Abstract. For many individuals with end-stage liver disease, the only treatment option is liver transplantation. However, liver transplant rejection is observed in 24%–80% of transplant patients and lifelong drug regimens that follow the transplant procedure lead to serious side effects. Furthermore, the pool of donor livers available for transplantation is far less than the demand. Well-characterized and physiologically relevant models of liver transplantation are crucial to a deeper understanding of the cellular processes governing the outcomes of liver transplantation and serve as a platform for testing new therapeutic strategies to enhance graft acceptance. Such a model has been found in the rat transplant model, which has an advantageous size for surgical procedures, similar postoperative immunological progression, and high genome match to the human liver. From rat liver transplant studies published in the last 5 years, it is clear that the rat model serves as a strong platform to elucidate transplant immunological mechanisms. Using the model, we have begun to uncover potential players and possible therapeutic targets to restore liver tolerance and preserve host immunocompetence. Here, we present an overview of recent literature for rat liver transplant models, with an aim to highlight the value of the models and to provide future perspectives on how these models could be further characterized to enhance the overall value of rat models to the field of liver transplantation.

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INTRODUCTION

The liver is an astonishing, unique organ capable of versatile immune responses and responsible for critical functions including detoxification, metabolism, and protein anabolism.¹ Today, prevalence of liver diseases is on the rise and liver transplantation (LT) remains the only treatment for end-stage liver disease, resulting in an increasing number of annual transplant operations of the liver.² Despite the immunologically privileged status of liver¹ and advances in LT surgical techniques and postoperative care since the first human LT in 1963, leading to an overall 5-year graft survival of 76% in adults,⁴,⁵ complications prevail. Long-term immunosuppression is part of the standard of care for LT patients with common immunosuppressive (IS) therapy including corticosteroids, antimetabolite, and specific inhibitors which block calcineurin and T-cell activation and proliferation.⁶,⁷ However, lifelong drug regimens significantly impact an individual’s quality of life and lead to higher risk of carcinomas, diabetes, viral infections, and opportunistic disease development.⁶,⁷ A serious LT complication, postoperative acute cellular rejection (ACR), develops in 24%–80% of patients, presents within days to weeks after surgery, and is a major contributor to graft failure leading to nearly 4% of posttransplant deaths.³ Currently, the management of ACR relies mainly on fine-tuned immunosuppression.⁵ By dissecting the mechanism of post-LT ACR and key players linked to immune tolerance, we may not only improve LT recipient’s survival but also quality of life.

An appropriate and relevant animal model that is able to recapitulate posttransplant liver immune microenvironment, delineate mechanisms involved in cellular rejection,
and explore promising therapeutics can help advance the field faster and in a more physiologically relevant way. The laboratory rat (Rattus norvegicus) model is the gold-standard animal model in orthotopic liver transplantation (OLT) for reasons such as optimal size for surgical operation and immunological similarity to the human liver. Many studies have explored the immunobiology and immunopathology of the liver or have tested for potential immunotargets using the rat model, as human liver samples are scarce. In this review, we will discuss how rat models are helping to advance the standard of care of LT patients, summarize the recent findings in the field, address the limitations of the models, and discuss the most promising way forward.

THE LANDSCAPE OF A HEALTHY LIVER

In the healthy state, parenchymal and nonparenchymal hepatic cells work cooperatively to establish the tolerogenic environment of the liver. Hepatocytes, liver sinusoidal endothelial cells (LSECs), hepatic stellate cells, liver resident macrophages (Kupffer cells [KCs]), regulatory T cells (Treg), and dendritic cells (DCs) among other hepatic cells work in synergy to create an IS microenvironment enriched in interleukin (IL)-10, transforming growth factor β (TGF-β), hepatocyte growth factor, retinoic acid, and prostaglandin E2 (PGE2). Professional antigen-presenting cells (APCs), such as DCs and KCs, and nonprofessional APCs, such as LSECs and hepatic stellate cells, not only favor the development of Treg in detriment of cytotoxic T lymphocytes but also promote clonal anergy and deletion of effector T and B cells. This response can be attributed mainly to the upregulation of inhibitory signals, such as programmed death ligand 1 (PD-L1) and cytotoxic T-lymphocyte–associated protein 4 (CTLA-4), and the low levels of the major histocompatibility complex (MHC) class I and of costimulatory molecules in the liver.

The inherent tolerogenic state of the liver reflects in less stringent criteria for matching LT donors and recipients when compared with other solid organs. As mentioned here, however, this perceived tolerogenic environment is not sufficient to control alloimmune responses, with most LT recipients requiring lifelong IS therapy. Interestingly, a small fraction of stable LT recipients have exhibited persistent graft tolerance following complete weaning of IS and are known as operationally tolerant.

Identifying biomarkers that can guide IS therapy withdrawal and understanding how this tolerogenic state can be achieved post-LT are key to develop drug-free, permanent tolerance that preserves host immunocompetence in LT patients. As part of this effort to identify reliable biomarkers that can determine whether LT patients are fit for IS withdrawal, Pérez-Sanz et al examined the predictive capacity of 20 biomarkers previously associated with LT tolerance and identified SENP6, FEM1C, miR31, and miR95 as promising candidates worthy of further investigation for their biological relevance and value in clinical practice.

OF RATS AND MEN

Rat Liver Transplant Models

A number of animal models have been used in LT research including the first animal OLT performed in canine species, pigs, mice, and rats. Of note, the rat model is especially valuable as it is small enough for easy handling and large enough to encompass complex microsurgical procedures, all the while presenting up to 90% genome match and post-LT immune progression similarity to humans. The rat OLT surgical procedure was first established in 1973 by Lee et al and has since served as a valuable experimental tool in rat LT research. Hindered by limitations of excessive bleeding and lengthy surgical time, the procedure was later modified by Kamada and Calne to include the cuff technique that significantly reduced the anhepatic phase and improved surgery outcomes. Besides instrumental when improving surgical procedures, organ preservation, and IS drug therapy in LT, as we will further discuss here, the rat model has been serving as a platform for the investigation of rejection and tolerance posttransplantation. Interestingly, liver allograft rejection is observed to have less severe manifestations or the graft is accepted without rejection when OLT involves donor-recipient pairs from certain rat strains, while it leads to fatal ACR in others. Table 1 shows the main combinations of rat strains used when studying tolerance or rejection post-LT. Analyzing 42 studies published within the past 5 years, the preferred methods for the investigation of ACR are (1) Lewis (LEW) as donor and Brown Norway (BN) as recipient (LEW→BN) (53%), and (2) Dark Agouti (DA) as donor and LEW as recipient (DA→LEW) (33%). It is important to highlight that by inverting the strains serving as donor and recipient, that is, BN→LEW and LEW→DA, we obtain instead a model for tolerance in LT studies (Table 1). Further understanding the mechanisms of rejection and tolerance in these models and their parallel/relevance to humans is critical for the design and testing of novel therapeutic strategies aimed at improving survival and health of LT recipients.

Similar to rats, select strain combinations in mice can also model both rejection and tolerance without IS treatment. For example, LT in the C3H→B10 mouse model presents rejection, while reversing the donor-recipient pair shows high tolerance rates despite mismatched MHC. Although mouse models have been helping to uncover molecular mechanisms and potential biomarkers of tolerance after LT (reviewed in Thomson et al), 1 major limitation of the model is its small size. LT surgeries comprise complex microvasculature reconnections involving sutures, cuff techniques, and lengthy operation times that are made difficult in small animals. Not only is anastomosis important in improving animal survival posttransplant, but the overall operation time also plays a large role in posttransplant complications and animal mortality. The establishment of surgical protocols in rats, which are 10 times the size of mice, allows for a more direct comparison with human LT by facilitating intricate microsurgery that mimics the human LT procedure, including hepatic artery reconstruction, which yields improved survival rates and reduced secondary complications from operational challenges. Higher precision in arterial and duct reconnections are especially important when trying to recapitulate the human physiological conditions in long-term tolerance modeling.

Assessment of Liver Allograft Acceptance

The use of rat models to explore ACR and tolerance in LT has provided valuable insights into postoperative
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rejection progression and served as a platform to test ACR treatments. Depending on the rat donor-recipient strain combinations used, specific levels and lengths of rejection vary. The degree of ACR can be determined with the help of laboratory-based and symptom-based tests. In laboratory-based histology, the Banff pattern is used to calculate Rejection Activity Index (RAI) based on changes in venous endothelial inflammation (E score), bile duct damage (B score), and portal inflammation (P score). In rejection rat combinations such as LEW → BN and DA → LEW, higher RAI values in tissue histology are indicative of increased rejection severity, hepatocyte necrosis, and worsened graft outcomes. In human LT, ACR normally presents in the first month after surgery and biopsies are taken to assess histology according to the Banff criteria.

### THE INTERCONNECTION BETWEEN THE LIVER AND THE BODY

The multifunctional, tolerogenic, and complex nature of the liver and the central position that it occupies in the physiology of the body must be taken into consideration when trying to decipher the phenomena of tolerance and rejection after LT. Further examining how the liver is seen once transplanted into another body and how it interacts and communicates with the body that it is within are the first steps toward a better understanding of LT.

### TABLE 1.

Combinations of rat strains as orthotopic liver transplantation donor and recipient to model postoperative acute cellular rejection and tolerance

| Donor strain (MHC) | Recipient strain (MHC) | References |
|--------------------|------------------------|------------|
| Acute cellular rejection | | |
| LEW (RT1) | BN (RT1) | 36-57 |
| DA (RT1 av1) | LEW (RT1) | 58-71 |
| LEW (RT1) | ACI (RT1) | 72 |
| ACI (RT1) | LEW (RT1) | 73,74 |
| Wistar (RT1/RT1) | SD (RT1/RT1) | 75,76 |
| SD (RT1/RT1) | Wistar (RT1/RT1) | 51 |
| Tolerance | | |
| BN (RT1) | LEW (RT1) | 50 |
| LEW (RT1) | DA (RT1) | 77-80 |
| DA (RT1 av1) | PVG (RT1) | 67 |
| Wistar (RT1/RT1) | August (RT1) | 81 |

ACI, AxC; BN, Brown Norway; DA, Dark Agouti; LEW, Lewis; MHC, major histocompatibility complex; PVG, piebald virol glaxo; SD, Sprague Dawley.

### FIGURE 1.

Development of ACR symptoms common to LEW→BN and DA→LEW rat OLT models. OLT rat recipients in alloimmune strain combinations that present ACR typically show lethargy and loss of appetite by POD3, and symptoms progressively worsen to development of severe jaundice and ascites from POD5. At the cellular level, assessment of RAI shows mononuclear cell infiltration as early as POD1, which continues to expand to liver parenchyma, with indication of severe ACR at POD7. LEW→BN rat OLT recipients typically survive for an average MST of 13.66 d, while DA→LEW models survive on average 11.22 d. ACR, acute cellular rejection; BN, Brown Norway; DA, Dark Agouti; IRI, ischemia-reperfusion injury; LEW, Lewis; MST, median survival time; OLT, orthotopic liver transplantation; POD, postoperative day; RAI, Rejection Activity Index.
The first days following LT are marked by upregulation of IL-2, interferon γ (IFN-γ), tumor necrosis factor (TNF), adhesion molecules such as vascular cell adhesion molecule-1 (VCAM-1), and intragraft leukocytes infiltration (Figure 2). This initial response can be attributed to (1) direct alloantigen recognition, where allograft APCs present intact alloantigen directly to host T cells, and (2) indirect alloantigen recognition, where host APCs process and present the allograft alloantigen to host T cells (Figure 2). The so-called passenger leukocytes (graft-derived immune cells) can also reach host lymphoid tissue and contribute to direct recognition of the alloantigen. The architecture of the liver combining a unique sinusoidal microvasculature marked by fenestrated endothelium,

![Diagram](image-url)

**FIGURE 2.** The days that follow an LT are marked by a dialogue between the liver graft and the host, through direct, indirect, and semidirect allorecognition. In this process of getting acquainted, the communication between different cells and the signals in the microenvironment of the liver will determine whether tolerance or rejection will follow. An initial inflammatory response, marked by IL-2, IFN-γ, TNF, and VCAM-1, leads to intragraft leukocyte infiltration independent of the final outcome and, in the presence of hepatic signals, is key to prime the response seen afterward. For instance, IFN-γ secretion is key to upregulate PD-L1 expression by LSECs, hepatocytes, and KCs, which subsequently contributes to the establishment of a tolerogenic environment. If, however, the alloantigen load or inflammatory signals, such as IL-2, are overexpressed, then the response is skewed toward acute cellular rejection. KCs are found in a spectrum from anti-inflammatory to proinflammatory phenotypes and are important contributors to the final outcome following LT. Activation of transcription factors, such as NF-κB, NFAT, RORγt, T-bet, leads to an environment enriched in proinflammatory cytokines and guide T-cell differentiation toward Th17 and Th1 responses. On the other hand, the engagement of PIK3/AKT and the upregulation of coinhibitory molecules, such as PD-L1, lead to a tolerogenic environment, with predominance of Treg and Th2 responses.

**Key Terms:**
- APC, antigen-presenting cell
- BD, bile duct
- CTL, cytotoxic T lymphocyte
- CTLA-4, cytotoxic T-lymphocyte–associated protein 4
- DA, Dark Agouti
- DC, dendritic cell
- FasL, Fas ligand
- FGL2, higher fibrinogen protein 2
- FOXP3, forkhead box P3
- GDF2, growth differentiation factor 2
- HA, hepatic artery
- HGF, hepatic growth factor
- HSC, hepatic stellate cell
- IFN-γ, interferon γ
- IFN-β, interferon β
- IL, interleukin
- KC, Kupffer cell
- LEW, Lewis
- LSEC, liver sinusoidal endothelial cell
- LT, liver transplant
- MHC, major histocompatibility complex
- NF-κB, nuclear factor of activated B cells
- NFAT, nuclear factor of activated T cells
- NK, natural killer
- PD-1, programmed cell death protein 1
- PI3K, phosphatidylinositol 3-kinase
- RORγt, retinoic acid receptor (RAR)–related orphan receptor (ROR)γt transcription factor
- T-bet, T-box transcription factor
- TGF-β, tumor growth factor β
- Th, T helper cell
- TNF, tumor necrosis factor
- Treg, regulatory T cells
- VCAM-1, vascular cell adhesion molecule-1
low blood pressure, and pattern of adhesion molecules facilitate the interaction between circulating immune cells and both parenchymal and nonparenchymal hepatic cells. Interestingly, in a mouse model of liver tolerance, a reduction in intragraft donor DCs was followed by an increase in host APCs expressing graft MHC molecule (also known as cross-dressing), which can present intact alloantigens to host T cells without further processing. Examining the cross-talk between donor hepatic cells, including parenchymal and immune cells, and the host immune cells is critical to determine the mechanisms associated with rejection and tolerance post-LT. Two aspects of LT that are especially intriguing are (1) the operational tolerance observed in a parcel of the LT recipients, and (2) the development of “infectious tolerance” post-LT. As mentioned, operational tolerant individuals can sustain transplant tolerance in the absence of IS therapy. Although the mechanisms associated with this state are poorly understood, there is evidence suggesting that increased frequency of Treg and hyporesponsive or exhausted T cells are important for sustaining tolerance in these individuals. “Infectious tolerance” on the other hand speaks to the extended tolerogenicity offered by liver allografts to other solid grafts from the same donor and has also been observed in rat models. Understanding the underlying mechanisms associated with spontaneous acceptance of graft is of great interest to medicine, and having animal models able to recapitulate the human LT procedures and outcomes can help advance the field in a more assertive way. Here, we review key mediators of rejection and tolerance, including recent findings uncovered by rat OLT models detailed in Table 2.

**Mechanisms Associated With Rejection of a Liver Graft**

As previously noted, the initial immune response post-LT is mediated through direct and indirect alloantigen presentation by APCs to host T cells, however little is known of the specific interactions within the graft immune microenvironment which dictate graft rejection versus tolerance. The experimental rat model with its high parallels to the human LT recipients, and (2) the development of “infectious tolerance” post-LT. As mentioned, operational tolerant individuals can sustain transplant tolerance in the absence of IS therapy. Although the mechanisms associated with this state are poorly understood, there is evidence suggesting that increased frequency of Treg and hyporesponsive or exhausted T cells are important for sustaining tolerance in these individuals. “Infectious tolerance” on the other hand speaks to the extended tolerogenicity offered by liver allografts to other solid grafts from the same donor and has also been observed in rat models. Understanding the underlying mechanisms associated with spontaneous acceptance of graft is of great interest to medicine, and having animal models able to recapitulate the human LT procedures and outcomes can help advance the field in a more assertive way. Here, we review key mediators of rejection and tolerance, including recent findings uncovered by rat OLT models detailed in Table 2.

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# Mechanisms of liver transplantation ACR and tolerance uncovered by rat OLT models

| Cell type | Mechanisms | Effect | Outcome | References |
|-----------|------------|--------|---------|------------|
| Cytotoxic T cells | Recognize alloantigen presented by APCs through direct or indirect recognition | Initiate downstream cell death pathways, inflammatory cytokine production, and immune cell recruitment to the liver graft | ACR | 36,49,73 |
| Th cells | Th1 and Th17 cells secrete IFN-γ, TNF, IL-2, IL-17, IL-22 | Promote proinflammatory microenvironment | ACR | 39,46,58,60,63,64,75 |
| Th2 cells produce IL-10, TGF-β, IL-4, IL-13 | | Promote anti-inflammatory microenvironment | Tolerance | 39,46,48,60 |
| Treg | Produce IL-10 and TGF-β | Modulate and maintain immature DC phenotype Inhibit CD8+ T-cell levels and function | Tolerance | 58,51 |
| DCs | Express FasL Secrete exosomes | Induce immunosuppressive T-cell (Treg, Th2) differentiation, reduce T-cell proliferation, induce antigen-specific T-cell apoptosis/anergy, reduce MHC and costimulatory molecule expression | Tolerance | 45,58,60,62,90 |
| Macrophages, KCs | Express costimulatory molecules | Activate lymphocytes and inflammatory cytokine secretion | ACR | 55 |
| | Produce IL-10 and TGF-β | Downregulate MHC and costimulatory molecule, upregulate MGAT5, which decreases T-cell activation and induces Treg differentiation | Tolerance | 41 |
| | Express PD-L1 | Inhibit T-cell proliferation, induce apoptosis, and suppress inflammatory cytokines | Tolerance | 37 |
| | Express TGF-β | Induce Treg expansion and reduce Th17 frequency | Tolerance | 43,63,72 |
| MSCs | Boost IL-10 produced by Treg | Promote anti-inflammatory microenvironment | Tolerance | 63,42,76 |
| | Express PD-L1 | Bind PD-1 to induce immunoregulatory signals | Tolerance | 48 |
| | Express HO-1 | Expand Treg population, maintain ACR attenuation past POD7 | Tolerance | |

| Molecule/p | Role in ACR | ACR/tolerance | ACR rat model | References |
|------------|-------------|---------------|--------------|------------|
| Gal-1/NF-κB/RelB | Gal-1 transfusion maintains tolerant DC phenotype and alleviates ACR | Tolerance | DA→LEW | 69,60 |
| | Limits DC-mediated CD4+ proliferation and increased Treg/Th17 ratio through NF-κB/RelB-IL-27 pathway | Tolerance | | |
| | Reduces NF-κB/RelB to maintain tolerant DC | Tolerance | | |
| Gal-9 | Perfusion with recombinant Ad-Gal-9 suppressed Th1 and Th17 differentiation | Tolerance | LEW→BN | 39 |
| Inflammasome activation pathway | Blocking ASC-mediated caspase-1 activation of IL-1β reduces ACR inflammation | Tolerance | DA→LEW | 68 |
| Ceruloplasmin/Nrf-2/ROS | Knock-down of ceruloplasmin diminished hepatocyte survival and lost protection from oxidative damage | Tolerance | DA→LEW | 67 |
| IL-22/STAT3 | Neutralization of IL-22 early in IRI stage worsened graft function | ACR | LEW→BN | 50 |
| | Neutralization of IL-22 late in ACR stage improved function | Tolerance | | |
| NFAT-BATF/JUN/IRF4-IL-21 | Inhibition of BATF/JUN/IRF4 complex attenuated rejection injury and decreased Bcl-6 and IL-21 in Th | ACR | DA→LEW | 66 |
| Fas/FasL | Mediates T-cell apoptosis to promote spontaneous acceptance of graft | Tolerance | DA→LEW | 62,90 |
| ERS-related IRE-1α/TRAF2/NF-κB | Suppression by gastrin inhibited in ACR attenuation and promotion of tolerogenic macrophage phenotype | ACR | LEW→BN | 40 |
| OX40/OX40L | Blocking OX40/OX40L resulted in less hepatic damage and longer survival | ACR | LEW→BN | 44 |
| HO-1 | Overexpression of HO-1 expanded Treg to maintain tolerance | Tolerance | LEW→BN | 48 |

Continued next page
with OX40L (CD252) on APCs to expand CD4⁺ T cells, showed that T-cell costimulatory molecule OX40 interacts
differentiation, respectively and levels of both transcrip-
t are measured. T-box transcription factor and ROR γ, IL-17, and IL-10 mRNA, and protein expression of IFN-
GATA-binding protein 3 and forkhead box P3 (FOXP3)
t is important in
γ secretion seems to upregulate PD-L1
ory macrophages and inhibited ACR
Stabilizing β-actin increases CD8⁺ T-cell apoptosis
and predicts ACR episodes
MSCs upregulate PD-L1 expression to attenuate
ACR
PD-L1 expression on KCs downregulated in ACR
Silencing PD-L1 increases inflammatory cytokines
and decreases tolerogenic cytokines
MSCs modified to express PD-L1/11g inhibited lymphocyte activity and attenuated ACR
(Tacrolimus) administration,
and predicted to obscure the underlying
mechanisms of graft tolerance.

As mentioned, an inflammatory response (marked by IL-2 and IFN-γ) is present early after human LT and is
common to both rejecting and tolerant rat models. In
particular, IFN-γ secretion seems to upregulate PD-L1 expression in LSECs and hepatocytes and
was shown as necessary for inducing tolerance.

The tolerogenic environment of the liver, marked by increased expression of IL-10 and TGF-β, weakens T-cell activation leading to short-lived T-cell populations incapable of assembling an effective immune response.
Specifically, liver APCs show low expression of costimulatory molecules (CD80/CD86) and elevated levels of inhibitory markers (PD-L1, CTLA-4) that impair T-cell activation (Figure 2). In addition, the liver microenvironment lacks stimulatory cytokines that act as the third signal in the events leading to T-cell activation, which include T-cell receptor (TCR) and CD28 interaction to MHC and B7 (also known as CD80/CD86) molecules on APCs as signals 1 and 2, respectively. The ability to induce Treg and mediate T-cell apoptosis is critical

### Mechanisms Associated With Acceptance of a Liver Graft

The genome similarity between rat and human, physiological relevance of rat OLT procedures, and spontaneous presentation of tolerance in select strain combinations of rat LT donors and recipients can help uncover the underlying mechanisms of graft tolerance.

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### TABLE 2. (Continued)

| Cell type | Mechanisms | Effect | Outcome | References |
|-----------|------------|--------|---------|------------|
| CTLA-4/B7 | Blocking CD28-B7 results in improved liver function and increased levels of immunomodulatory cytokines | Tolerance | LEW→BN | 45,55 |
| IDO       | Overexpression of IDO led to fewer ACR symptoms and better survival | Tolerance | LEW→BN | 55 |
| Autophagy pathway | Blocking autophagy decreased CD8⁺ T function, prolonging graft survival | ACR | LEW→BN | 36 |
| FGL/STAT1/ NF-κB | Overexpression of sFGL2 ameliorates ACR through anti-inflammatory cytokines and inhibit STAT1, NF-κB signaling, and induces immunoregulatory macrophage polarization | Tolerance | LEW→BN | 52 |
| IL-34/Pi3K/Akt | Overexpression of IL-34 polarized immunoregulatory macrophages and inhibited ACR | Tolerance | LEW→BN | 53 |
| β-actin   | Stabilizing β-actin increases CD8⁺ T-cell apoptosis and predicts ACR episodes | ACR | LEW→BN | 49 |
| PD-L1     | MSCs upregulate PD-L1 expression to attenuate ACR | Tolerance | LEW→BN | 42 |
| PD-L1 expression on KCs downregulated in ACR | Silencing PD-L1 increases inflammatory cytokines and decreases tolerogenic cytokines | Tolerance | | 37,37 |
| PD-L1 modified to express PD-L1/11g inhibited lymphocyte activity and attenuated ACR | | | | |
| | | | | |
| BN 76 | Tolerance | LEW→BN | | |
| BN 45,55 | Tolerance | LEW→BN | | 78 |
| BN 52 | Tolerance | LEW→BN | | |
| BN 55 | Tolerance | LEW→BN | | |
| BN 45,55 | Tolerance | LEW→BN | | |
| BN 36 | Tolerance | LEW→BN | | |
| BN 53 | Tolerance | LEW→BN | | |
| BN 52 | Tolerance | LEW→BN | | |
| BN 55 | Tolerance | LEW→BN | | |
| BN 45,55 | Tolerance | LEW→BN | | |
| BN 36 | Tolerance | LEW→BN | | |
| BN 53 | Tolerance | LEW→BN | | |
| BN 52 | Tolerance | LEW→BN | | |
| BN 55 | Tolerance | LEW→BN | | |
| BN 45,55 | Tolerance | LEW→BN | | |
| BN 36 | Tolerance | LEW→BN | | |
| BN 53 | Tolerance | LEW→BN | | |
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| BN 45,55 | Tolerance | LEW→BN | | |
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to liver graft acceptance. Blockade of PD-L1-PD-1 or B7-CTLA-4 in mice prevented apoptosis of alloreactive cells and converted spontaneous liver allograft tolerance into ACR. Meanwhile, Fas ligand (FasL) on APCs interact with Fas on T cells to induce specific T-cell apoptosis and promote immune tolerance.

Very much like in humans, the tolerance-promoting immune microenvironment in rats consists of primarily IS cytokines TGF-β, and IL-10, as depicted in Figure 2. In LEW→BN and DA→LEW rat combinations within the 42 rat model-based articles, TGF-β and IL-10 levels were observably diminished post-LT, correlating with higher rejection and shorter survival times. In assessing the role of TGF-β in tolerance promotion, Qiu et al cotransfected immature DCs (imDCs) with TGF-β and FasL and injected the modified imDCs into DA→LEW ACR recipients 5 days before LT. Post-LT, groups treated with single and cotransfections showed reduction in costimulatory molecules, diminished T-cell proliferation, and increased liver function and survival. Similarly, another study, through overexpression of TGF-β on mesenchymal stem cells (MSCs) in DA→LEW rats, saw ACR attenuation, improved mortality, and increased Treg/Th17 ratio within the graft. In contrast to Th1 cells, the main helper T cells population driving tolerogenic states in the rat liver are Th2 cells that secrete IL-4 and IL-13 along with TGF-β and IL-10, and evaluation of these IS cytokine expressions are often reported in parallel to Th1-associated cytokines in the literature.

Other soluble factors such as fibrinogen-like protein 2 (FGL2), discussed below, and acute-phase protein ceruloplasmin can also inform a tolerogenic microenvironment in the liver. Ceruloplasmin is produced by hepatocytes to mediate attenuation of reactive oxygen species damage and has been implicated to enhance hepatocyte survival post-transplantation. Ceruloplasmin levels are significantly higher in tolerant DA→Piebald Virol Glaxo rat models at POD63 than in DA→LEW models that expire at POD14, where systematic profiling showed that ceruloplasmin may be induced by IL-1β through ERK1/2-, JNK1-, and p39 MAPK activation of the AP-1 pathway which may contribute to operational tolerance.

LSECs and KCs are key cellular mediators of liver tolerance. They hold potent endocytic capacity and are able to present alloantigens to both CD8+ and CD4+ T cells. However, the low expression of costimulatory molecules (CD80/CD86) and high expression of the coinhibitory molecule PD-L1 promote alloreactive T-cell apoptosis, IL-10 and TGF-β secretion and Treg differentiation. The latter is one of the most accepted agents in liver acceptance. Treg (CD4+CD25highFOXP3+) direct the immune microenvironment toward a more tolerant state and are known to inhibit DC maturation through IL-10, TGF-β, PD-L1, and IDO induction. The bidirectional interaction between DCs and Treg is important in orchestrating graft tolerance, and the balance between Tregs and Th17 populations has also been suggested to maintain tolerance post-LT for individuals with liver disease. Furthermore, Treg have been associated with the mechanism of “infectious tolerance,” mentioned previously and are currently being assessed as a therapy in clinical trials, with most trials testing Treg infusion near the time of transplantation, as a way to induce tolerance (reviewed in Tang et al and Tanimine et al).

In the context of rat OLT, Treg have shown similar importance in inducing IS environments. In a study where Treg isolated from LEW rat spleens were infused into DA→LEW recipients 5 days before LT, rejection characterized by RAI values were significantly lower on POD3 and POD7 compared with control. Additionally, IS cytokines TGF-β and IL-10 were heightened, while proinflammatory IL-12 levels were inhibited. This immunotolerance effects of Treg can be moreover combined with the administration of immature DCs, further described below, to produce synergistic ACR attenuation effects. Furthermore, exosomes from Treg have also been implicated in tolerance induction. Exosomes were collected from CD4+CD25+ Treg culture medium and injected into SD→Wistar LT recipients on select days post-LT and resulted in an increased survival time from 28 to 90 days.

In vitro, CD8+ T-cell proliferation was inhibited by both Treg and Treg-derived exosomes, for which the effect could be reversed with exosome inhibitor administration. One of the challenges involving adoptive transfer of Tregs as cell therapy, however, is clinical-grade manufacturing, especially for ex vivo expansion Treg. Pharmacological and immunological treatments that favor in vivo expansion of this population are also being considered.

Myeloid-derived DCs (mDCs) in the liver microenvironment differentiate into a more tolerogenic phenotype. The presence of hepatocyte growth factor upregulates IL-10 expression by DCs monocytes, and the strong interaction established between Ag-specific Treg and mDCs can result in the removal of the MHC-antigen complex from the surface of the DCs, compromising their ability to present antigens. DCs have been extensively explored in rat models of LT. More specifically, DCs in the immature state (imDCs) or tolerogenic DCs, which express lower levels of MHC-II and costimulatory surface molecules, have been explored as local immunosuppression inducers in the liver post-transplantation. As previously mentioned, mDCs confer synergistic IS effects with Treg through upregulating tolerance-promoting cytokines, reducing T-cell proliferation, and inducing antigen-specific T-cell apoptosis. When constructed to overexpress IL-10 and FasL and injected into LEW→BN recipients 5 days before LT, DCs were able to reduce expression of MHC-II and costimulatory molecules (CD80/CD86), decrease T-cell expansion, and induce an IS environment. In particular, cotransduction with both IL-10 and FasL led to better results than single transductions alone. Tolerogenic DCs can be generated through induction by Gal-1 and other agents such as dexamethasone, VitD3, and rapamycin. Gal-1 regulates activated T-cell apoptosis, increases Treg, and suppresses direct DC recognition. A previous study by Peng et al explored treating DA→LEW recipients with Gal-1-induced DCs (DCgal-1) and apoptotic lymphocytes, both individually and synergistically. Proinflammatory cytokines declined on POD7 and an increase in tolerant cytokines was observed in the long term, while the transfusion promoted recipient T-cell hyporesponsiveness and Treg expansion. Treatment with both Gal-1 and apoptotic lymphocyte rendered better results compared with individual treatments alone and increased MST from 10 to 43.5 days.
Liver resident macrophages (KCs) have emerged as a critical determinant of posttransplant outcomes and are traditionally classified as anti-inflammatory or proinflammatory where polarization is dynamically influenced by cells such as Treg that moderate IS microenvironments. Activated KCs in the early ischemia-reperfusion injury stage release inflammatory IL-1, IL-6, and TNF cytokines and promote Th1 subtype differentiation, while anti-inflammatory KCs regulate Th2 differentiation and increase T-cell apoptosis. Previous rat studies, described below, show that proinflammatory phenotype can be directed through the NF-κB and MAPK pathway, while regulatory phenotype is promoted by higher FGL2 and IL-34 exposure and acts through the critical mechanistic target of rapamycin intermediate step, which informed current IS therapies of rapamycin inhibitors (eg, mechanistic target of rapamycin-Is). To explore the role macrophages play in immune tolerance induction post-LT and assess whether polarized macrophages direct tolerance, Yang et al performed an adoptive transfer experiment of CD163+, TGF-β- and IL-10-secreting anti-inflammatory macrophages into LEW→BN ACR recipients through the portal vein during OLT. The study resulted in diminished CD8+ T-cell infiltration, increased cellular apoptosis, and improved graft function and survival (25.0 ± 12.2 and 26.0 ± 12.4 d in BN- and LEW-derived anti-inflammatory macrophage-treated group, compared with 16.6 ± 2.3 d in PBS-treated group). The anti-inflammatory macrophage-treated rat group also saw a decreased expression of MHC-II+ cells and increased anti-N-acetylglicosaminyltransferase V expression, which has been previously implicated as a downstream target of NF-κB. These observations are consistent with in vivo studies that showed a phenotype switch from proinflammatory to anti-inflammatory KCs in the early ischemia-reperfusion injury stage, filling the niche thereafter are presumed to drive ACR or tolerance. Through monitoring changes in the LMP2 and LMP7 subunits of the immunoproteasome responsible for alloantigen recognition and processing, it was suggested that subunit levels may associate with anti-inflammatory macrophage activation through regulating antigen presentation and promote tolerance induction in phase 2.

In the search for ways to regulate macrophage polarization toward anti-inflammatory character, FGL2, which has previously been shown to possess immunoregulatory function and ability to promote tolerance in mouse and rat cardiac transplant models, was examined in the rat OLT model. Pan et al first observed upregulation of FGL2 mRNA and protein levels in the tolerant BN→LEW model and found that in vitro, FGL2 levels correlated with decreased STAT1 and NF-κB phosphorylation levels important for macrophage polarization. In vivo, overexpression of soluble FGL2 through injection of adenovirus expressing FGL2 (AAV-FGL2) into recipient rats of the ACR LEW→BN model before LT, induced polarization toward tolerogenic KC, promoted an IL-10 and TGF-β-rich immunotolerant microenvironment, and led to improved graft function and survival. Another recently explored cytokine associated with IS ability is IL-34. IL-34 is secreted by human and rodent Treg and has previously been associated with transplant tolerance in a rat cardiac transplant model. To determine IL-34 effects in the rat liver model, Zhao et al first noticed lower levels of IL-34 in LEW→BN ACR rat LT. Then through AAV-mediated overexpression of IL-34 and administration 30 days before LT in LEW→BN recipients, they observed attenuation of ACR in the treatment group. In vitro, IL-34 induced a phenotype switch from inflammatory KCs to noninflammatory KCs through activation of the PI3K/Akt pathway. Further adoptive transfer of lipopolysaccharide-stimulated and unstimulated KCs into AAV-IL-34-treated rats showed improved graft acceptance compared with no-KC transfer and control LEW→BN groups, suggesting that the ACR attenuation by IL-34 was mediated through noninflammatory KCs. Additionally, an increase of NK T-cell population was associated with acceptance of liver graft contributing to downregulating inflammatory responses and promoting a tolerogenic environment in mice. Further studies exploring the role of hepatic NK T cells in the induction of peripheral tolerance are also warranted.

As discussed here, different strategies able to induce tolerance in solid organ transplantation, including chimerism-based tolerance, and the main obstacles in the field are topic of great interest. Chimerism marks the establishment of donor hematopoietic cells within the recipient after transplantation. This state can be either transient or permanent and encompass >1% of hematopoietic cell lineages from donor origin (macrochimerism) or <1% (microchimerism). Several chimerism tolerance protocols based on bone marrow transplantation or hematopoietic stem cell transplantation have been tested both in preclinical and clinical settings and focuses specially on living-related renal transplantation. These protocols include different conditioning regimens of irradiation, IS, and T-cell deple-
constraints, including safety, feasibility, and applicability to deceased donor grafts, HLA-mismatched patients, and limited exploration in solid organs other than kidney. Chaudhry et al tested whether the induction of transient donor chimerism would lead to liver tolerance posttransplant using cynomolgus macaques as a model. Liver rejection was observed shortly after IS withdrawal (POD28) and suggested that interventions able to act during the inflammatory phase post-LT, such as depletion of CD8\(^+\) memory T cells, may also be required. Bone marrow cells have the capability of differentiating into liver KCs, endothelial cells, and hepatocytes and have been frequently explored as a way to induce tolerance in rat ACR models. In the DA→LEW model, replacing donor KCs with recipient bone marrow cells after total body irradiation pre-LT led to a decrease in IL-2 and IFN-\(\gamma\) levels at POD7 and attenuation of ACR. Multiple studies in LEW→BN, DA→LEW, and LEW→ACI rat models have also shown that MSCs, a subpopulation of cells within the BM, specifically function to attenuate ACR and improve survival rates in allogeneic transplants, with clear reductions in both liver enzymes and RAI at POD3 and POD7. MSCs originate from the bone marrow, are capable of self-renewal and pluripotency, and are speculated to attenuate ACR either through soluble immunomodulators or through cell to cell contact mechanisms. Treatment with MSCs in some cases may effectively hinder symptom progression by weeks or altogether. In particular, MSCs seem to mediate tolerance mechanisms through upregulating PD-L1 coinhibitory molecule expression to mitigate ACR progression. Assessment of helper T cells in MSC-treated conditions indicated that infusion of MSCs prefers anti-inflammatory-associated Th2 and T\(_{\text{reg}}\) differentiation over Th1 and Th17 cells. Aligning with these observations, proinflammatory cytokine levels were reduced, while anti-inflammatory cytokine levels of IL-10 and TGF-\(\beta\) increased. MSCs were also associated with an upregulation of splenocyte T\(_{\text{reg}}\) and these IS effects were enhanced when MSCs were transduced with heme oxygenase-1, an immunoregulatory player previously implicated in tolerance (Figure 2).

**Limitations**

Despite the key benefits of employing rat models of OLT, including optimal size for microsurgery, similarity in postoperative immunological progression, and high genome match of up to 90% resemblance to the human liver, the model is not without limitations. For one, although resembling human genetics, the MHC of the rat, RT1, and the rat genome is not as thoroughly characterized compared with the mouse model. Recent efforts, however, have advanced genome characterization in the rat model. As expected, differences also exist between rat and human genomes where certain genes map to different chromosomes and chromosomal locations, such as with the RT1 MHC class I gene. Moreover, although bigger in size compared with mouse models, rats are comparably smaller animals to humans, which render microsurgical techniques demanding for surgeons. Often, physiological relevance for long-term studies obtained from anastomoses that mimic human transplantation need to be balanced with simpler procedures and shortened surgical time.

**UNDERSTANDING THE LIVER IMMUNE MICROENVIRONMENT—CHALLENGES AND FUTURE DIRECTIONS**

Understanding the dynamic interaction between host and graft during rejection and tolerance post-LT and how different cell populations adapt to the evolving microenvironment and shape this environment is critical to the development of novel therapies. The systemic events that follow a liver transplant, in addition to events within the liver graft, are strong evidence of the ability of the liver to not only neutralize circulatory donor-specific antibodies but also to reshape itself and immune responses in different organs. Strategies able to instruct the recipient’s immune system to recognize donor antigens as self, eliminating the need for IS and maintaining their immune competence, are the ultimate goal. Grasping this complex network can certainly shed light on how posttransplant immunotolerance can be achieved and perhaps how to make more organs suitable for transplantation. The combination of single-cell transcriptomic techniques and functional assays can provide a more accurate description of the complex network of the liver, its interaction with the body, and key molecular pathways associated with rejection/acceptance. The use of single-cell RNA sequencing is revolutionizing the way we look at the cellular composition and function of different organs and has provided in-depth characterization of the liver landscape. This framework should be applied to the study of LT.

LT offers a unique opportunity to study mechanisms of immunosuppression, specially aimed at reprogramming the recipient immune system to establish and sustain tolerance in the absence of IS therapy. The characterization of rat OLT models helps to fully explore their potential as platforms to perform mechanistic studies of rejection and tolerance during transplantation. This may also contribute to new therapies that can help recover or improve outcomes with marginal organs, making them suitable for transplantation.

**CONCLUSION**

Through elegant rat models that recapitulate aspects of ACR and tolerance and can be modified to promote or treat either, we have begun to understand the underlying mechanisms of ACR and spontaneously developed operational tolerance, with hopes to inform therapeutics that will restore liver tolerance, obliterate IS therapy, and improve quality of postoperative life. However, the role of specific cell populations (donor and recipient) in transplant rejection remains unknown, which is limiting the ability to specifically target hepatic populations as part of IS strategies. Employing more specific strategies rather than systemic immunosuppression would limit off-target effects in patients. The overall goal of a deep characterization of immunological changes posttransplantation in rejection and tolerant individuals is to enable uncovering of the mechanistic basis that will guide the development of new therapeutic strategies to restore liver tolerance and preserve host immunocompetence.
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