5. Preoperative desensitization

To perform organ transplantation between a sensitized recipient and donor pair, preoperative desensitization is required. Generally, crossmatch positive organ transplantation is mainly performed in kidney and liver transplantation. Desensitization can be divided into three main treatments: depletion/modification of B cells, apheresis for antibodies, and inhibition of reaction between DSA and HLA. The golden standard of B-cell depletion therapy is rituximab (anti-CD20 antibodies) administration [32–35]. There are a wide variety of rituximab administration protocols in terms of a dosage and schedule. In our institution, generally, rituximab (375 mg/m$^2$) is administered in 2 weeks prior to organ transplantation [20]. Subsequent B-cell count in the peripheral blood is measured by flow cytometry. In addition to rituximab, to deplete B cells, anti-CD52 antibodies (alemtuzumab) also can be used, because higher than 95% B cells express CD52 on the cell surface [36]. It has been reported that alemtuzumab-combined regimens are safe and effective for highly sensitized recipients [36, 37]. Other antibodies’ introduction would be expected regarding induction regimens.

To decrease DSA production and reaction, the importance of intravenous immune globulin (IVIG) has been reported. IVIG infusion significantly decreased the baseline flowPRA levels [38]. There is a variety of reports regarding the dosage and duration of IVIG (100 mg/kg/day–4 g/kg/day, etc.), partially due to the cost problems.

Regarding apheresis therapy, double filtration plasmapheresis (DFPP) or plasma exchange (PE) is generally performed. Usually, as an index of DSA titer, MFI has gained its popularity. Given the fact of g-DSA and s-DSA assessments, DSA with MFI < 2500 might not deposit and cause clinical damages to allografts. Thus, although it depends on each institution, acceptable DSA MFI prior to surgery might be estimated around 2000 [20] in renal transplantation. However, on the other hand, intensive posttransplant desensitization also can result in
comparable outcomes with non-sensitized transplantation [39]. For highly sensitized recipients, a phased desensitization protocol by using rituximab and bortezomib was advocated [34]. In fact, it is true that rituximab administration can deplete B cells but not plasma cells. Thus, it is reasonable to believe that rituximab and bortezomib combination therapy eradicates the B-cell lineage which is potentially associated with AMR.

Furthermore, the idea only relying on MFI might be unrefined. In other words, the quality of DSA is also important to determine the impact of DSA. Regarding the quality, IgG subclasses [1–4, 40] and complement fixing ability [41] seem to be paid attention, considering the severity of subsequent AMR and graft survival.

On the other hand, in liver transplantation, there seems to be no concrete evidence regarding DSA MFI just prior to transplantation. A large amount of hemorrhage during surgery and liver allograft resistance against DSA, etc., might complicate to set a MFI threshold. Nevertheless, Yoshizawa et al. [42] reported that class I DSA MFI > 10,000 has a negative impact on graft survival. Thus, it is important to keep circumstances where allograft injury due to remnant DSA is minimum and additional DSA production inhibited.

6. Treatment for antibody-mediated rejection

In other words, this is a treatment for B cells/plasma cells and DSA and reaction between DSA and HLA. Both for acute AMR and chronic active AMR, generally clinical managements also can be divided into medications and apheresis: steroid pulse, IVIG, Rituximab, etc., and DFPP or PE. In severe cases, it is true that splenectomy has a certain effect on AMR [43]. Regarding treatment for acute AMR, the concepts are the same: depletion of B cells, reduction of DSA, and inhibition of reaction between DSA and HLA. The core agents and methods are summarized in Table. It is true that high-dose steroid administration is effective on all aspects of AMR treatment. There is no fixed data to determine the dosage and duration of steroid pulse therapy. However, generally, 10–100 mg/kg/day equivalent dosage of hydrocortisone is administered as steroid pulse therapy, depending on the severity of AMR. For depletion of B cells, rituximab, alemtuzumab, or splenectomy is utilized. To reduce DSA, DFPP, or PE, apheresis methods are commonly used in the same way as desensitization. As immunomodulation, IVIG administration also plays an important role in controlling acute AMR. Given the fact of AMR pathogenesis, complement activation should be paid attention. Final tissue injury due to AMR would occur following activation of antibody-induced terminal complement cascade. Albeit limited evidence, it has been reported that eculizumab C5 inhibitor is effective to rescue an AMR allograft [44–47].

Recently, chronic active AMR has been paid strong attention, because this pathologic condition directly deteriorates the long-term graft survival. Despite the recognition of chronic active AMR, diagnostic criteria are only established in kidney [21] and liver transplantation [22]. There seems to be no therapeutic consensus on this condition. Furthermore, it has generally resistance to ordinal AMR managements discussed above [48–50], although limited effectiveness was observed in few studies [51, 52]. Bachelet et al. [50] reported a treatment for chronic active AMR (mean eGFR 30.6 mL/min/1.73 m²) by utilizing rituximab (375 mg/m²) and IVIG (1 g/kg/week × 4 weeks). Although serum MFI tends to decrease, there is no difference in

Human Leukocyte Antigen (HLA)
2-year graft survival between the treatment (47%) and without treatment groups (40%). These reports suggest a difficulty in the management for chronic changes in allografts. In addition to acute AMR, eculizumab was also challenged for chronic active AMR. Although there were no notable differences in eGFR between treatment and control groups, C1q-positive recipients demonstrated significant better eGFR than recipients with C1q-negative status. Inhibition of complement-dependent allograft injury would bring benefits on certain population. Recent reports discussing treatment for chronic active AMR are summarized in Table 3.

In total, it is of vital importance to prevent from developing into chronic lesions and initiate appropriate treatment in the early stages of AMR, because fully established chronic lesions are irreversible. It is reasonable to believe that these managements lead to improvements of the long-term allograft survival.

### 7. Conclusion

Under the present circumstances, it is of vital importance to control AMR in advance in order to improve graft survival rate in all fields of organ transplantation. To detect early-stage AMR, clinicians need to be aware of recent advances in DSA analyses, including graft ICFA and an acid elution method to assess intra-graft DSA status. Preoperative desensitization therapies and management plans are decided depending on classes of DSA and s-DSA MFI. Conversely,
even in crossmatch-negative cases, there is a possibility that memory B cells might evoke severe AMR 1 week following transplantation. In addition, there might be a discrepancy between s-DSA and g-DSA. It is also true that only relying on s-DSA MFI is difficult to determine appropriate managements. Further research is required for addressing these issues.

Conflict of interest

None.

Abbreviation

AMR antibody-mediated rejection  
DSA donor-specific antibodies  
ICFA immunocomplex capture fluorescence analysis  
LCT lymphocyte cytotoxic test  
FCXM flow cytometry crossmatch  
SAB single antigen bead assay  
MFI mean fluorescence intensity  
g-DSA intra-graft DSA  
s-DSA serum DSA  
ptc peritubular capillaries

Author details

Tsukasa Nakamura*, Hidetaka Ushigome¹, Takayuki Shirouzu² and Norio Yoshimura¹  
*Address all correspondence to: tsukasa@koto.kpu-m.ac.jp  
1 Department of Organ Transplantation and General Surgery, Kyoto Prefectural University of Medicine, Kyoto-prefecture, Japan  
2 Wakunaga Pharmaceutical Co., Ltd. Molecular Diagnostics Division, Japan

References


