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**Revisiting liver’s role in transplant alloimmunity**

Abrol N *et al*. Liver’s role in alloimmunity

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**Abstract**

The transplanted liver can modulate the recipient immune system to induce tolerance after transplantation. This phenomenon was observed nearly five decades ago. Subsequently, liver’s role in multi-visceral transplantation was recognized, as it has a protective role in preventing rejection of simultaneously transplanted kidney, heart, or other solid organs. The liver has a unique architecture and is home to many cells involved in the immunity and inflammation. After transplantation, these cells can migrate from the liver into the recipient. Early studies pointed out towards chimerism as an important mechanism by which the liver modulates human immune system. Recent studies on human T-cell subtypes, cytokine expression, and gene expression in the allograft have expanded our knowledge on the potential mechanisms underlying immunomodulation. In this article, we discuss the privileged state of liver transplantation compared to other solid organ transplantation, the liver allograft’s role in multivisceral transplantation, various cells in the liver involved in immune responses, and the potential mechanisms underlying immunomodulation of host alloresponses.

**Key words:**Liver transplantation; Alloimmunity; Liver-kidney transplant; Tolerance; Rejection

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**Core tip:** The liver does not only protect itself from the host alloimmune responses, but also modulates alloimmune responses to simultaneously transplanted other solid organs like heart or kidney. The titer of donor specific alloantibodies decreases after liver transplantation, making transplantation of other solid organs possible even in the highly sensitized high risk patients. The immune cells from the liver allograft cross-talk with recipient immune cells and modulate the immune system towards tolerance. The cross-talk between these cells suppress the genes involved in alloimmunity and upregulate the genes involved in tissue repair and metabolism.

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**INTRODUCTION**

The liver has baffled researchers for decades because of its complex set of functions and unique architecture. From a metabolic and anatomic standpoint, it has a dual blood supply with the portal vein carrying blood from the gastrointestinal tract and the hepatic artery carrying systemic blood. From an immunological standpoint, the liver is home to many cells of the lymphoid system. Together, the liver’s unique architecture and resident immune cells, allow it to play a key role in transplant alloimmunity. It is well recognized that the liver is an immunologically privileged organ, compared to other organs that are commonly transplanted. The liver allograft not only protects itself from the host immune system, but this protection also extends to other simultaneously transplanted solid organs from the same donor. Many researchers have investigated potential mechanisms of this donor specific hypo-responsiveness. Recent studies on the host T-cell subtypes and gene expression in the allograft after multi-visceral transplants that include the liver, have expanded our knowledge on the liver’s role in transplant immunity[[1-3](#_ENREF_1)]. In this article, we revisit the liver allograft’s role in modulating host alloimmunity with special emphasis on combined organ transplants.

**LITERATURE SEARCH**

For purpose of this review, the Embase and Ovid MEDLINE databases were searched from year 2000 till January 2019 using keywords “Liver transplant\*” and “Alloimmunity”. The search included Epub ahead of print, in process, and other non-indexed citations. After removing duplicate publications, 242 studies were finally reviewed by title and abstract for selecting full text articles for current review. The studies describing liver based modulation of cells of the immune system in solid organ solitary liver or multi-visceral transplantation were selected for review.

**LIVER IS AN IMMUNOLOGICALLY PRIVILEGED ORGAN**

The liver’s immune-privileged status was first recognized in the porcine liver transplantation model[[4](#_ENREF_4)]. As early as 1965, it was observed that pigs undergoing liver transplantation survived for prolonged periods with limited immunosuppression, whereas other organs, including skin, heart, and kidneys were quickly rejected[[4](#_ENREF_4)]. This phenomenon has since been observed also in other animal models[[5](#_ENREF_5)].

The first reports of tolerance in human liver transplantation came from Thomas Starzl and the Pittsburgh group[[6](#_ENREF_6),[7](#_ENREF_7)]. Their early experience showed that 27% of liver transplant recipients could be weaned from all immunosuppression[[8](#_ENREF_8)]. Subsequently, many other groups tried immunosuppression weaning in patients with stable liver function[[6](#_ENREF_6),[9-11](#_ENREF_9)]. In a pilot study, 60% of carefully selected pediatric liver transplant recipients could be successfully weaned off immunosuppression[[10](#_ENREF_10)]. However, this approach was associated with increased rejection in other recipients[[9-11](#_ENREF_9)]. Nevertheless, most liver transplant patients require less maintenance immunosuppression than recipient of other solid organs. Likewise, induction immunosuppression, other than steroids, is rarely needed in liver transplantation.

***Antibody-mediated rejection***

Compared to other solid organ transplants, liver transplant recipients have fewer episodes of antibody-mediated rejection (AMR). While donor specific alloantibodies can cause antibody-mediated hyperacute or acute rejection in other solid organs, their role in liver transplantation remains unclear[[12-17](#_ENREF_12)]. Liver transplantation is often performed without a prospective cross-match and outcomes do not appear to be related to pre-transplant positive cross-match[[18](#_ENREF_18),[19](#_ENREF_19)]. The majority of recipients with preformed donor specific antibodies (DSA) show decline in their DSA levels after liver transplantation (Figure 1). In patients with persistent post-transplant DSA, there does not appear to be any negative impact on allograft survival within the first year after transplantation[[20](#_ENREF_20)]. The observed decline in DSA post-transplant appears to be linked to the overall health of the liver allograft, as persistence of DSA or development of de novo DSA are observed more commonly in patients with allograft fibrosis or recurrent disease. However, this protection is not complete and there is evidence of complement fixation in recipients with persistent DSA in protocol liver biopsies[[20](#_ENREF_20)]. In patients with de novo DSA against class II human leukocyte antigens (HLA), the overall survival is inferior to those with no DSA[[21](#_ENREF_21)]. Importantly, de novo DSA are not uncommon in patients whom immunosuppression withdrawal attempted, suggesting that most liver transplant patients require some, albeit minimal immunosuppression to counter the host alloimmune responses[[22](#_ENREF_22)].

Several factors are felt to play a role in the liver’s resistance to antibody-mediated hyperacute rejection. These factors include the liver’s dual blood supply, its fenestrated sinusoidal complex, secretion of soluble major histocompatibility complex antigens, and its ability to absorb antibody (Figure 2). In contrast to other solid organs, the microvascular network of the liver is sinusoidal and lined by fenestrated endothelium with a scant underlying basement membrane (Figure 2, g)[[23](#_ENREF_23)]. This sinusoidal network is in contrast to other organs that not only have a single afferent blood supply, but also have standard capillary microvasculature that results in ischemia when occluded by complement activated immune complexes. In the liver, only the biliary system is truly dependent on capillary microvasculature. This histological variation may result in a more limited, biliary-specific, form of injury in liver transplantation compared to other solid organs[[24](#_ENREF_24)].

***T cell-mediated rejection (TCMR)***

Unlike other solid organs, cellular (T-cell mediated) rejection (TCMR) in liver transplantation follows a bimodal pattern of distribution with the majority of cellular rejections occurring very early (< 6 wk) post-transplant[[25](#_ENREF_25)]. When early cellular rejection episodes occur in liver transplant patients, these episodes require much less immunosuppression compared to TCMR in heart, pancreas, lungs, or kidney. Similarly, unlike other solid organs, these early episodes of T-cell mediated rejection do not appear to have any long-term impact of patient or allograft survival[[25](#_ENREF_25)]. In liver transplantation, TCMR can largely be treated by increasing the dose of immunosuppression or by pulse steroids without requiring lymphocyte depleting antibody-based treatment.

**LIVER ROLE IN MULTI-VISCERAL TRANSPLANTATION**

Liver-induced immunological tolerance to other allografts was first recognized in pigs, when liver allografts were noted to prevent rapid rejection of skin, kidney, and heart from the same donor[[4](#_ENREF_4)]. This phenomenon was observed to be true for both orthotopic and auxiliary liver transplants[[4](#_ENREF_4)]. Since these initial animal models, the same observation has been made in human multivisceral transplantations[[1](#_ENREF_1),[2](#_ENREF_2),[26-32](#_ENREF_26)]. Patients who undergo a combined liver-kidney transplantation (LKT) experience lower number of kidney TCMR episodes compared to a matched group of solitary kidney transplant alone (KTA) recipients (4.2% *vs* 32.6%)[[33](#_ENREF_33)]. The protective effect of the liver allograft on simultaneously transplanted kidney persists for a long time[[2](#_ENREF_2),[34](#_ENREF_34)]. In a study comparing kidney transplantation after liver or heart/lung transplantation, recipients who previously had liver transplantation had fewer episodes of TCMR in the kidneys (20% *vs* 36%)[[34](#_ENREF_34)]. In addition, the observed rejection episodes were less severe (all rejection episodes were grade IA/IB), and grade II or grade III rejections were seen only after heart/lung transplantation (0% *vs* 16%)[[34](#_ENREF_34)]. Similar protective effects against AMR have been reported by several groups. In Sweden, auxiliary liver transplantation was done in a group of highly sensitized kidney patients who were otherwise deemed too risky to transplant, to facilitate kidney transplantation, with partial success[[35](#_ENREF_35)]. We have also demonstrated protection of the heart allograft from AMR in highly sensitized patients by initial liver transplantation from the same donor[[27](#_ENREF_27)]. All patients in this cohort had pre-existing DSA and positive cross-match. There was immediate decrease in DSA and stable cardiac and liver allograft function at mean follow up of nearly 2 years[[27](#_ENREF_27)]. Furthermore, cardiac allograft vasculopathy (assessed with 3D volumetric intravascular ultrasound) and the overall plaque volume was lower, and the plaque progression rate slower in the cardiac allografts of patients who underwent combined liver-heart transplantation[[36](#_ENREF_36)]. In a series of 13 combined liver-lung transplants, only 3 patients experienced early rejection that was successfully treated with methylprednisolone[[28](#_ENREF_28)], and this rate is much lower than that seen after solitary lung transplantation[[28](#_ENREF_28)]. Similarly, liver was protective in combined liver-intestine transplantation[[26](#_ENREF_26),[37](#_ENREF_37)].

**OVERVIEW OF ALLOIMMUNITY**

Detailed discussion of alloimmunity and downstream pathways after antigen presentation is beyond the scope of this article. Briefly, alloantigens from the transplanted organ are recognized by the host lymphocytes in the secondary lymphoid organs. Dendritic cells (DC), macrophages, B lymphocytes, and endothelial cells (EC) can play the role of antigen-presenting cells (APC) under various circumstances. Allorecognition occurs via three main pathways: (1) the direct pathway where T cell receptors on host T cells directly interact with the HLA molecules on the surface of donor APC; (2) the indirect pathway where host APC process donor peptides (mostly derived from donor HLA) and present to host T cells; and (3) the semi-direct pathway that involves membrane exchange between donor and host cells or extracellular vesicles[[38](#_ENREF_38),[39](#_ENREF_39)]. T cell activation after antigen presentation (Signal 1) requires two additional signals after the cognate HLA: T cell receptor interaction occurs; binding of costimulatory molecules on T cells (CD40, CD28) with corresponding ligands on the APCs (CD40L, CD80, CD86) (Signal 2), and the presence of T cell stimulatory cytokines in the microenvironment (Signal 3) resulting in T cell proliferation (Figure 3).

**LIVER AS A LYMPHOID AND IMMUNE-REGULATORY ORGAN**

Liver architecture is uniquely adapted to provide immunomodulation after exposure to the foreign antigens from the gastrointestinal tract. The liver receives a dual blood supply from the high-pressure systemic and the low-pressure portal circulation. These two circulations meet in the hepatic sinusoids resulting in low oxygen saturation, low pressure, and irregular flow facilitating interaction between antigens, T cells and other resident immune cells[[40](#_ENREF_40)]. Although cell migration occurs in all types of solid organ transplants, the large population of migratory cells in liver allografts may explain the privileged tolerogenicity of the liver compared to other organs (Figure 2)[[41](#_ENREF_41)]. The hepatocytes are arranged as sheets around the sinusoids. The liver is constantly exposed to microbial antigens carried through the portal circulation. In order to avoid immune activation in response to microbial antigens, liver has developed many molecular modifications. This is evident from high levels of lipopolysaccharide in the portal blood when none is detected in the systemic circulation under normal conditions[[42](#_ENREF_42)]. Therefore, there is evolutionary advantage to the immunomodulatory role of liver parenchyma. In fact, the liver has been described as a “lymphoid”, ‘immunoregulatory” and “immunomodulatory” organ with various cells playing active role in supporting this function[[13](#_ENREF_13),[42](#_ENREF_42),40].

Immune cells of both lymphoid and myeloid lineage line the thin walled sinusoids, mostly in the space of Disse (Figure 2, e)[[42](#_ENREF_42)]. These cells include Kupffer cells (KC), DC, T cells, B cells, natural killer (NK) cells, natural killer T (NKT) cells, hepatic stellate cells (HSC), and hematopoietic stem cells[[42](#_ENREF_42),[43](#_ENREF_43)]. The phenotype of hepatic T cells also differs considerably from that observed in the periphery as reflected by a higher ratio (3.5:1 *vs* 1:2) of CD8+ *vs* CD4+ cells[[44](#_ENREF_44)]. The unique architecture of liver sinusoids (low pressure, fenestrated system, expression of adhesion molecules) allows direct contact of circulating T cells with these cells. These alloantigen recognizing T cells are exposed to IL-10, PD-L1, and lack of co-stimulation leading to their destruction in the liver[[42](#_ENREF_42)].

***Endothelial cells***

The sinusoidal EC comprise of 50% of non-parenchymal liver cells (Figure 2, g)[[40](#_ENREF_40),[43](#_ENREF_43)]. The sinusoidal EC uniquely lack a basement membrane, are fenestrated, and express scavenger receptors that remove circulatory antigens[[42](#_ENREF_42)]. ECs also express class I and II HLA and costimulatory molecules, making them potent APCs. However, their main role seems to be induction of tolerance because they respond to antigen stimulation by IL-10 secretion[[42](#_ENREF_42)]. ECs increase their expression of FasL upon exposure to antigen and induce apoptosis of activated CD4+ T cells[[45](#_ENREF_45)]. ECs also induce apoptosis of reactive CD8+ T cell via a pro-apoptotic Bcl-2 family member Bim[[46](#_ENREF_46)].

***Dendritic cells***

The DC are professional APCs derived from bone marrow (Figure 2, f). The liver contains two types of DCs-plasmacytoid (pDC) and myeloid (mDC)[[42](#_ENREF_42)]. The observed frequency of pDC in the liver is more than that in the lymph nodes[[42](#_ENREF_42)]. The liver contains these cells in immature form. Under normal circumstances, these cells have low expression of the costimulatory molecule CD80[[47](#_ENREF_47)]. Liver pDC play an important role in innate immunity, as they can produce and secrete IFN-γ[[42](#_ENREF_42)]. On the other hand, pDC can express PD-L1 on their cell surface, and increased expression of PD-L1 on pDC in tolerant liver transplant patients has been correlated with elevated Tregs[[48](#_ENREF_48)]. Liver mDC, unlike their counterparts isolated from other organs, appear to have a more inherent tolerant phenotype. For example, under normal circumstances, liver derived mDCs secrete IL-10 and mediate differentiation of T cells into Tregs[[49](#_ENREF_49)]. MDC interactions with HSC may play a role in downregulating immune response[[50](#_ENREF_50)]. HSC regulate mDC function by inducing signal transducer and activator of transcription 3 and upregulation of indolamine 2,3-dioxygenase (IDO)[[50](#_ENREF_50)]. Therefore, mDC primed by HSC have impaired ability to induce allogenic T cell response[[50](#_ENREF_50)].

***Kupffer cells***

The KC are macrophages present in the intra-sinusoidal space and comprise of 15% of all liver cells and 20% of non-parenchymal liver cells (Figure 2, d)[[40](#_ENREF_40),[43](#_ENREF_43)]. Their main role is phagocytosis and cytokine secretion[[40](#_ENREF_40)]. They also express HLA and costimulatory molecules, therefore they can present antigen to T cells[[42](#_ENREF_42)]. However, compared to DC, the expression of HLA and costimulatory molecules is low[[40](#_ENREF_40)]. KC secrete IL-10 and downregulate secretion of proinflammatory cytokines IL-6 and TNF-γafter exposure to lipopolysaccharide[[51](#_ENREF_51)]. KC have also been found to secrete prostaglandin E2 (PGE2) and 15-deoxy-delta 12,14-PGJ2 (15d-PGJ2)[[52](#_ENREF_52)]. PGE2 and 15d-PGJ2 inhibit activation of CD4+ T cells[[52](#_ENREF_52)]. KC can also stimulate Tregs to secrete IL-10[[53](#_ENREF_53)].

***Natural killer cells***

The liver contains a high percentage of NK cells (50% of liver lymphocytes) compared to peripheral blood (Figure 2, b)[[40](#_ENREF_40),[42](#_ENREF_42)]. Two type of NK cells exist in the liver: CD3-CD56dimCD16+CD27- (cytotoxic phenotype) and CD3-CD56brightCD16- CD27+ (cytokine secreting phenotype)[[40](#_ENREF_40)]. Their role in alloimmunity and rejection appears to be influenced by their origin, such that the NK cells derived from the recipient are involved in rejection while donor-derived NK cells induce tolerance[[54](#_ENREF_54)]. NK cells have been found to overexpress certain genes in tolerant liver transplant recipients signifying their important role in tolerance induction[[55](#_ENREF_55)]. This is consistent with the upregulation of NK cell transcripts in tolerant liver transplant patients[[56](#_ENREF_56)]. In addition, NKT cells, which express markers of NK cells along with T cell receptor Vα chain, appear to have a role in liver-induced tolerance, as tolerance is reversed in mice deficient in Vα14 NKT cells [[57](#_ENREF_57)].

***Hepatic stellate cells***

HSC are located in the subendothelial space and comprise 10% of the liver cells[[43](#_ENREF_43)]. Known also as Ito cells, HSC store vitamin A and are involved in various fibrotic processes[[43](#_ENREF_43),[58](#_ENREF_58)]. They express HLA class I, HLA class II and can activate T cells[[58](#_ENREF_58)]. However, they also express PD-L1 that can lead to tolerance by inactivating activated T cells[[42](#_ENREF_42),[59](#_ENREF_59),[60](#_ENREF_60)]. There is evidence that both parenchymal and non-parenchymal cells in the liver cause activation followed by apoptosis of the T cells in the liver allograft as well as in vitro[[42](#_ENREF_42),[61](#_ENREF_61)]. Allogenic HSC can migrate to lymph nodes and induce expression of Tregs[[62](#_ENREF_62)].

***Mesenchymal stromal cells***

The mesenchymal stromal cells (MSC) have also been localized in the liver[[63](#_ENREF_63)]. MSC were first described by Friedenstein et al as fibroblast like colonies in the bone marrow cultures[[64](#_ENREF_64)]. Subsequently, MSC have been identified in various organs like bone marrow, adipose tissues, as well as the liver. These cells are characterized by their ability of trilineage differentiation, plastic adherence, and expression of certain markers on their surface[[65](#_ENREF_65)]. Though liver derived MSC have not been well characterized yet, extensive research on the bone marrow and adipose tissue-derived MSC shows that MSC have the ability to modulate every cell of the immune system including macrophages, DC, NK cells, B cells, and T cells[[66](#_ENREF_66)]. Interaction of APC with MSC program the former towards tolerant phenotype as evident from increased IL-10 secretion[[66](#_ENREF_66)]. MSC may modulate these responses by IDO[[67](#_ENREF_67)]. Liver MSC appear to be more potent than the bone marrow- and adipose-derived MSC in their capacity to modulate alloimmune T cell responses, at least in vitro[[68](#_ENREF_68)].

**LIVER’S ROLE IN TOLERANCE DEVELOPMENT AND THE UNDERLYING MECHANISMS**

“True tolerance” is long-term acceptance of the allograft in the absence of any immunosuppression and without evidence of any DSA or signs of lymphocyte activation on biopsy[[38](#_ENREF_38)]. True tolerance in human beings is a rare phenomenon. More common scenario in clinical transplantation is the stable graft function for at least 1 year in the absence of immunosuppression (“operational tolerance”) or with minimal immunosuppression (“prope tolerance”)[[38](#_ENREF_38),[69-71](#_ENREF_69)]. Nearly 25% of adult and 60% of pediatric liver transplantation recipients can achieve operational tolerance[[55](#_ENREF_55),[72](#_ENREF_72)]. While different tolerance mechanisms have been demonstrated in animal models and limited clinical studies, how exactly the liver allograft dampens the host alloimmune responses remains unknown.

***Chimerism***

The liver contains a population of hematopoietic stem cells[[73](#_ENREF_73)]. In fact, liver transplants can behave like bone marrow transplants and rare cases of graft versus host disease have been described after liver transplantation[[74-76](#_ENREF_74)]. In the earliest era of clinical liver transplantation, presence of donor cells in the recipient circulation was observed (chimerism), and through circumstantial evidence, chimerism was thought to lead to tolerance, however despite these early clues, these findings were not further investigated until the late 1980s. Chimerism was first demonstrated in 1968, with karyotyping studies of male donor livers that had been transplanted into female recipients[[77](#_ENREF_77)]. It was observed that the majority of the allograft retained its donor specificity, but the bone marrow derived passenger leukocytes, including KCs, were largely replaced with recipient female cells within 100 d. There was also evidence of adoptive immunity, with demonstration of new acquired immunoglobulin types of donor specificity and donor derived anti-erythrocyte isoagglutinin-associated hemolysis. Despite these subtle clues, it would not be until almost two decades later, that the conviction that donor cells were wholly eliminated by the immune system would be challenged[[7](#_ENREF_7)]. In 1992, decisive steps were taken to search for donor leukocytes in the blood and tissue of thirty human recipients of successful liver transplants up to 29 years prior. Female recipients from male donor were found to have microchimerism in their allografts and extrahepatic tissues 10 to 19 years post-transplant[[7](#_ENREF_7)].

The early alloresponse after liver transplantation is characterized by recruitment of CD4 T cells to the allograft and by their proliferation and IFN-γ production[[78](#_ENREF_78)]. However, later there is selective reduction of T cells in the recipient[[78](#_ENREF_78)]. At the same time the donor hematopoietic and T cells migrate from the allograft into the recipient. These donor-derived cells may survive in the recipient for a prolonged period of time and lead to chimerism observed after liver transplantation[[79](#_ENREF_79)]. If donor hematopoietic cells constitute more than 1% of the recipient tissue, this is termed macrochimerism, and if they are < 1%, microchimerism[[79](#_ENREF_79)]. In one study, all patients showed chimerism initially after liver transplantation, however chimerism decreased to variable degree in the first year[[79](#_ENREF_79)]. The rejection episodes in this study correlated with the lower degree of chimerism[[79](#_ENREF_79)]. The patients with high degree of chimerism had measurable in vitro alloreactive response after one year suggesting that chimerism did not lead to complete depletion of cytotoxic T cells[[79](#_ENREF_79)]. The passenger cells in the liver may be playing role in tolerance induction[[6](#_ENREF_6),[7](#_ENREF_7),[79](#_ENREF_79),[80](#_ENREF_80)], as strategies to reduce number of these passenger cells before transplantation prevent tolerance induction in experimental models[[81](#_ENREF_81)].

***T cell deletion***

The unique architecture of the liver and the cross-talk between alloreactive T cells and liver inhabitant cells may play a significant role in tolerance induction by destroying host T cells[[42](#_ENREF_42),[45](#_ENREF_45),[61](#_ENREF_61)]. The fenestrated endothelium of hepatic sinusoids facilitates direct contact between T cells and parenchymal cells leading to T cell deletion[[74](#_ENREF_74)]. There is distinct expression of genes for T cell recruiting cytokines after LKT in tolerant patients[[82](#_ENREF_82)]. This study found large number of CD3+ T cells and macrophages in the liver allograft but only few in the simultaneously implanted kidney allograft[[82](#_ENREF_82)]. It seems that increased expression of chemokines in the liver attracts alloreactive T cells that are subsequently destroyed by coming in contact with various liver cells inherently programmed towards tolerance induction. Another study found donor specific hypo-responsiveness, down regulation of T helper type I cytokine (IFN-γ) and no change in T helper type 2 cytokine (IL10) in the in vitro mixed lymphocyte reaction in recipients who achieved operational tolerance[[83](#_ENREF_83)]. Similar cytokine pattern was found in the allograft on real time reverse transcriptase polymerase reaction (RT-PCR)[[83](#_ENREF_83)]. Animal experiments have shown that T cells activated in the lymph nodes are capable of mediating immune response but T cells activated in the liver are short lived, defective, and are not able to mount immune response[[84](#_ENREF_84)].

***Peripheral Tregs***

An alternative model is development of regulatory T cells (Treg) that actively regulate alloreactive T cells[[62](#_ENREF_62)]. The liver cells secrete cytokines after antigen presentation that differentiates host T cells into regulatory phenotype[[48](#_ENREF_48),[49](#_ENREF_49),[53](#_ENREF_53),[62](#_ENREF_62)].

***DSA neutralization***

Under normal circumstances, liver has strong expression of class I HLA, secretes class I HLA antigens, and has weak class II expression[[13](#_ENREF_13),[85](#_ENREF_85)]. This might absorb anti HLA type I DSA leading to lower risk of antibody mediated rejection. We studied DSA levels in the serum of liver transplant recipients who did not receive any antibody targeting induction[[20](#_ENREF_20)]. Nearly 20% recipients had pre-formed DSA that markedly decreased in all but three recipients 7 days after transplantation[[20](#_ENREF_20)]. In rare instance, when DSA persists, there is compliment activation and C4d deposition in the liver[[20](#_ENREF_20)]. One year follow up showed stable function despite antibody mediated complement activation in patients with persistent DSA[[20](#_ENREF_20)]. The unique architecture of hepatic sinusoids–fenestrated endothelium, lack of basement membrane, wider lumen–may confer resistance to complement activity. When endothelium injury does occur, it is seen in microvasculature but not in the sinusoids[[86](#_ENREF_86)]. This might be a reason for high susceptibility of peribiliary plexus to immunological or ischemic damage that is derived from hepatic artery[[43](#_ENREF_43),[86](#_ENREF_86)]. DSA level in the recipient seem to be the net result of two opposing factors–host memory cells mounting immune attack and liver mediated neutralization of alloantibodies. Though protection against de novo class II DSA is less, incidence of de novo class II DSA is less in liver transplantation compared to kidney transplantation[[13](#_ENREF_13)].

At Mayo Clinic, we perform nearly 400 solid organ transplants in a year and many are combined liver-kidney transplants. Our group has investigated the liver’s role in modulating host alloimmune response in these combined transplant recipients. Our program also employs protocol kidney biopsies to investigate the extent of subclinical and chronic alloreactivity. In our work, we have found that liver allografts from LKT protect the kidney from hyperacute/acute antibody mediated rejection [odds ratio 0.11, 95% confidence interval (CI) 0.03-0.32] and acute cellular rejection (odds ratio 0.13; 95%CI 0.06-0.27). Moreover, in assessing variables, the presence of a functioning liver allograft was the most predictive factor for protecting kidney allograft from the chronic injury (odds ratio 0.22, 95%CI 0.06-0.59)[[1](#_ENREF_1)]. The KTA patients with positive DSA had 44% decline in GFR by 5 years while LKT patients with positive DSA had stable GFR[[1](#_ENREF_1)]. The LKT recipients have lower frequency of circulating CD8+, activated CD4+, and effector memory T cells, compared to KTA recipients[[2](#_ENREF_2)]. Moreover, surviving T cells in LKT patients had lower proliferative response to the donor cells (11.9% *vs* 42.9%), although the response to third party was not altered[[2](#_ENREF_2),[87](#_ENREF_87)]. This donor specific hypo-responsiveness persisted after first year of transplant[[2](#_ENREF_2)]. We further compared molecular changes in the kidney allograft after LKT and KTA by doing RT-PCR on the protocol kidney biopsies[[3](#_ENREF_3)]. We found that mechanisms underlying liver’s protective role do not only operate inside the liver but extend to the kidney as there were distinct gene expressions seen on the RT-PCR[[3](#_ENREF_3)]. The kidneys in LKT showed markedly increased expression of genes associated with tissue integrity/metabolism, even in cross-match positive transplants[[3](#_ENREF_3)]. We hypothesize that liver inhabitant cells migrate into the circulation after liver transplant transplantation to home at the site of inflammation in the second co-transplanted solid organ and modulate host immune cells. While the key cell type is unknown yet, this hypothesis is supported by our work and previously published studies[[2](#_ENREF_2),[3](#_ENREF_3),[7](#_ENREF_7),[80](#_ENREF_80),[88](#_ENREF_88)].

Tolerance can be conceptualized as a state of fine balance between two opposing forces: host immune system and liver mediated immune-regulation. This balance can be tilted towards rejection by stimulation of immune system by tissue damage[[89](#_ENREF_89)]. After exposure to endotoxins from infectious agents or Toll Like Receptors resulting from ischemia reperfusion injury, there is upregulation of class II HLA on hepatic EC[[13](#_ENREF_13)]. While operational tolerance has been demonstrated in few patients, other liver transplant patients require minimal immunosuppression to counter the effect of host alloimmune response.

**CONCLUSION**

The liver microenvironment is inherently programmed towards induction of tolerance as a result of evolution to avoid immune activation on exposure to the gut delivered antigens. This has important implications in the alloimmunity in the context of liver transplantation alone or in multi-visceral transplants. Liver induced protection against host immune system is likely the result of multitude of effects including microchimerism, deficient antigen presentation due to lack of costimulation and expression of inhibitory molecules, deletion of activated recipient T cells in the liver, large antigen load in liver and active secretion of HLA molecules neutralizing alloantibody, and generation of Tregs in peripheral lymph nodes. Liver parenchymal as well as non-parenchymal cells including MSC may be playing a crucial role in some or all of the effects of liver on the host immune system.

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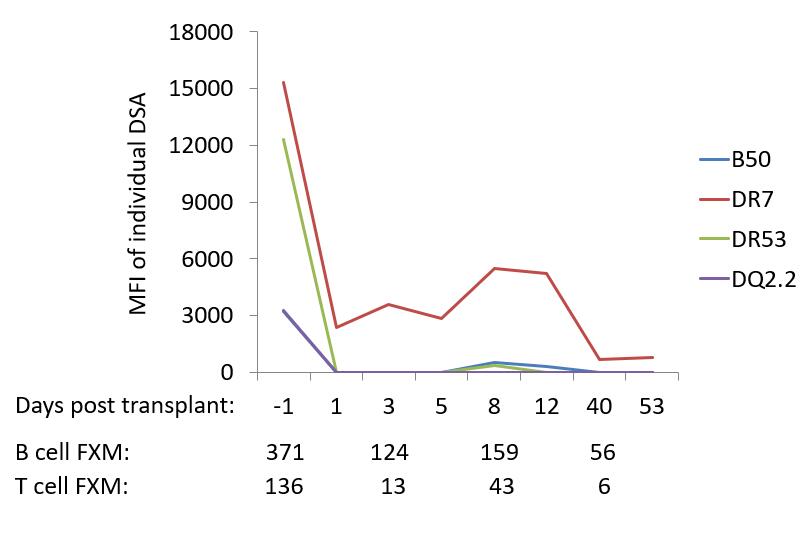
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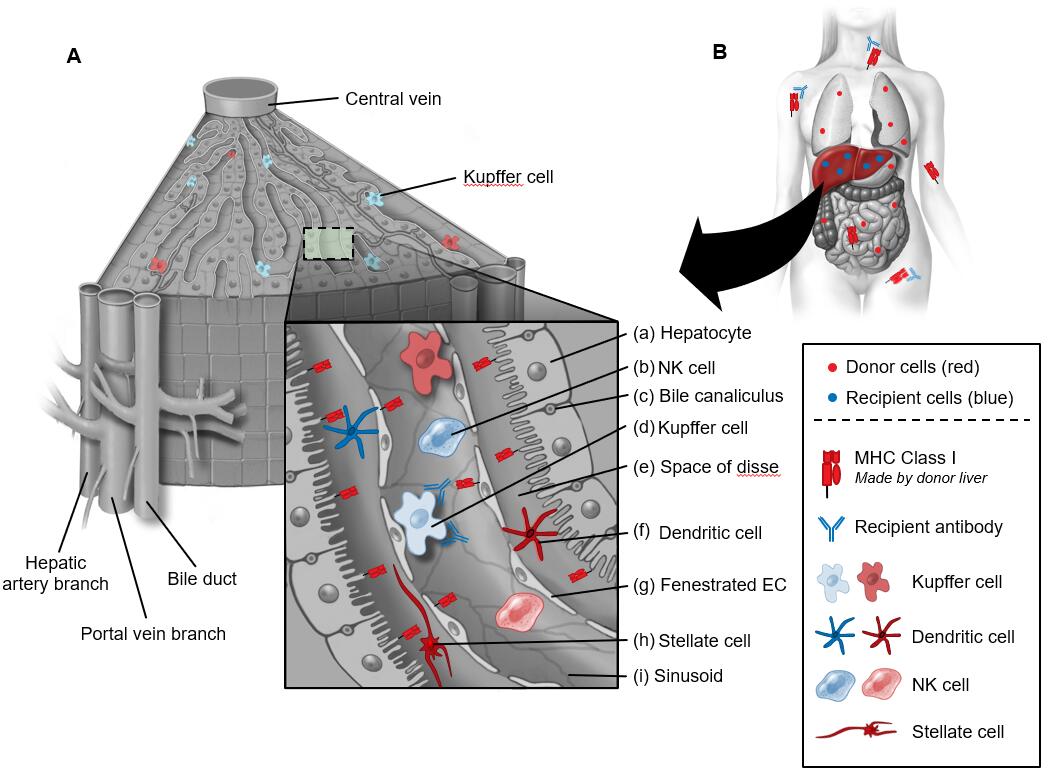
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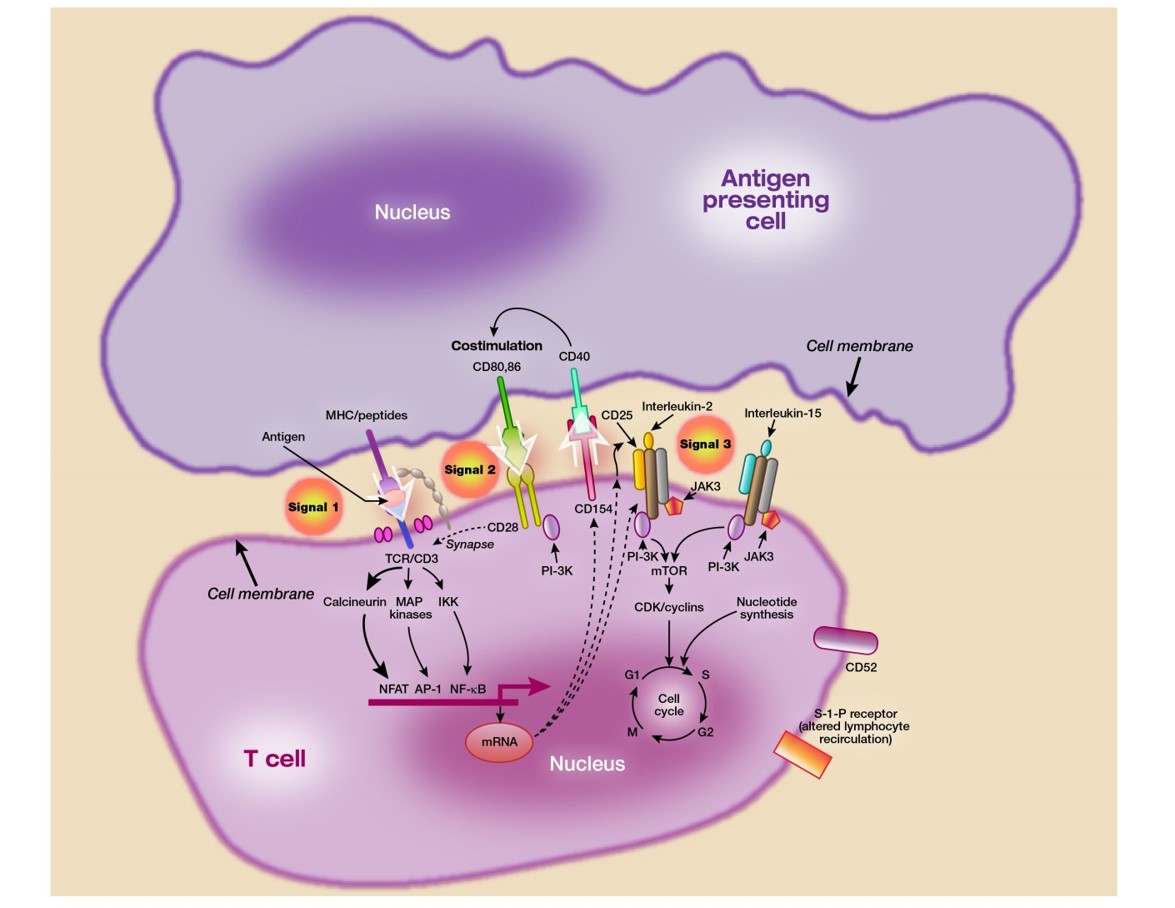
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**Figure 1 Typical course of donor-specific antibodies and flow cytometric cross match after liver transplant in a patient with fully functional liver allograft who is maintained on triple regimen immunosuppression (tacrolimus, mycophenolate, and prednisone).** DSA: Donor specific antibodies; FXM:Flow cytometric cross match.

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**Figure 2 Liver architecture and resident immune cells.** A:The liver’s unique architecture and the large number of passenger immune cells that accompany it during transplant likely play a role in its immunologic activity. Class I major histocompatibility (MHC) antigens are strongly expressed on bile ducts (c) and to a lesser extent on sinusoidal and endothelial cells (g). By contrast, Class II MHC antigens are primarily expressed on capillary endothelium, sinusoidal cells and dendritic cells (f). It is also recognized that cell surface MHC antigens are not static and can change in response to host and allograft dynamics such as infection and rejection; B: Liver transplants secrete soluble class I MHC antigens that bind and neutralize systemically circulating antibodies. Kupffer cells (d) also are involved in neutralization of antibodies. As such, liver allografts are thought to function as sinks for circulating immune complexes. EC: Endothelial cell; NK: Natural killer; MHC: Major histocompatibility complex.



**Figure 3 Activation of naïve helper T cells is thought to occur through a three signal pathway.** Signal 1, antigen recognition by the T cell receptor complex. Antigens are presented by major histocompatibility complex II cells [antigen presenting cells (APC) such dendritic cells]. Signal 2, co-stimulation, the interaction between the APC (CD80 and CD86) and the T cell (CD28). Signal 3, cellular proliferation and T cell differentiation into effector phenotypes (Th1, Th2), through cytokine stimulation. MHC: Major histocompatibility complex; APC: Antigen presenting cells.