Therapeutic Targeting of Hepatic Macrophages

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Abstract
Purpose of Review This review outlines the current knowledge about hepatic macrophages and provides an overview of therapeutic approaches to target hepatic macrophages for the treatment of liver diseases.
Recent Findings In recent years, it has been increasingly recognized that hepatic macrophages (resident macrophages, Kupffer cells, or circulating bone marrow monocyte-derived macrophages) are implicated in liver homeostasis as well as in disease progression and resolution. More recently, different populations of hepatic macrophages with distinct phenotypes and functions have been identified that have shown to play distinct roles in the pathogenesis of various acute and chronic liver diseases. The understanding of the role of hepatic macrophages in initiation, progression, and resolution of liver diseases has given rise to the development of therapeutics that can target different phenotypes of hepatic macrophages. Innovative strategies comprises of microRNA (miRNA), small interfering RNA (siRNA), therapeutic proteins, and small-molecule inhibitors.
Summary Evidence from recent in vitro and in vivo studies support the fact that hepatic macrophages can be efficiently targeted using miRNA/siRNA-based approaches, protein-based approaches, and small-molecule inhibitors for the treatment of liver diseases. However, more in-depth understanding underlying the roles of distinct macrophage phenotypes in different liver diseases is required for the translation of novel targeted therapeutics to the clinic.

Keywords Hepatic macrophages · Targeted therapeutics · Liver diseases · Monocytes · Kupffer cells

Introduction
The liver, the largest organ in the human body, exerts multiple vital bodily functions including blood detoxification, bile production, and synthesis, storage, and redistribution of different biomolecules like lipids, proteins, and carbohydrates [1, 2]. The liver also plays an important role in maintaining liver immune homeostasis and employs different mechanisms to suppress immune system and create immune tolerance [1–3]. Numerous studies have shown that hepatic macrophages [resident Kupffer cells (KCs) and monocyte-derived macrophages (MoMFs)] represent a diversified population of immune cells that are derived from varied sources and play a pivotal (and distinct) role in the initiation, progression, and restoration of liver diseases [4]. In recent years, incredible heterogeneity in these hepatic macrophages has been revealed, highlighting their complexity and involvement in liver diseases [2, 5–8]. While KCs maintain liver homeostasis, acute liver damage can result in activation of KCs into inflammatory phenotypes and instigate rapid infiltration of circulating monocytes into the injured liver resulting in intrahepatic macrophage imbalance. This imbalance results in increased liver inflammation, which can eventually deteriorate normal liver functions [8, 9]. Hepatic macrophages also engage closely with parenchymal (hepatocytes) and non-parenchymal cells, namely, hepatic stellate cells (HSCs) and liver sinusoidal endothelial cells (LSECs). HSCs are perisinusoidal cells located in a space between the LSECs and hepatocytes, called the space of Disse, and exert multiple functions such as vitamin A storage, immune-regulation, and remodeling of extracellular matrix [10–12]. Liver injury causes hepatic macrophages to release several inflammatory cytokines and chemokines that can activate these HSCs, thereby fostering not only inflammation but also liver fibrogenesis characterized by an excessive accumulation of ECM proteins, mainly produced...
by activated HSCs, leading to formation of scar tissue with
distorted liver architecture and loss of liver function [13, 14].
This review presents the brief overview of hepatic macro-
phages and discusses the advancements that have been made
in their understanding that has led to novel therapies that are
currently been examined in preclinical models and clinical
trials. Promising therapeutic approaches that have been sum-
marized in this review provide new perspectives towards the
effective and efficient treatment of liver diseases.

Origin and Heterogeneity of Hepatic
Macrophages

The liver comprises of about 80–90% of the macrophages in
the body [7]. Hepatic macrophages originate from different
sources, resulting in cellular heterogeneity in the liver [2].
Hepatic macrophages can be derived from resident hepatic
macrophages (KCs) or from distinct populations of infiltrating
macrophages, i.e., bone marrow (BM)-derived macrophages,
avascular peritoneal macrophages (PMs) that reside in subcap-
ular regions of the liver, and/or splenic monocytes [2, 4, 9,15,16]. Resident hepatic macrophages, the so-called KCs, are
non-migrating and self-renewing cells located at the luminal
side of the hepatic sinusoidal endothelium [15, 17]. Owing to
hepatic metabolic or toxic insults, circulating monocytes/macrophages known as bone marrow monocyte-derived macro-
phages (MoMFs) rapidly infiltrate the liver tissue [6].

Traditionally, macrophages depending on microenvironment-
mental signals have been classified into two distinct sub-
groups referred to as classically-activated M1 and alternative-
ly activated M2 macrophages [18, 19]. Macrophages can be
polarized towards the pro-inflammatory (and anti-
tumorinic) M1 phenotype in response to lipopolysaccha-
rides (LPS) and interferon gamma (IFN-γ) stimulation, secret-
ing high levels of pro-inflammatory cytokines. Alternatively,
macrophages can be polarized towards restorative or anti-
-inflammatory (and pro-tumorinic) M2 phenotype in re-
sponse to IL-4 and IL-13 stimulation. In contrast to M1, M2
macrophages suppress inflammatory responses and facilitate
tissue repair [18, 20]. M2 macrophages can be further subclassified into M2a, M2b, M2c, and M2d based on differ-
ent stimuli, distinct gene expression profiles, and functions
[21, 22]. However, recent studies have identified tremendous
diversity in macrophage phenotypes, attributed to their plas-
ticity, capable of adapting their phenotype in response to di-
verse microenvironmental as reviewed extensively elsewhere
[4, 5]. It is therefore imperative to gain mechanistic insights
into these distinct phenotypes of macrophages to develop
smart approaches to target selectively pathogenic phenotype
of macrophages for the treatment of pathological diseases
without affecting other phenotypic macrophages.

Role of Hepatic Macrophages in Liver
Diseases

Hepatic macrophages are important determinant in the patho-
genesis of liver diseases. They play a central role in promoting
inflammation, the hallmark of nearly all liver diseases includ-
ing acute liver failure (ALF), liver fibrosis, non-alcoholic fatty
liver disease (NAFLD)/non-alcoholic steatohepatitis (NASH),
alcoholic liver disease (ALD), viral hepatitis, and hepatocel-
lar carcinoma (HCC) [2]. When liver injury ensues, trig-
gered by multiple factors including alcohol/drug abuse, viral
infections, fatty diet, or auto-immune disorders [23], hepato-
cyes are injured and release mediators leading to the activation
of resident KCs. This results in the production of inflam-
matory cytokines and chemokines, e.g., C-C motif chemokine
ligands (CCLs) that induce massive infiltration of inflamma-
tory monocytes into the injured liver, leading to higher levels
of secreted cytokines and chemokines promoting chronic inflam-
mentation [9]. The role of hepatic macrophages in different
liver diseases has been extensively reviewed elsewhere [2, 8,9,15].

As mentioned earlier, hepatic macrophages interact with
HSCs, whereby, upon activation, HSCs undergo functional
and morphological changes resulting in proliferative myofibroblasts that produce excessive ECM and secrete nu-
erous pro-inflammatory and pro-fibrotic factors [12, 24].
Since it is known that these events promote inflammation and contribute to liver fibrogenesis, HSCs are considered to
be the main effector cells in hepatic fibrosis [25]. It has also
been shown that these myofibroblasts can differentiate hepatic
macrophages into pro-inflammatory and pro-fibrotic phen-
typic macrophages [26], therefore playing a central role in
liver fibrogenesis together with hepatic macrophages.

The plasticity and the capability of hepatic macrophages to
switch phenotypes according to microenvironmental signaling
account for their numerous functions in liver diseases.
Because of this, manifold challenges remain in understanding
the full spectrum of mechanisms of hepatic macrophages and
their activation, recruitment, and involvement in promoting
liver disease progression or in restoring damaged liver tissue
[9]. This knowledge will open promising opportunities to-
wards the treatment of liver diseases.

Therapeutic Targeting of Hepatic
Macrophages

Over past years, due to the growing knowledge about hepatic
macrophages, multiple mechanisms and pathways have been
identified and investigated that regulate their recruitment, ac-
tivation, differentiation, and polarization, based on which, a
number of novel therapies have been developed and examined
in various (pre-)clinical studies [2]. These novel therapies to
treat liver diseases mainly aim at targeting hepatic macrophages. Increasing evidence from mouse models and clinical studies in patients with liver diseases support the potential of hepatic macrophage targeting attributed to reduced inflammation thereby attenuation of liver fibrosis. Nowadays, targeting therapies are predominantly focused on (i) reducing the activation of KCs thereby reducing the release of pro-inflammatory mediators, (ii) promoting a switch from a pathogenic macrophage phenotype towards a restorative macrophage phenotype promoting disease resolution and liver regeneration, and/or (iii) reducing inflammatory monocyte recruitment into the injured liver to attenuate liver inflammation [2, 7, 15].

However, treatment options for most liver diseases still remain limited, owing to several challenges that are hampering the development of potential therapeutics: (1) discrepancy in macrophage phenotypes in humans and in animal models resulting in poor translation of therapeutics from animal models to human patients; (2) greater heterogeneity in humans as compared with inbred mouse strains due to several intrinsic (genetics, ethnicity, sex, and age) and extrinsic (microbiota, infections, medications) factors; (3) inadequate knowledge about human macrophages as compared with mouse models. Importantly, macrophages display tremendous heterogeneity, as shown in animal models and human liver diseases, with distinct functions in disease initiation and progression as well as disease regression and homeostasis [2, 8, 9, 15]. Therefore, it is crucial to target the pathogenic phenotypes of macrophages therapeutically without hindering the functions of so-called restorative or homeostatic macrophages [2].

We review multiple approaches that have been investigated for therapeutic targeting of hepatic macrophages. These approaches are divided into three categories as depicted in Table 1: (a) miRNA/siRNA-based approaches, (b) protein-based approaches, and (c) small-molecule inhibitors. We have also discussed synthetic nanoparticles (NPs) and extracellular vesicles (EVs) as efficient nanocarriers of therapeutics. While NPs mainly comprises of liposomes, polymeric NPs, and organic or inorganic NPs, EVs are biologically formed membrane-derived vesicles that carry multiple bioactive molecules to regulate cellular responses [41, 42]. NPs and EVs can both be selectively engineered for the delivery of several therapeutics including miRNA/siRNA, proteins, peptides, and small-molecule inhibitors to target hepatic macrophages for the treatment of liver diseases [43]. The major advantages of these delivery vehicles are improved stability, site-specific delivery, and thus increased pharmacokinetics of the encapsulated therapeutics [44].

**miRNA/siRNA-Based Approaches**

The increasing understanding of the role of molecular pathways in pathogenesis of many diseases has given rise to in-depth research for strategies to manipulate the expression of specific genes that are involved in liver pathogenesis. Gene expression is biologically regulated via RNA interference, wherein microRNAs (miRNAs) and small interfering RNAs (siRNAs) are fundamental. miRNAs and siRNAs are naturally occurring small non-coding RNA molecules composed of about 22 nucleotides that regulate gene expression via gene silencing at the posttranscriptional level [42, 43]. New insights illustrate that miRNA and siRNA expression is specifically altered in nearly all liver diseases, suggesting the great involvement of dysregulation of these non-coding RNAs in liver pathology [45, 46]. Therefore, extensive research has been performed into the possibilities of miRNAs and siRNAs to serve as a therapeutic target, whereby both inhibition and overexpression of specific miRNAs and siRNAs are therapeutically in reach [47]. However, the therapeutic approaches of miRNA and siRNA are quite different. This is mainly because miRNA is known to have multiple targets, whereas siRNA is highly specific with only one mRNA target [48]. Momen-Heravi et al. showed in vitro that EVs can successfully deliver a miR-155 inhibitor to macrophages. miR-155 is known to be involved in liver inflammation and fibrogenesis, and delivery of a miR-155 inhibitor to RAW macrophages has shown to cause >50% suppression of LPS-induced tumor necrosis factor alpha (TNF-α) production, a critical pro-inflammatory mediator involved in liver diseases [27, 48, 49]. He et al. have investigated the potential of mannose-modified trimethyl chitosan-cysteine (MTC)-conjugated NPs containing TNF-α siRNA for targeted inhibition of TNF-α expression by macrophages. Intriguingly, in vitro studies using RAW macrophages showed ~70% TNF-α silencing efficiency, requiring only 0.15 pm/well siRNA. These results were confirmed in an in vivo study, whereby orally delivered MTC NPs that contained a small amount of TNF-α siRNA inhibited TNF-α production in macrophages and protected mice from liver inflammation and damage caused by acute hepatic injury [28].

Besides using miRNA/siRNA to inhibit the secretion of pro-inflammatory mediators such as TNF-α by macrophages, there is another strategy that involves the inhibition of recruitment of circulating monocytes-derived macrophages. Upon liver injury, hepatic recruitment of MoMFs is mainly driven by C-C motif chemokine receptor/ligand interactions such as CCR2/CCL2 [7, 50, 51]. Since the infiltration of MoMFs contributes to inflammatory processes and thus to the progression of liver diseases, obstructing chemokine signaling can be an attractive potential therapeutic target to dampen the recruitment of monocytes into the liver [52]. Kim et al. showed that silencing the expression of CCR2 through delivery of anti-CCR2 siRNA to RAW macrophages induced ~60% reduction in CCR2 expression levels. Similar effects were observed in vivo in diet-induced obese mice, which showed decreased numbers of infiltrated monocytes into the injured liver. In
addition, ~ 70% reduction in TNF-α levels and complete reversal of steatosis was evidenced [29]. These results indicate that strategies comprising cell-targeted gene knockdown through miRNA/siRNA are promising therapeutic approaches for the treatment of various liver diseases. However, the implementation of these two RNA molecules as novel therapeutics currently face some limitations with regard to clinical translation due to potential off-target effects, delivery efficacy, and poor stability in vivo.

However with successful implementation of Onpattro, first siRNA-based lipid nanoparticles for the treatment of hereditary transthyretin amyloidosis [53], siRNA/miRNA-based (nano) therapeutics offer exciting and daring platforms for scientists to overcome these challenges.

### Protein-Based Approaches

Proteins, especially monoclonal antibodies, and peptides are highly promising therapeutics due to their high biocompatibility, non-toxicity, biodegradability, and selectivity [54]. Nowadays, monoclonal antibodies are receiving more and more attention from scientists to serve as therapeutic agents due to their specificity and capability of blocking the activity of specific signaling molecules, cell surface markers, or enzymes [55, 56]. Lundbäck and co-workers explored the therapeutic application of a partially humanized anti-high mobility group box 1 (HMGB1) monoclonal antibody in drug-induced liver injury (DILI) and ALF.

**Table 1** Therapeutic targeting strategies of hepatic macrophages

| Strategy               | (nano) Therapeutics                                      | Mechanism                                                                 | Reference(s)                      |
|------------------------|----------------------------------------------------------|---------------------------------------------------------------------------|----------------------------------|
| miRNA/siRNA-based      | EV-based delivery of miR-155 inhibitor                   | Inhibition of KC activation                                               | (Momen-Heravi et al., 2014) [27] |
| approaches             | Mannose-modified trimethyl chitosan-cysteine (MTC) TNFα siRNA NPs | Inhibition of KC activation                                               | (He et al., 2013) [28]           |
|                        | Anti-CCR2 siRNA                                          | Dampening of monocyte recruitment                                        | (Kim et al., 2016) [29]          |
| Protein-based          | Partially humanized anti-HMGB1 monoclonal antibody      | Inhibition of KC activation, dampening of monocyte recruitment            | (Lundbäck et al., 2016) [30]     |
| approaches             | IL-4/anti-IL-4 monoclonal antibody complex (IL-4c)       | Modulation of macrophage polarization                                     | (Lynch et al., 2019) [31]        |
|                        | Human monoclonal antibody BTT-1023                       | Dampening of monocyte recruitment                                        | (Amditz et al., 2017) [32]       |
| Small-molecule         | ASK1 inhibitor selonsertib                               | Inhibition of KC activation                                               | (Loomba et al., 2018) [33]       |
| inhibitors             | NIK inhibitor B022                                       | Inhibition of KC activation                                               | (Xiaomeng et al., 2016) [34]     |
|                        | CCR2/CCR5 antagonist cenicriviroc                        | Dampening of monocyte recruitment                                        | (Lefebvre et al., 2016; Krenkel et al., 2018) [35, 36] |
|                        | CTD-002                                                  | Inhibition of KC activation, dampening of monocyte recruitment            | (Khranana et al., 2019) [37]     |
|                        | SYK pathway inhibitor R406                               | Inhibition of KC activation, dampening of monocyte recruitment            | (Bukong et al., 2016) [38]       |
|                        | R406-PLGA NPs                                            | Inhibition of KC activation, dampening of monocyte recruitment            | (Kurniawan et al., 2018) [39]    |
|                        | Docosahexanoic acid (DHA)                                | Modulation of macrophage polarization                                     | (Carpino et al., 2016) [40]      |

*Abbreviations used are ASK-1, apoptosis signal-regulating kinase 1; CCR2, chemokine (C-C motif) receptor 2; CCR5, chemokine (C-C motif) receptor 5; CTD, cathepsin D; DHA, docosahexanoic acid; EV, extracellular vesicle; HMGB1, high mobility group box 1; KC, Kupffer cell(s); MTC, mannose-modified trimethyl chitosan-cysteine; NF-κB, nuclear factor kappa-light-chain-enhancer of activated B cells; NIK, NF-κB-inducing kinase; NPs, nanoparticles; PLGA, poly(lactic-co-glycolic acid); SYK, spleen tyrosine kinase.

HMGB1 is one of the most well-known damage-associated molecular patterns (DAMPS), and the release of HMGB1 by macrophages and other immune cells is correlated with liver injury and inflammation with poor prognosis. Interestingly, mice treated with HMGB1-neutralizing antibody showed reduced liver injury and inflammation [30, 57]. More recently, Lynch et al. reported that administration of IL-4/anti-IL-4 monoclonal antibody complexes (IL-4c) reduced the number of pro-inflammatory macrophages in C57Bl/6 mice with acute liver injury. IL-4c therefore were capable of fostering liver regeneration upon acute injury through promoting a switch from macrophages with a pathogenic phenotype towards macrophages with a restorative phenotype [31].

Furthermore, human monoclonal antibody BTT-1023, also known as Timolumab that targets vascular adhesion protein-1 (VAP-1), is currently examined as novel treatment option for the progressive inflammatory liver disease, primary sclerosing cholangitis (PSC). VAP-1 is an endothelial glycoprotein that is highly implicated in leukocyte migration towards the inflamed site, thereby driving inflammation and fibrosis. Blockade of VAP-1 via BTT-1023 can therefore be an interesting approach to attenuate leukocyte trafficking and thus inhibit inflammatory responses. At present, BTT-1023 is intensively studied to determine its safety, dose, and efficacy [32, 58, 59].
Small-Molecule Inhibitors

Small-molecule inhibitors are compounds with low molecular weight (< 1000 Da), which allows them to translocate through cell membranes to reach and interact with intracellular targets to disrupt specific pathological pathways [60]. Since hepatic macrophages are known to be regulated by multiple inflammatory signaling pathways including apoptosis signal-regulating kinase 1 (ASK1) and nuclear factor kappa-light-chain-enhancer of activated B cells (NF-κB) [61], small-molecule inhibitors targeting these signaling pathways have been developed and explored for the treatment of liver diseases. Looma and co-workers has evaluated a small-molecule ASK1 inhibitor, selonsertib, which has shown to inhibit hepatocyte metabolism and macrophage activation. In patients with NASH and fibrosis (randomized phase 2 trial), selonsertib evidenced an improvement in fibrosis, lobular inflammation, and serum biomarkers of apoptosis and necrosis [33], while in STELLAR-3 and STELLAR-4 randomized double-blind placebo-controlled phase 3 trials in NASH patients with F3 fibrosis and F4 cirrhosis, respectively, selonsertib failed to show a significant improvement in fibrosis. In another study, Xiaomeng et al. found a potent small-molecule inhibitor (B022) that efficiently inhibits NF-κB-inducing kinase (NIK). Inhibiting this inflammatory NIK pathway through B022 has shown to ameliorate macrophage-induced liver inflammation, as macrophages (mainly KCs) are strong activators of NF-κB [34, 62].

Another strategy besides inhibition of macrophage activation via specific inflammatory signaling pathways involves, as mentioned earlier, the dampening of inflammatory monocyte recruitment via CCR2/CCR5 [15]. Studies have demonstrated that blocking of CCL2-mediated monocyte recruitment using a selective CCR2/CCR5 small-molecule inhibitor, cenicriviroc, ameliorates inflammation, and fibrosis in the liver and kidney fibrosis mouse models [35, 36]. Interestingly, Friedman et al., in patients with NASH (NAFLD activity score, NAS ≥ 4) and stage 1–3 liver fibrosis, has shown ≥ 2-point improvement in NAS and ≥ 1 stage improvement in fibrosis with favorable safety and tolerability after 1-year treatment with cenicriviroc in a CENTAUR phase 2b clinical study [63].

Khurana and co-workers investigated a highly specific small-molecule inhibitor CTD-002 of extracellular cathepsin D (CTSD), a lysosomal enzyme that is known to be associated with lipid-related disorders such as NAFLD. In this study, bone marrow-derived macrophages (BMDMs) were treated with CTD-002 and incubation with CTD-002-treated BMDMs-conditioned medium on human hepatocellular carcinoma (HepG2) cells resulted in reduced levels of inflammation and improved cholesterol metabolism. In addition, high fat diet-fed rats treated with CTD-002 showed attenuated hepatic steatosis [37]. Bukong et al., and Kurniawan and colleagues demonstrated the potential of a small-molecule inhibitor (R406) that inhibits spleen tyrosine kinase (SYK) [38, 39]. SYK signaling pathway has been shown to play an important role in the pathogenesis of alcoholic and non-alcoholic steatohepatitis driving liver inflammation and steatosis [38, 39]. SYK inhibitor, R406 ameliorated hepatic injury, steatosis and inflammation in alcoholic steatohepatitis in mice by inhibition of recruitment of immune cells macrophage and inflammasome activation [38]. R406, when delivered via poly(lactoid-co-glycolic acid) (PLGA) NPs, showed significant inhibition of M1-specific inflammatory markers in RAW and BMDMs in vitro and attenuated liver inflammation, fibrosis, and steatosis in vivo in methionine–choline-deficient (MCD) diet-induced NASH mouse model [39].

Furthermore, Carpino et al. demonstrated that docosahexanoic acid (DHA), an omega-3 fatty acid, abrogated pediatric NAFLD through the interaction with G protein-coupled receptor 120 (GPR120) and reduced expression of NF-κB. GPR120 is known to be involved in macrophage polarization towards the M1/M2 phenotype. Treatment with DHA resulted in decreased polarization of macrophages towards the inflammatory M1 phenotype and increased polarization towards M2 macrophages, along with the improvement in NAS and reduction in hepatic progenitor cell activation, thereby suppressing inflammatory responses and fostering tissue repair respectively [40, 64].

Conclusions

Our improved fundamental understanding about the initiation, progression, and resolution of liver diseases has provided us the tools for the development of promising therapeutic approaches for the treatment of liver diseases. Hepatic macrophages play an essential role in maintaining liver homeostasis; however, numerous studies have found that when imbalance in their functioning occurs owing to liver injury, they are also involved in the initiation and progression of different etiological liver diseases. Intriguingly, hepatic macrophages possess tremendous heterogeneity and therefore possess diverse phenotypes with discrete functions in the liver. In the past decades, multiple approaches have been developed to target hepatic macrophages for the treatment of liver diseases. These approaches mainly aim at inhibiting KC activation, influencing macrophage polarization, and inhibiting monocyte infiltration, and great progress is being made in the application of specifically modified EVs and synthetic NPs for the delivery of therapeutics. As described in this review, innovative therapeutic strategies entail the application of miRNA/siRNA to alter expression levels of specific genes that are correlated with disease progression. Another strategy involves protein-based macrophage targeting, wherein monoclonal antibodies
appear as increasingly interesting therapeutic agents due to their high specificity. Small-molecule inhibitors are the most extensively studied therapeutics, and plenty of them are currently advancing to clinical trials. Nevertheless, also miRNA/ siRNA- and protein-based approaches have already shown promising results from in vitro studies and in experimental animal models. However, there are still several obstacles that are hampering the translation from animal models to the clinic, e.g., disparity in macrophage phenotypes and the fact that there is larger macrophage heterogeneity in humans in comparison with animal models. Furthermore, it still remains not fully understood, yet a subject of intensive research, what the underlying mechanisms of hepatic macrophage phenotype switching are during the diverse stages of liver diseases. Therefore, in-depth investigation of distinct macrophage phenotypes and their exact role in liver diseases is required to help unravel the complex liver pathobiology and to develop novel therapeutic targets to treat liver diseases in the clinic.

Compliance with Ethical Standards

Conflict of Interest Nijland and Dr. Bansal declare that they have no conflicts of interest.

Human and Animal Rights and Informed Consent This article does not contain any studies with human or animal subjects performed by any of the authors.

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