Liver, rejection and tolerance
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INTRODUCTION

Clinical and experimental observations indicate that liver occupies a singular position relatively to other solid organs such as kidney, heart or lung. In some animal species as pig or some breeds of rat, liver transplantation can be performed without immunosuppression [1-3].

In humans, studies have been conducted where LT recipients were completely weaned off pharmacological immunosuppression [4] without rejection. Moreover, LT performed simultaneously with other solid organ of the same donor, prevent the rejection of this organ [5,6]. Liver can even reverse the rejection of another allograft, of another donor, transplanted a few days before the liver [7,8]. All these observations make us consider the liver to have inherent tolerogenic properties [9]. While this phenomenon was better understood, we discovered in the same time, that antibodies mediated rejection (AMR) was perhaps not as seldom in LT [10] as we thought, contrary to its fundamental role in other solid organ transplantations [11,12]. In this review, we will first expose the mechanisms of rejection and tolerance in LT, then we will describe in its complexity, the present knowledge of AMR in LT. At last, we will discuss the immunomodulatory role of the liver in multiple transplantations and expose the different hypothesis about its mechanism.

I - PRELIMINARY CONSIDERATIONS ABOUT REJECTION IN LIVER TRANSPLANTATION

1. Mechanisms and types of rejection in liver transplant

After solid organ transplantation, hyperacute vasculitis rejection can...
occur in animal or human with preformed antibodies (Ab) against the donor major histocompatibility complex (MHC) class I-encoded antigens\[13,14\]. This Ab mediated attack of the recipients’ cells leads to the necrosis of the organ in the hours after the reperfusion. In the case of the liver, this hyperacute rejection could occur in ABO incompatible transplantation because ABO antigens are present on the endothelial cells. In most other clinical situations, acute allograft rejection is initiated by recipient T cells that recognize donor antigens, mostly alloantigens encoded by the polymorphic MHC. Donor MHC molecules are internalized by donor and recipient antigen-presenting cells (APC). Molecular and cellular basis of graft rejection have been exhaustively reported elsewhere\[15\]. Rapidly, three different non exclusive pathways could be implicated in the allograft rejection\[16\]: first, the direct pathway: recipient T cells recognize intact allogenic MHC molecules (class I and II) on the surface of donor APCs, secondly the indirect pathway: recipients T-cell recognize donor antigen in recipient MHC molecule on the surface of recipient APC. These donors Ag are the product of the phagocytosis and trafficking of necrotic donor cells material in recipient APCs. Third, the semi direct pathway in which recipient APCs acquire intact donor MHC molecules following direct contact with donor APCs or through fusion with donor APC-derived exosomes. Acute cellular rejection is the best-characterized graft-specified presentation of immune rejection, especially in liver transplantation. It occurs most frequently in the period between 5 and 21 days after transplantation, its onset may be as early as three to four days or at least as several years following transplantation, in this case most frequently after interruption of immunosuppressive therapy. It is defined by an often –sudden deterioration in allograft function. Biopsy analysis of the transplanted tissue makes the diagnosis: in the liver, it shows infiltration by host T cells and other mononuclear leukocytes in portal veins, bile ducts and central vein endothelium\[17\]. Although patients with documented acute cellular rejection would undergo improvement in liver function tests and exhibit biopsy-proven reversal of acute rejection without administration of specific antirejection therapy\[18\]. Acute rejection should usually be treated, even in liver transplanted patients, to prevent the irreversible deterioration of the organ.

Beside the cellular mechanism of rejection, the production of antibodies is also associated with acute and chronic rejection in organ transplantation. The role of anti-donor MHC class I and II has been well identified in the past five years in the kidney\[19,20\], cardiac\[21\] and pancreatic\[22\] transplantation. Antibodies can injure the graft by activating complement and mononuclear cells with Fe receptors that recognize the heavy chain of antibody. Fe receptor-expressing leukocytes can thereby be activated by antibodies-coated donor cells. Anti-donor antibody can also inhibit signaling cascades within endothelial cells\[23\] resulting in the common form of chronic rejection, the transplant vasculopathy. Over time, transplanted organ present inexorable deterioration of the function. Although this process is called chronic rejection, the donor-specific immune process is probably not the sole, and even not the primary cause in liver transplant\[24\]. In liver transplant, chronic rejection differs from acute rejection, in that it is more insidious in onset, does not respond to methylprednisolone boluses, and has a histopathologic picture that is characterized by loss of bile ducts and arteriole obstruction by foamy macrophages. Among the factors that are thought to increase the likelihood of chronic rejection are prior episodes of acute cellular rejection, chronic ischemia secondary to hepatic artery insufficiency or thrombosis; cytomegalovirus infection or recurrent infection with hepatitis C virus; and perhaps chronic antibody-mediated injury as we will discuss this point. It is likely that chronic rejection occurs in grafts that are damaged or fail for many non-immunologic reasons and histologic analysis reveals fibrosis in the absence of immune cell infiltration.

### 2. Interaction between ischemia-reperfusion, inflammation and immune reaction

Newly engrafted organ is subject to intense inflammation. The injury to the graft, caused by donor disease (steatosis, stroke)\[25\] or hemodynamic instability, organ procurement or reanimation, cold and warm preservation, surgical trauma and reperfusion injury, leads to release of reactive oxygen species (ROS) and proinflammatory cytokines. Following liver ischemia reperfusion, there is activation of tumor necrosis factor-α (TNF-α) by chemokine (CXCL) 10, interferon regulatory factor (IRF) and toll-like receptor (TLR) (innate immunity) in parallel. TNF-α activates downstream hepatocyte/sinusoidal endothelial cells (SEC) nuclear factor kb (NFkb) and CD4+ T cells separately which activate c-Jun N-terminal protein kinase-2 (JNK-2) and signal transducer activator of transcription-4 (STAT4), respectively leading to increased cell injury\[26\]. In innate natural killer, nuclophilic polynuclears, macrophages, platelets and alloimmune (lymphocytes T and B, antigen presenting cells, dendritic cells) reactions are both stimulated. It has been demonstrated that ischemia-reperfusion injury of liver allografts was associated with a reduction in the expression of immune response genes and promotion of those involved in protection and repair\[27\].

### II-TOLERANCE AND LIVER TRANSPLANTATION

#### 1. Definitions

Immunologically, tolerance is defined as the absence of an immune response towards a specific antigen in the absence of immunosuppression. In contrast to clinical transplantation, animal models of organ transplantation allows a straightforward mean of detecting immunological tolerance\[27\]. An animal is formally proven to be tolerant when, in the absence of immunosuppression, a second graft from the same donor is accepted, while a graft from a third-party donor is rejected\[28\]. Since such unambiguous assays clearly cannot be used in clinical organ transplantation, the surrogate definition of operational tolerance has been established. Operational tolerance is defined as the absence of graft rejection without the use of immunosuppressive drugs\[28\].

#### 2. Mechanisms of tolerance

**2.1. Central tolerance:** In central thymic tolerance, thymocytes CD4+CD8+ are initially positively selected to become CD4+ or CD8+ depending on their TCR interactions with thymic stroma cells (dendritic cells, macrophages, thymic epithelial cells) which strongly express MC class I (CD8) and II (CD4) molecules. If MHC selection fails, the double positive thymocytes undergo apoptosis in 95 percent of case. Negative selection follows for single positive cells, in which the most self-reactive CD4+ or CD8+ cells undergo TCR-induced death if they interact with MHC molecules on APCs carrying self-peptides. Alloreactive cells can also undergo negative selection in the thymus during the interaction of antigen-specific T cells with tolerogenic dendritic cells expressing varying levels of costimulatory molecules.

**2. 2. Peripheral tolerance:** However, some CD4+ or CD8+ cells can escape negative selection and become auto/alloreactive T lymphocytes in the periphery (lymph nodes, blood, spleen). That can cause autoimmune disease or allograft rejection. Fortunately these high-reactive cells, escaping thymic deletion may be deleted
or regulated in the periphery, by activation-induced cell death, apoptosis or the suppressive action of T regulatory lymphocytes (Treg) and cytokines. Activated lymphocytes can also undergo apoptosis because of the absence of the antigen or the costimulatory factors. Moreover, cell surface receptors PD-1 and its inducible ligand PD-L1 down-regulate T cell activation, inducing tolerance. Another mechanism of peripheral concerns immature APCs such immature DC or plasmacytoid DC, that are poor presenters of alloantigens on MHC molecules, especially in tolerant environment (IL-10, IDO, PD-L1, Fas L, HO-1, HLA-G) resulting in deletion of Teff, induction of Treg and anergy/deletion of memory T [39]. However, a major drawback in vivo is the potential of tolerogenic DCs to mature during infections or inflammation, which would convert them into immunogenic cells [31].

In vitro propagation of tolerogenic donor- or recipient-derived DC has been used extensively as an experimental approach to target the pathways of allore cognition, with the aim of prolonging transplant survival, while reducing dependency on immunosuppressive drugs. The development of techniques to propagate large numbers of TOL (operationally tolerant patients) DC in vitro has provided the basis for ascertaining the ability of these cells to down regulate both host-versus-graft and graft-versus-host immune responses mediated by T cells (in the context of hematopoietic stem cell transplantation). DCs can be treated with pharmacologic agents before injection, which may attenuate their maturation in vivo. Genetically engineered DCs have also been tested. There have been numerous reports of indefinite murine allograft survival following infusion of either donor- or recipient-derived TOL DC [32-34]. Clinical trials in human including kidney transplantation are in progress.

Interestingly, in a recent study [35], B cell phenotypes were followed in blood of operationally tolerant kidney transplanted patients (TOL) and compared patients with stable graft function treated with immunosuppressive therapy. Apoptosis, proliferation, cytokine, immunoglobulin production and markers of differentiation were followed in blood of these operationally tolerant patients (TOL). Tolerant recipients show a higher frequency of CD20+CD24hiCD38lo transitional and CD20+CD24hiCD38hi naive B cells compared to patients with stable graft function, correlating with a decreased frequency of CD20+CD38hiCD138lo differentiated plasma cells, suggestive of abnormal B cell differentiation. B cells from TOL proliferate normally but produce more IL-10. In addition, B cells from tolerant recipients exhibit a defective expression of factors of the end step of differentiation into plasma cells and show a higher propensity for cell death apoptosis compared to patients with stable graft function. This in vitro profile is consistent with down-regulation of B cell differentiation genes and anti-apoptotic B cell genes in these patients in vivo. These data suggest that a balance between B cells producing IL-10 and a deficiency in plasma cells may encourage an environment favorable to the tolerance maintenance.

Other authors studied the difference of the immune response in tolerant liver or kidney transplanted patients [32]. Whereas in liver transplant tolerance the major cell types involved appear to be NK cells and γδTCR+T cells, in kidney transplantation the predominant cell type appears to be B cells. The term of B regulatory cells (Breg) for this type of B cells with tolerant properties, already defined for auto-immune pathologies, was applied to organ transplantation [37]. The induction and maintenance of immune tolerance is a complex mechanism involving not one cell subset alone but many types of cells (Figure 1), especially Tregs and iDC, with intricate interactions. Moreover, the immunological and inflammatory environment is crucial to determine the state of each cell type which will drive a tolerogenic or immunologic response. A major point in clinical care for operationally tolerant transplant recipients is the ability to monitor for its stability. Several papers have already described transplant patients that lost their tolerant state towards the allograft and developed a rejection response [33,34]. Brouard et al showed that in two kidney transplanted patients the occurrence of rejection followed a loss of the tolerogenic phenotype. Some situation like infection may reverse tolerance but the mechanism responsible for the switch from tolerance to immunogeneity is not yet clarified in the field of liver immunology. For example, why any liver recipients with recurrent HCV (hepatitis C virus) infection and HCV specific T cells do not usually reject despite infiltration of the liver graft? In some of these cases, operational tolerance may have even occurred [40].

The effects of the reurrences of HBV (hepatitis B virus), HCV or combined hepatitis virus & human immunodeficiency virus (HIV) on liver immunology are another interesting question. Concerning HBV recurrence, individualized modern prophylaxis regimen, based on an integrated approach, modern molecules (entecavir, tenofovir) and risk-assessment (genetic variations in the HBV) allow eradicating recurrences [41]. For HCV, the first difficulty is the overlapping features of clinical and histo-pathological characteristics between ACR and HCV recurrence. Regev et al evaluated more than 100 LT biopsies conducted by five liver expert pathologists and concluded that the histopathological differentiation of HCV recurrence and ACR post-LT had relatively low inter-observer and intra-observer agreement rates, showing concerning low reliability. More recently authors used interesting strategy, proteomic analysis in plasma samples, distributions of different types of immune cells in tissue samples, microarray technology allowing studying molecular profiles and regulatory mechanism, miRNA, permitting important advances in the identification of several potentially useful biomarkers to differentiate ACR and HCV recurrence [43]. Despite these interesting advances, large studies correlating tissue analysis, serum molecular biomarkers and clinical observations are lacking.
The second difficulty is possible interaction between HCV recurrence and ACR. Many data confirm the pivotal role of T-cells, Tregs and DCs in the post-LT RHC setting[44]. So, the host immune response (mainly cellular mediated) appears to be crucial both in the control of HCV infection and in the genesis of rejection (paragraph I), and it is also strongly influenced by immunosuppressive treatment. No clear immunosuppressive strategy could be strongly recommended in HCV-positive recipients to prevent HCV recurrence. Nonetheless it seems that episodes of rejection and over-immunosuppression are more likely to enhance the risk of HCV recurrence through immunological mechanisms[45]. The past most common situation was an ACR episode (or a so-called post-LT increased in hepatic enzymes!), which was treated by increase in immunosuppressive therapy, with consecutive severe acute HCV recurrence.

Both complete prevention of rejection and optimization of immunosuppression represented the main goals towards reducing the rate of graft HCV re-infection. The present situation is much easier to manage with new hepatitis C treatments. These new interferon –free combinations can be used, either in pre transplant decompensated liver disease[46] or at every time after LT[47]. The eradication of the hepatitis C virus have favorable effect on the liver graft in double way: it permits to eliminate the differential diagnosis between ACR and HCV recurrence, and to increase IS therapy if ACR suspicion occurs, without the risk to trigger or worsen HCV recurrence concomitant with ACR.

III—ANTIBODY-MEDIATED REACTION AND LIVER TRANSPLANTATION

Antibody-mediated rejection (AMR) is an increasingly problematic entity in solid organ transplantation that leads to graft dysfunction and loss. AMR often occurs late (>1 year) after kidney transplantation with an incidence of approximately 30%, and was recently attributed to cause half of all renal allograft failures (48). In liver transplantation, AMR is usually diagnosed on the basis of the presence of all 4 of the following strict criteria: (1) DSAs (DSA, donor specific antigen) in serum; (2) histopathological evidence of diffuse microvascular injury/microvasculitis consistent with antibody-mediated injury; (3) diffuse C4d staining in the portal microvasculature with or without staining in the sinusoids or central veins in at least 1 sample; and (4) the exclusion of other causes of a similar type of injury[49].

One of the main reasons of renewed interest, especially in liver transplantation in which we considered this issue as minor or confined to ABO non compatible transplantation, is a recent technological advance that uses fast solid-phase assays and immobilized HLA to characterize HLA-specific antibodies[50]. The most widely adopted form of this is the single antigen bead assay (SABA), which although not perfect, has distinct advantages over the complement-dependent lymphocytotoxicity assay with which it is compared. SABAs identify antibody specificities to individual HLA alleles and can separate reactivity against different HLA loci, which is difficult to achieve using assays based on whole cells expressing multiple HLA in set combinations due to linkage disequilibrium. SABA also make sequential testing of multiple samples much easier to do.

Although the frequency and the impact of AMR in liver allograft is much lesser than in other organ transplantations, < 10% of sensitized recipients, emerging literature suggests that DSA can be associated with more rapidly progressive fibrosis, especially in hepatitis C-positive recipients with recurrent hepatitis, diminished long-term graft and patient survival[51]. An understanding of how antibodies cause graft injury and promote acute and chronic rejection is critical for the management of sensitized recipients and improvement of therapeutics. Studies have previously demonstrated that HLA I antibodies act as agonists, inducing molecular aggregation of HLA class I molecules to trigger intracellular signaling pathways that are critical in the regulation of cell survival, proliferation and migration in graft vascular cells[52]. Recent investigations have revealed additional functional responses of vascular cells to antibodies, including induction of inflammatory mediators by endothelial cells and recruitment of immune cells. Moreover, the relevance of complement fixing capacity of donor specific antibodies has received increasing attention[53] but some interesting studies counteract the relevance of sinusoidal deposition of complement 4d in liver allografts as proof of AMR[54]. We summarized in the table I the presumed role of different types of antibodies in liver AMR. However, the question of whether these antibodies are a cause or a consequence of the rejection process is unproven[55]. To confirm this point in liver transplantation, we should add to the definition[56] one last parameter: the therapeutic decrease in serum DSA improves liver function and decreases histologic injuries. Only, rare observation, in combined transplantation, demonstrated this last point[57]. In table 2, we summarize the possible role of DSA, according the different types of clinical rejection.

In state of the art, many meetings or consensus reports[48,50,51] insisted on different technical (especially, importance to detect HLA antibodies with solid phase immunoassay test) and no technical followed points: (1) The liver allograft may be partially resistant to antibody mediated damage; however, high-level DSA antibody may be associated with inferior outcomes and should be considered as a risk factor for graft dysfunction; (2) Pretransplantation screening for HLA antibodies is recommended in liver retransplant recipients and cross-match should be performed in sensitized recipients[58]; (3) Donor blood and tissue should be collected and stored in liver transplantation to a posteriori research DSA or complement deposition; (4) In sensitized recipients of combined liver-kidney transplantation, the liver may not confer full protection for preventing AMR in the kidney and should be included in risk assessment (confer next chapter).

IV—THE LIVER AS IMMUNOREGULATORY ORGAN IN COMBINED ORGAN TRANSPLANTATION

Liver has been recognized as an immunoregulatory solid organ in the field of transplantation for a long time[59]. Some strains of pigs can tolerate allogenic MHC mismatched liver without

| Table 1 | Different types of antibody related to liver graft injury and their relevance. |
|---------|-------------------------------------------------|
| Auto versus allo | ++ | ++ |
| HLA vs non HLA | ++ | +? |
| De novo vs predefined | + | ++ |
| Class I vs Class II | ++ | (++ preformed) |
| IgG-complement fixing vs not (IgG1,3 vs other Ig) | ++ | ? |
| Flow cytometry crossmatches T vs B (MFI) | ++ if > 5000 | ? if < 3000 |

| Table 2 | Antibody-mediated rejection (AMR) and type of allograft rejection in liver or liver kidney transplantation. DSA: donor specific antigen. |
|---------|-------------------------------------------------|
| Circumstances | | |
| Hyperacute | ABO incompatible, very seldom |
| Acute | <1% recipients, in patients with predefined DSA |
| Chronic | Possible, in some patients after complete weaning immunosuppression (de novo DSA) |
| C4d-negative | Probably, under development |
immunosuppressive therapy (IS). Moreover, in certain strains of rodents, instead of sensitizing the recipient, the enduring liver graft induce a state of donor-specific unresponsiveness in which subsequent grafts (skin or other organs) are accepted permanently without IS\(^\text{60}\). Tolerant recipients of liver allografts have a high blood concentration of Class I MHC molecules of donor type and injections of serum and lymph from such animals given daily successfully prolonged the survival of skin or heart grafts in naïve animals of appropriate strains\(^\text{61,62}\).

In human, many studies\(^\text{60}\) reported that complete withdrawal of IS maintenance therapy is possible in a little percentage of well-selected recipients < 20%, with non-viral, nonimmune disease. The protective immunologic role of the liver was confirmed in combined heart-liver, intestine-liver, pancreas liver and kidney-liver transplantation\(^\text{641}\). However, in this large cohort provided by the United Network of Organ Sharing (UNOS), the reduction in allograft rejection was also observed in patients co-transplanted with two organs not including a liver: rejection-free survival of patients simultaneously kidney-heart or kidney-liver transplanted (from the same donor) was higher than the rejection-free survival of the patients transplanted with heart or liver alone; rejection-free survival of patients simultaneously transplanted with two lungs or two kidneys (from the same donor) was also better than patients transplanted with one single organ. Authors focused their explanations on the importance of high antigen load brought by simultaneous combined transplantation. In a more recent study\(^\text{622}\), also provided by the UNOS and interesting specifically in mortality and rejection, authors found that the mortality at 1, 3 or 5 years of patients undergoing combined heart-liver transplantation (respectively 84, 74 and 72%) was not different from patients undergoing isolated liver transplantation (respectively 85, 78 and 72%) and significantly better than patients undergoing isolated heart transplantation (respectively 83, 73 and 65%). Moreover, the incidence of acute cardiac rejection within 1 year for patients with cardiac graft survival greater than 1 year was 9% for those undergoing simultaneous transplantation compared with 24% for those undergoing heart transplantation alone \((P=0.002)\). The incidence of acute liver rejection within 1 year for patients with liver graft survival greater than 1 year was 5% for those undergoing simultaneous transplantation versus 12% for those undergoing liver transplantation alone \((P=0.060)\).

Concerning liver and kidney transplantation, Simpson et al\(^\text{623}\) showed that rejection-free graft survival of kidney after liver transplantation (KALT) patients was lesser than combined liver kidney transplantation (CLKT) patients. This difference was even more statistically significant when they considered patients with HLA mismatched grafts. Another study from the UNOS database\(^\text{624}\) showed that there was no difference in recipient mortality between liver transplantation alone (LTA) and simultaneous liver-kidney transplantation (SLK). Recipient and graft survival in SLK was higher compared to both kidneys after liver transplantation (KALT) and liver transplantation after kidney (LAKT).

All these clinical studies have limitations: retrospective, comparison of medical selected patients and treatment and not randomized. However, all observed a better rejection free survival and a better “no liver” graft survival in patients simultaneously heart-liver or kidney-liver transplanted and seem to confirm experimental data. Many mechanisms have been reported in the past. Concerning acute cellular rejection, the main invoked mechanisms are: (1) Large cellular compartment with hematopoietic regulators (γδT lymphocytes, NK and NKT cells, DC) and dilutional mass effect; (2) Mixed hematopoietic microchimerism; (3) Regulatory proteins/cytokines secretion, for example HLA-G.

The first mechanism observed is that the liver is an especially large alloantigen transfer in recipient. The increased mass of tissue transplanted may exhaust the recipients’ immune response. One mechanism for this exhaustion could be that there is a limited clone size of graft-reactive T cells which are unable to establish “critical mass”. This is analogous to a nuclear chain reaction, where fission does not occur until there is sufficient density of free neutrons\(^\text{625}\). Many experimental studies confirm this non-specific liver effect. Recently authors\(^\text{666}\) studied in mice if recombinant adenoassociated virus (rAAV) mediated expression of donor major histocompatibility complex in recipient livers could induce tolerance to donor-strain grafts. High-level expression of donor major histocompatibility complex in recipient livers promotes tolerance to skin allograft while low level accelerated the rejection. This effect of high dose was already described if donor MHC gene were transferred in hematopoietic cells before cardiac graft of the same donor\(^\text{67,68}\) or into skeletal muscle before skin graft\(^\text{69}\).

The second main mechanism is microchimerism\(^\text{69}\). Microchimerism arises as a result of migration of passenger leukocytes from a transplanted allograft into an unconditioned recipient and donor pluripotent HSCs do not engraft, but alternatively hematopoietic-derived cells from the donor organ are produced and migrate systemically. Although this phenomenon probably does exist with all solid organ transplantsations, but it is higher\(^\text{70}\) longer\(^\text{71}\) and more frequent in liver transplantation\(^\text{72}\). Stable levels of donor chimerism may be a marker of transplantation tolerance, and may help to tailor immunosuppressive treatment in liver transplantation\(^\text{73}\). As low as 1% donor chimerism is sufficient to induce tolerance to donor-specific organs, cells, and tissues\(^\text{74}\). This level may occur more often when liver is present in combined solid organ transplantation. Already, Kawai et al\(^\text{75}\) achieved the establishment of chimerism in primates, without bone marrow transplantation, in a non myeloablative preconditioning protocol, opening promising possibility in human.

Concerning AMR, many reasons have been given to explain why liver can withstand and protect other organs from DSA\(^\text{76}\): (1) Dilution: Increase area of distribution: HLA Class I antigens are present on all liver cells, especially endothelial cells and liver capillary area is 100 times that of the heart or the kidney. HLA Class II is constitutively expressed only on dendritic cells and on other cells in the setting of inflammation. Moreover, platelets and complement factors are decreased in cirrhotic patients; (2) Inactivation: Role of Kupffer cells, able to removal immune complexes, activated complement and platelet aggregates, formed during AMR\(^\text{77}\); (3) Characteristics of HLA alloantibodies: class and subclass, Fe binding, complement fixation and antigen affinity, likely affect their injurious potential.

A last propriety of the liver to explain it withstand to both humoral and cellular rejection is its exceptional regenerative capacity. While other solid organs presenting acute rejection, are rapidly suffering from organ dysfunction, liver is able to function unless a very severe rejection occurs. Moreover, rapid treatment of the rejection will avoid anatomic and functional aftereffects.

**CONCLUSION**

In conclusion, recent researches permitted a progression in the understanding of the better tolerance of the liver than other solid organs. The three major mechanisms, at this date, are microchimerism and tremendous antigen load, overlapping the
lymphocytes suppressing LT response, in cellular rejection and trapping of different products like, antibodies, immune complex and platelet aggregates in AMR. This better comprehension could offer perspectives not only in liver transplantation and combined liver-other solid organ transplantation but also in the other solid organs transplantation.

**CONFLICT OF INTERESTS**

The authors declare that they do not have conflict of interests.

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Peer reviewer: