The field of adeno-associated virus (AAV) gene therapy has progressed rapidly over the past decade, with the advent of novel capsid serotype and organ-specific promoters, and an increasing understanding of the immune response to AAV administration. In particular, liver-directed therapy has made remarkable strides, with a number of clinical trials currently planned and ongoing in hemophilia A and B, as well as other liver disorders. This review focuses on liver-directed AAV gene therapy, including historic context, current challenges, and future developments.

Keywords: gene therapy, hemophilia, AAV, liver, clinical trials, metabolic

INTRODUCTION

Today, there is an intense interest in developing gene therapy products for monogenic diseases, with the liver garnering significant attention due to the large number of conditions affecting hepatocytes and liver function. The scope of this review is limited to advances in gene therapy applications employing vectors based on adeno-associated virus (AAV), and the clinical and commercial applications of these vectors for the treatment of liver-based diseases. Multiple advances in the field have led to the continued refinement of AAV vector properties, the discovery of new versions of AAV capsids, and the completion of numerous robust therapeutic studies in relevant animal models of inherited diseases in animals, both small and large. To enable a full discussion of this field, the properties of the liver that make it amenable to AAV gene therapy and the properties of AAV vectors that make them especially relevant to liver gene therapy are reviewed. In addition, aspects of nonclinical studies that support clinical studies, as well as those that might offer insight into the field, are reviewed. Early and ongoing clinical trials are examined, attempting to draw some conclusions across the field, and, finally, the challenges for the broad clinical application of AAV gene therapy vectors for liver diseases are examined.

Much of the current enthusiasm for liver-directed AAV gene therapy product development stems from preclinical and clinical successes in the field of hemophilia B. Numerous studies in classic mouse and dog models of hemophilia A and B have demonstrated clear and robust long-term benefit from administration of AAV vectors encoding the relevant clotting factors, with the vector trafficking to the liver for gene expression. However, the path to clear clinical benefit for AAV hemophilia gene therapy has been a winding one. The first liver-directed clinical trial in hemophilia B, run by Manno et al., showed initial success, achieving remarkable factor IX (FIX) expression levels of ~11% of normal, only to have levels subsequently return to the patient’s baseline of <1% of normal within weeks. The concomitant rise in liver enzymes stimulated the hypothesis that an immune response to the vector capsid was the culprit for the decline in expression and triggered an intense period of research focusing on immune responses to AAV.

A portion of the limitation of the Manno trial was placed on the vector capsid, AAV2, which in turn stimulated a period of concentrated development of alternative AAV capsids to overcome these AAV2 restrictions. After a 5-year clinical hiatus in the field, Nathwani et al. published the results of a
study in which an AAV8 vector was able to deliver clinically meaningful levels of FIX to hemophilia B patients. The salient achievements of the Nathwani trial included demonstration of clinical benefit with restoration of as low as ~2% of normal FIX expression, sustained expression of FIX for >5 years at ≥5%, and the demonstration that intervention into the presumptive immune response with steroid administration could abort the increases in liver enzymes and preserve FIX expression.6

These encouraging results have sparked multiple AAV gene therapy trials for hemophilia B, as well as other liver and metabolic diseases. Additional improvements in delivery, capsid and vector genome designs, and better understanding of the immune response are building upon these encouraging results, and will potentially lead to treatments for other rare disorders.

PROPERTIES OF THE LIVER

The liver is a vital organ and plays an important role in human metabolism and other key physiologic functions.7,8 Hepatocytes, the most common type of liver cell, synthesize and metabolize a large number of proteins, including intracellular and secreted proteins responsible for a diverse range of critical functions in the body.9,10 These functions include carbohydrate use and storage, lipid metabolism, and hemostasis. In addition, the liver plays an important role in detoxifying naturally occurring metabolites such as ammonia and bilirubin. Numerous inherited metabolic disorders (IMDs) have their origin in the liver. Disruption of key metabolic pathways modulated by the liver can lead to the accumulation of toxic products and subsequent liver damage, or to the inappropriate or insufficient production of key proteins for proper metabolic function, both of which can have a range of deleterious systemic effects. Today, there are >400 described rare monogenic disorders associated with the liver.11–17

The liver also possesses unique anatomic properties that make it a preferred target for AAV-based gene therapy.18 Due to its unique dual blood supply, an adult human liver receives nearly 25% of the cardiac output of blood, filtering approximately 1 L of blood per minute, and accounts for 10–15% of the blood volume at any given moment.19 This means that administration of gene therapy systemically leads to rapid accumulation of high levels of vector particles within the liver. Specialized fenestrated endothelium along hepatic sinusoids allows for free passage of vector particles from the blood directly to the hepatocytes, making transduction of cells much more efficient than in organs with continuous endothelium (see Fig. 1). In part due to these anatomic features, the liver has the highest recorded AAV genome copies per tissue of natural isolates outside of primary immune organs.20,21

Because AAV vectors are largely non-integrating, an important property to consider with liver-directed gene therapy is the longevity of the transduced hepatocyte. In a fully developed liver, under normal conditions, <1–2% of hepatocytes are turning over at any given time, while the rest remain in a quiescent state.14,15,22 In addition, turnover of mature hepatocytes occurs relatively slowly, with the average life span of non-resting hepatocytes ranging from 200 to 300 days.16 Given the relative lack of hepatocyte turnover, gene therapy administered to these cells should experience minimal dilutional effects due to new hepatocyte generation. In the Nathwani study, published in the New England Journal of Medicine, hemophilia B patients are still experiencing durable FIX expression and clinical benefit for >5 years from the time of vector administration.6,23

The ideal goal of treatment for liver-associated monogenic diseases, including IMDs, is to revert a harmful or disease-causing genetic condition to its normal state. Substantial progress has been made in the treatment of these patients over the past six decades, beginning with the first report of dietary control of phenylketonuria in the early 1950s.24 However, many of these approaches are inadequate for long-term control. Monogenic diseases associated with the liver are particularly well-suited to gene therapy, as the nature of these diseases permits the use of well-described, and often clinically validated, biomarkers. Because gene therapy aims to revert an abnormal genetic condition toward a more normal state, the use of well-understood biomarkers can inform dose selection, enable evaluation of therapeutic efficacy, and serve as a surrogate for early signals of clinical benefit.

AAV VECTORS IN GENE THERAPY

There are several reasons why AAV vectors are useful for liver-directed gene therapy applications, especially when considering commercial drug development. An AAV vector consists of only two components: the protein coat, or vector capsid, and the vector DNA genome. Differences in the vector capsid are responsible for serological classification of vectors (i.e., AAV1 through AAV6) and structural classification of vectors with the advent of AAV7 and beyond.20,21 Despite differences in capsid protein amino acid sequence, all AAV capsids adhere to certain core structural themes,25 which to a large degree dictate the size and therefore the
DNA-carrying capacity of AAV vectors. Considering AAV vectors as biologic products, the conserved vector capsid structure and the inherent stability thereof offer advantages. Because no other proteins participate in the structure of the vector capsid, the analytical requirements for purity and identity of the final AAV vector product are simpler compared with other vector systems. The inherent stability of an AAV vector capsid also enables the application of robust downstream purification unit operations such as heat inactivation of potential adventitious agents, ion exchange chromatography, and various filtration and liquid handling operations that are standard within the biologics manufacturing realm.26–28

Using the well-studied AAV2 as the canonical representative, the AAV virus genome is limited to 4,679 nucleotides with 290 nucleotides reserved for the ITRs, the latter being the only cis-acting requirement for packaging of an AAV vector genome within the capsid.29 Thus, in the ideal case, the carrying capacity of an AAV vector genome, including transcriptional control sequences, any 5’ and 3’ mRNA untranslated sequences, protein (or RNA) coding sequences, and poly-A addition signals is limited to approximately 4,400 nucleotides. Therefore, in each case, the precise nucleotide sequence of the AAV vector is known and analytical tools exist to enable sequencing of the plasmids entering the manufacturing process and of the AAV vector that emerges as the final purified product. For large protein coding sequences, there are stringent size limitations on each of the noncoding elements within the AAV vector genome, and attempts to produce oversized vectors have resulted in inefficient packaging and truncated genomes.30,31 To overcome the size limitation challenge, dual vector systems have been proposed, where the larger genome would be recreated through recombination of the partial genomes.32 In addition, the effort to develop cassettes for factor VIII (FVIII) expression serves as an object lesson in setting expectations for both efficacy and quality of large-genome rAAV. Bearing in mind the well-established packaging “cliff” of approximately 5.2 kb, success has only come after considerable effort from individual groups to minimize the size of the protein itself (deletion of the B-domain), establish the bare minimum regulatory sequences required for a reasonable level of tissue-specific expression, and test alternate serotypes for greater size tolerance.31,33–35 For small protein coding sequences, there is ample space within an AAV vector genome, but the opposite problem, having a vector genome that is too small,
can also occur. Engineering a vector genome that is too small can lead to inclusion of the dimeric replication intermediates, adjacent plasmid sequences, or higher levels of host cell DNA.\textsuperscript{36} Taken together, the above considerations indicate that an AAV vector product can be manufactured and characterized as a well-controlled biologics product, employing systems that are currently established within the biologics manufacturing world.

**CHOICE OF SEROTYPE**

Given the anatomical features of the liver reviewed in the introduction, it might seem that achieving effective liver transduction following systemic delivery would be serotype independent. However, it was apparent from early clinical trials seeking to deliver FIX that improvement in AAV2 was needed, which stimulated the development of new AAV capsids for clinical use.

In 2002, the serotype AAV8 was discovered in rhesus macaques.\textsuperscript{37} Reports from multiple groups confirmed that AAV8, a member of the Clade E family of AAV, outperformed AAV2 and others in liver transduction and expression, demonstrating that serotype choice plays a significant role in successful transgene delivery.\textsuperscript{37–39} Key advantages of vectors based on AAV8 and other members of the Clade E family include the ability to achieve a high level of hepatocyte gene transfer via peripheral vein administration, presence of a lower proportion of neutralizing antibodies in the human population, and commercially useful manufacturing properties such as high yield and release of vector into the cell culture medium.\textsuperscript{40,41} The research that led to the early clinical development of AAV8 vectors culminated in the aforementioned successful clinical studies by Nathwani et al., in which all hemophilia B subjects in the trial appeared to derive clinical benefit from the experimental gene transfer protocol. Importantly, a smaller proportion of potential subjects failed the neutralizing antibody screening test, allowing for more facile patient enrollment. The peripheral vein administration route was successful in delivering the AAV8 FIX vector to the liver, and the elicitation of ALT after vector infusion was reduced and readily controlled by a course of prednisone treatment.\textsuperscript{6}

Concurrent with ongoing searches for totally new AAV capsids, many groups have sought to engineer current serotypes to improve liver transduction and/or avoid preexisting immunity. These include alteration of surface residues to reduce phosphorylation and improve trafficking, peptide insertion, ligand modification, and so-called “directed evolution” or capsid shuffling.\textsuperscript{42–47} One engineered serotype, LK03, was identified using the latter method in the context of a rodent model in which mice livers were repopulated with human hepatocytes.\textsuperscript{48} Reports on the in vivo transduction superiority of LK03 versus AAV8 and AAV3B are mixed, probably due to the models used. Regardless, LK03’s most promising characteristics may be that it seems to be more resistant than AAV2 and AAV3B to pooled human IVIG, a surrogate for preexisting neutralizing antibodies in the human population, and therefore may be available to a greater number of patients.\textsuperscript{48} However, noting that LK03 shares the majority of its variable domains with AAV3B (and HGFR receptor usage), this conclusion is also in dispute. Well-structured trials may be needed to resolve the benefits of the current generation of engineered capsids.

**LIVER-SPECIFIC PROMOTERS**

A future goal of AAV vector capsid engineering is eventually to limit gene transfer solely to the organ of interest, in this case the liver. Until this goal is fully realized, however, efforts must focus on the expression of the therapeutic transgene, which should largely be limited to hepatocytes to improve safety and efficacy profiles. A rich body of work has been generated on liver-specific promoters (LSPs), from deconstructed single-locus cassettes defined by “promoter bashing,” to modern chimeric cassettes, to next-generation cassettes identified by deep sequencing techniques and high-throughput screening, and finally to transgene-specific regulation.

Early LSPs drew directly from work aimed at understanding the regulation of high abundance liver secreted proteins such as human serum albumin and alpha-1-antitrypsin.\textsuperscript{49,50} Seeking to tailor and control transcriptional activity further, chimeric cassettes were developed. These LSPs are essentially strings of modular building blocks that have been identified and characterized independently. Among the best-known examples of chimeric promoters are those that have been used in the hemophilia A and B trials. Those used by Kay et al. and Nathwani et al. began as a combination of the Apolipoprotein E/C-I hepatic control region combined with the human alpha-1-antitrypsin core promoter.\textsuperscript{51–54} At 754 bp, this combination, though potent, proved to be too large to fit into a self-complementary vector. What followed was a stepwise reduction in LSP size to first accommodate self-complementary FIX, called LP1, and even smaller to drive the expression of FVIII, called HLP, based upon closer analysis of the two core elements.\textsuperscript{55,56} The cassette used by the Verma et al. and Wilson et al. groups uses a different set of mod-
ules, specifically two copies of alpha 1 microglobulin/bikunin enhancer coupled to the core promoter of human thyroxine-binding globulin (TBG). Expression is further stabilized by the inclusion of a woodchuck hepatitis virus post-transcriptional regulatory element (WPRE).57–60 Finally, Naldini et al. made their promoter, designated “ET,” and described as randomly assembled hepatocyte-specific transcription factor binding sites linked to the murine transthyretin promoter.61 Though it does not drive expression as well as those that entered clinical trials, it presages next-generation cassette design, exemplified by an alternative approach employing in silico analysis of microarray data to identify liver-specific control elements. Interestingly, this approach re-identified the core enhancer of alpha-1-antitrypsin, suggesting that another avenue for engineering tissue-specific gene expression exists.62

CODON OPTIMIZATION

While restriction of gene expression to hepatocytes is critical, equally important is the property of prolonged gene expression from a single dose of vector. Prolonged gene expression, meaning expression at therapeutic levels over a time span measured in years, is a key part of the value proposition for AAV liver-directed gene therapy. The data from large animal studies with AAV vectors, particularly in hemophilia dog models, suggest that multi-year therapeutic benefit, perhaps ≥10 years, can be achieved with a single vector dose.63 Fortunately, the clinical data available to date indicate that prolonged therapeutic benefit to hemophilia B patients can also be achieved with a single vector dose.64 Obviously, more clinical results are needed before firm general conclusions can be drawn regarding the duration of therapeutic benefit from a single dose.

While selection of enhancer and promoter elements seeks to endow AAV vectors with hepatocyte selective expression, the high vector dose requirements of liver-directed gene therapy make it useful to attempt to maximize the level of gene expression per dose. Transgene codon optimization is an option to maximize expression and therapeutic potential. Though the candidate selection process for transgenes is often not reported, codon optimization appears to be the default for most vectors used in research studies and clinical trials.64–67 Briefly, the goal of codon optimization is to match the codon usage in the target therapeutic transgene with the abundance of transfer RNA (tRNA) for each codon in a particular organism and/or cell type. There are examples where this process can result in a significant increase in gene expression,68 presumably by increasing the overall rate of translation via eliminating pauses created by codons that utilize low-abundance tRNAs. Codon optimization applied indiscriminately could potentially be detrimental versus relying on native sequence. Many proteins have complex structural topologies, where translational pauses might allow proper folding and/or interaction with translocation machinery; eliminating these pauses could lead to misfolding and loss-of-function, or aggregation.69,70 However, as hepatocytes function to synthesize and secrete massive levels of protein on a continuous basis, this potential limitation may be less of a concern for liver-directed therapies.

MANUFACTURING CHALLENGES FOR AAV LIVER-BASED GENE THERAPY

The challenges of manufacturing AAV vectors at a scale and quality level appropriate for clinical trials held back product development for many years. However, significant improvements have been made in recent years, enabling the current expansion of the number and spectrum of clinical trials employing AAV vectors.71–73

Clinical and eventual commercial manufacturing of AAV vectors for liver-based gene therapy products is an especially daunting challenge due to the large dose requirements required for systemic administration to reach the desired hepatocytes. Comparison of recent gene therapy programs illustrates the challenge of liver-based gene therapy versus other targets for vector delivery. A clinical gene therapy for Leber congenital amaurosis type 2 (LCA2), for example, utilized doses of approximately 6×10^10 vg of recombinant AAV2 vector, altered to carry the human RPE65 gene (rAAV2-CBSB-hRPE65) in a volume of 150 μL per eye.74 In contrast, the high dose of recombinant AAV8 encoding FIX used in the Nathwani clinical trial used doses of 2×10^12 vg/kg, or 1.61×10^14 vg for the average 80.7 kg patient in the trial. Thus, the amount of vector for a single dose by Nathwani was on average 271 times higher than for the LCA2 trial for a single eye, or 135 times higher for both eyes. Similar differences are relevant for ear, brain, or other local only treatments versus systemic delivery for liver-directed therapies. To meet these high dose requirements for liver-directed gene therapy, suspension bioreactor platforms are under development.72,73,75–78

VECTOR DISTRIBUTION IN NONCLINICAL MODELS

The distribution and transgene expression of several vectors after intravascular delivery has been studied in mice, rabbits, dogs, and nonhuman
primates. The overall tissue distribution between various nonclinical species is generally similar. In mice, AAV8 has been demonstrated to transduce liver efficiently, with levels of transduction 10–100 times greater than early generation AAV2 vectors, and approximately 9- and 16-fold higher than AAV7 and AAV5, respectively. This is consistent with the superior liver-transducing properties of Clade E AAV vectors. In a subsequent study in mice, a targeted group of liver-specific vectors were evaluated (AAV8, AAV6.2, AAV7, AAVrh.64R1, AAVhu.37, AAVrh.8, AAVrh.32.33), with biodistribution to liver and non-hepatic tissues as a criterion. Of the vectors tested, AAV8, AAVrh.8, and AAVhu.37 demonstrated the greatest future potential based on liver transduction and durability of expression. In terms of biodistribution to extrahepatic tissues (heart, lung, kidney, spleen, pancreas, brain, muscle, and testes), AAV6.2 and AAVrh.32.33 had the highest levels of extrahepatic vector, with AAV8, AAV7, AAVrh.64R1, AAVhu.37, and AAVrh.8 having similar levels of significantly lower levels in non-liver tissues.

There are a limited number of studies that directly compare results of AAV vectors between rodents and non-rodents, but these can provide important information to understand differences between species and potential allometric scaling. The biodistribution to liver and expression from AAV2, AAV5, AAV6, and AAV8 vectors encoding FVIII were evaluated in hemophilia A mice and dogs. Again, AAV8 had dramatically higher levels of transduction in the livers of mice, approximately twofold higher than AAV6, sevenfold higher than AAV2, and 100-fold higher than AAV5. The relative expression of hFIX between vectors showed a similar pattern, albeit with a slightly reduced magnitude of difference between serotypes. In comparison, in hemophilia A dogs, the levels of transduction and expression of canine FVIII was similar between each vector, each producing approximately 2–5% of normal canine FVIII activity, in contrast to the results in mice, which varied from 20% for AAV2, 50% for AAV6, and 110% for AAV8.

In preclinical work for hemophilia B, AAV8 also produced the highest level of FIX in C57BL/6 mice, approximately 17- to 84-fold higher than AAV2 and AAV5 vectors encoding hFIX after administration of a given dose, with the same pattern of transduction levels for each vector. In rhesus monkeys, animals treated with AAV8 had levels of hFIX that were approximately twofold higher than AAV5 at steady state.

In both dogs and monkeys, the differences in transduction efficiency between different serotypes is less pronounced than in mice, with a lower transduction efficiency overall. The partitioning of vector in tissues of AAV8 in rhesus monkeys is similar to that of mice and rabbits, however, with the liver having the highest level, at least 100-fold more than other tissues, and vector detected in all tissues examined, including the brain, testes, muscle, kidneys, lungs, heart, and spleen (in ascending order). A robust data set comparing the efficiency of gene transfer of the same AAV vector product between monkeys and humans is lacking, but data from ongoing clinical trials should begin to fill this gap.

LOCALIZATION OF HEPATOCYTE GENE TRANSFER

Although nonclinical and now clinical experiences in hemophilia offer important insights into liver gene therapy of secreted proteins, gene therapy for non-secreted proteins brings about a separate set of preclinical challenges. As mentioned previously, transduction rates and transgene expression can vary between species, and so can the site of vector predilection of cell transduction within the liver. In a 2011 paper, Bell et al. compared the expression pattern of AAV8 expressing green fluorescent protein (GFP) in liver between mice, dogs and nonhuman primates. They demonstrated a predominantly pericentral expression of GFP in both murine and canine liver, regardless of whether a liver specific promoter (TBG) or a ubiquitous promoter (CMV) was used. The expression pattern was also independent of the site of vector administration, with pericentral expression occurring in dogs after injection in either the cephalic vein or the hepatic artery. Transduction within the liver of adult rhesus and cynomolgus macaques, in contrast, occurred mainly in periportal regions. Interestingly, livers from mice and rhesus monkeys treated with AAV8 GFP as neonates or infants showed a more uniform distribution of GFP expression, lacking a predilection for either periportal or central venous areas. Transduction of neonatal animals, however, often leads to loss of transgene expression over time, due to the diluted effects of replicating hepatocytes. A more recent paper demonstrated that localization in mice can be capsid dependent, with AAV2 localizing to periportal regions, while AAV 7, 8, 9, and rb10 localized to pericentral regions, as seen previously.

The reason for the differences in localization between species is not known. However, it is speculated that it may be due to receptor localization and abundance, presence or absence of Kupffer cells, or differences in sinusoidal width between species. It is interesting to note that both hydrodynamic
injection of plasmids, as well as injection of dyes such as Evans Blue, result in accumulation of these substances within central venous regions of murine livers. The localization of these compounds within nonhuman primate livers is not known. Further studies are needed to understand the mechanism of localization for these viral vectors in animal models and how they compare to human patients.

Although less important for the production of secreted proteins such as FIX or FVIII, localization of vector transduction within the liver may be significant in the treatment of inborn errors of metabolism. Using gene therapy for restoring metabolic function, as in the case of ornithine transcarbamylase deficiency (OTC) or methylmalonic acidemia (MMA), is well summarized in the 2008 review by Alexander et al. Treating these disorders in human patients will likely require a higher percentage of transduction of hepatocytes than for secreted proteins, as well as careful consideration of hepatocyte zonality. It is well known that the liver is divided into functional zones, and although the secretion of many proteins from hepatocytes does not appear to be zonally specific, many metabolic processes, including ammonia detoxification, are localized to specific regions of hepatic architecture. The effects of enhancing or disrupting this pattern on restoring metabolic function will need to be further explored in the context of AAV gene therapy as these treatments move toward the clinic.

**PRECLINICAL IMMUNE RESPONSES TO AAV GENE THERAPY**

As mentioned in previous sections, early work with AAV in liver-directed transduction was hindered by unexpected immune responses to AAV, namely in the form of neutralizing antibodies and cytotoxic T-cell responses to the AAV capsid that were not predicted by preclinical studies, highlighting one of the major preclinical challenges when working with AAV. To date, an animal model completely predictive of the immune response seen in humans to this virus has yet to be discovered.

Both mice and nonhuman primates have proven to be successful models for the effect of neutralizing antibodies (NAbs) on transduction. Studies in both species have demonstrated that NAb titers as low as 1:5 can block AAV vector transduction of the liver completely, and that vector remains susceptible to antibody-mediated neutralization for several hours after intravascular delivery. Due to the strong conservation in capsid proteins between AAV serotypes, further work needs to be done to examine the cross-reactivity of antibodies and the potential for re-dosing patients with alternative serotypes in the future.

Although animal models predicted many aspects of the human immune response to transgene products, they largely failed to predict responses to AAV capsid. The first indication that human CD8+ T cells could limit efficacy of gene transfer was uncovered in the initial trial of AAV gene transfer to liver for hemophilia. In this trial, AAV2 was used to deliver human FIX under the control of a liver-specific promoter. Initially, the first subject infused at the high dose, 2×10^{12} vg/kg, expressed FIX at levels in the range of 10–12%, concurrent with what had been seen in the dog model of hemophilia. After a period of several weeks, however, FIX expression disappeared, accompanied by a self-limited and asymptomatic transaminase elevation in the blood. This series of events, not observed in the experimental animal models, spurred a large-scale effort to determine the underlying cause of this loss of expression. Ultimately, it was hypothesized that declining transgene expression was due to anti-capsid cytotoxic T-cell mediated destruction of transduced hepatocytes displaying capsid antigen via major histocompatibility complex class I pathways.

Multiple efforts have been made to develop preclinical models of the human immune response to the AAV capsid. In early studies, mice were immunized with AAV capsid protein or viral vector and then exposed to AAV expressing FIX. Despite the presence of cytotoxic CD8+ T cells specific for the AAV capsid protein within the liver, hepatocyte production of FIX was preserved to the same levels as seen in naive mice. Further studies in mice confirmed that although AAV did induce a capsid-specific cytotoxic CD8+ T-cell response, transduced hepatocytes did not become cytolytic targets. Recent attempts to boost this immune response in animals, including the use of adjuvants or incorporation of SIINFEKL peptide epitopes into the AAV vector capsid concurrent with the use of OT-1 mice, have recapitulated some of the immune response, but the utility and relativity of such contrived circumstances to administration in humans is questionable. Work done in nonhuman primates has not met with any additional success, and researchers have been unable to mimic reliably the cytolytic T-cell response and subsequent rise in alanine aminotransferase and aspartate aminotransferase to date. The reason for these species-specific differences is unknown. However, speculation suggests it may be due to the loss of a subset of inhibitory immunoglobulin-like lectins on human T cells during evolution. Further proof of this remains to be demonstrated, however.
LIVER-MEDIATED TOLERANCE

Although the liver is responsible for making many pro-inflammatory components of the immune system, it also plays a critical role in inducing tolerance to dietary and self-antigens. This tolerance induction is especially intriguing in the context of liver-directed gene therapy. Pastore et al. first described the phenomenon of limitation of immune responses to an otherwise immunogenic protein by use of a promoter sequence derived from a liver-specific gene.93 Direct comparison of two adenovirus vectors expressing human alpha 1-antitrypsin (hAAT) from the ubiquitous mouse phosphoglycerate kinase 1 (mPGK) promoter or from the liver-specific albumin promoter indicated that C3H/HeJ mice produced antibodies to hAAT when it was expressed from the mPGK promoter but not when it was expressed from the albumin promoter.93 Since then, a number of studies have demonstrated the induction of tolerance to different proteins following in vivo hepatocyte transduction and transgene expression. Work by Cao et al. demonstrated a role for regulatory T cells in liver tolerance.94 The authors used Rag2−/− BalbC mice transgenic for a T-cell receptor reactive against a peptide from chicken ovalbumin (OVA). These mice lack regulatory T cells normally. However, upon administration of AAV-OVA, they develop tolerance to the OVA transgene, accompanied by formation of regulatory T cells expressing CD4+CD25+FOXP3+. A role for regulatory T cells in induction of tolerance has been demonstrated in nonhuman primates as well. A paper from Mingozzi et al. demonstrated that the administration of the anti-interleukin-2 receptor antibody daclizumab, in the context of an immunosuppressive regimen in nonhuman primates administered AAV-FIX, led to decreased tolerance of the transgene versus mycophenolate mofetil (MMF) and sirolimus alone.95 The authors went further to discover that daclizumab inhibited the formation of CD4+ CD25+FOXP3+ T cells, which are essential to the development of liver tolerance to FIX.95

In addition to regulatory T cells, Kupffer cells and liver sinusoidal endothelial cells (LSECs) have also been shown to contribute to development of tolerance, potentially by the secretion of immunomodulatory factors such as nitric oxide and IL-10.96,97 Mouse LSECs function as a specialized form of antigen-presenting cell and express receptors that promote antigen uptake, including the mannose receptor and the scavenger receptor, and molecules that promote antigen presentation, including CD40, CD80, and CD86. Although mouse LSECs can prime both CD4 and CD8 T cells, the most common consequence of this is tolerance induction versus activation.96 The reasons for this are still unclear, as is whether this translates to human LSECs. What is clear, however, is that the presence of specialized T cells, in conjunction with Kupffer cells and LSECs, establishes a unique locally suppressive microenvironment, contributing to the development of tolerance to transgenes expressed in the liver.98,99

Although transduction of the liver versus other organs may lead to advantages in establishment of tolerance to gene therapy vectors and transgene production,100,101 considerations of transgene immuno- nity are still necessary, especially in cases of proteins of high immunogenicity, such as FVIII. The immunogenicity of a delivered transgene depends on a number of factors, including vector selection, route of administration, targeted transgene expression levels, and the genotype of the host.102 Despite the complexity of attempting to suppress immune responses to therapeutic transgenes while avoiding unwanted side effects, pharmacological interventions are still being pursued. Sack et al. explored the use of B-cell depletion in a model of AAV mediated FVIII gene delivery with some success, although the level of FVIII expression from different vectors was a confounding factor.103 Other pharmacological interventions include cyclophosphamide in a null mutant canine model of hemophilia B,104 as well as cyclophosphamide and methylprednisolone in murine models of hemophilia B.105 Immune responses to therapeutic proteins are not limited to gene therapy, in that development of anti-drug antibodies is also an issue in the protein therapeutics realm. Ellinwood et al. successfully employed a combined immunosuppression regime of mycophenolate mofetil plus cyclosporine to treat Sanfilippo and Hurler disease in affected dogs carrying null mutations of the relevant genes, and a relatively short course of a widely prescribed immunosuppressive drug, methotrexate, has been successfully employed in animal models of Pompe disease and in Pompe patients.106–108 Through studies such as these, an armamentarium of immunosuppressive drugs and protocols potentially useful in the context of gene therapy is being assembled and may prove useful in future gene therapy work.

REMAINING CLINICAL CHALLENGES FOR LIVER-BASED AAV GENE THERAPY

Assessment of the remaining challenges for broad commercial application of AAV gene therapy vectors across the full spectrum of liver-based diseases can clearly be debated. Two commonly discussed challenges have been selected for discussion.
below: re-administration of vector and risk of hepatocellular carcinoma.

### AAV vector re-administration

It is well-appreciated that systemic administration of AAV vectors to seropositive individuals will be blocked by the presence of neutralizing antibodies in subjects. It is therefore understood that simple re-administration of the same AAV vector product will not be successful due to blocking neutralizing antibodies, and there are ample data to suggest that re-administration of closely related AAV vector products will also not be successful. Nonetheless, the issue of vector re-administration is an important one and is the subject of active research. Three different avenues for re-administration have been published: capsid switching, removal of antibodies from serum, and administration of decoys to adsorb antibodies. An example of capsid switching relevant to the discussion of hemophilia gene therapy was published by Wang et al. In this study, hemophilia B dogs were given an intravenous dose of AAV2 FIX that resulted in modest therapeutic benefit. To test the capsid switch re-administration hypothesis, a second dose of an AAV FIX vector, this one packaged in an AAV8 capsid, was given. This second dose resulted in a dramatic improvement in FIX expression and correction of the bleeding disorder, confirming that it is possible to re-dose animals with alternative serotypes of AAV.

Antibody removal or reduction to enable AAV re-administration was tested using the process of plasmapheresis. These studies were conducted in rhesus macaques with AAV vectors encoding a microdystrophin gene. Two cycles of plasmapheresis were shown to reduce serum antibodies in animals with high titers to levels equivalent to animals in which only low levels of antibodies were detected. Importantly, gene transfer to muscle was blocked in animals with high antibody titers but was successful in animals that had undergone the plasmapheresis procedure.

Finally, Mingozzi et al. confirmed the ability of co-administration of empty AAV capsids to serve as decoys to block the action of existing neutralizing antibodies temporarily. A large excess of empty capsids (~100-fold) was able to allow gene transfer to the liver with the cognate capsid while also not blocking gene transfer by competition with cell surface receptors. The same results were obtained in cell-based assays and in vivo in mice and monkeys. Obviously each of these potential avenues for vector re-administration will have to be confirmed in clinical studies before coming into general use.

### Risk of hepatocellular carcinoma

Multiple studies have been conducted examining the potential effects of AAV integration on the development of hepatocellular carcinoma (HCC), a correlation currently hotly debated within the gene therapy community. This potential risk was first highlighted by Donsante et al. in 2001, utilizing recombinant AAV (rAAV) gene therapy to treat mucopolysaccharidosis type VII in newborn betaglucuronidase (GUSB) deficient mice. The authors reported an increased incidence of HCC in mice treated with rAAV compared with untreated controls. Their initial conclusion, based on quantitative polymerase chain reaction analysis of tumor samples, was that tumorigenesis was unlikely related to integration of rAAV due to the low numbers of vector genome copies in tumor tissue. However, a follow-up study of tumors from these mice identified four examples of AAV integration sites within the delta-like homologue 1-deiodinase type 3 (Dlk1-Dio3) region. Two of these insertions were within MIR341, an insertional location mirrored in later studies as well. It is important to note that there is no human orthologue to MIR341. However, upregulation of other transcripts within the orthologous Rian region in humans have been implicated in a poor prognosis for patients with HCC.

A study conducted in 2006 on the utility of gene therapy in the treatment of ornithine transcarbamylase (OTC) deficiency examined the frequency of HCC in spf ash, a model of this disease, and B6C3F1 control mice. Animals were treated with various AAV serotypes (2, 7, 8, and 9) encoding murine OTC (mOTC) or a control vector, AAV2/8-LacZ. All experimental groups developed liver nodules. However, mice that received AAV with mOTC as the transgene showed lower levels of nodules than either untreated control animals or animals treated with AAV-LacZ. This suggests that administration of AAV-OTC and correction of metabolic abnormalities with gene therapy may have actually been protective against the development of HCC in this mouse strain. This would not be surprising considering the metabolic alterations that occur in hepatocytes with disrupted urea cycles.

In a follow-up study, liver nodules or tumors from these same animals receiving AAV-mOTC, as described above, were collected and analyzed for integration sites. Mice were chosen with the highest numbers of vector DNA copies in the liver, and integration sites were compared between tumor and normal liver sections from five animals. Integration sites were distributed along the length of each chromosome without obvious clustering in hot spots. Integration sites were generally enriched...
in gene-dense regions and near gene 5’ ends, compared with computationally random distributions. A total of five sites in the Dlk1-Dio3 region could be verified, one from normal tissue and four from tumor-nodule tissue. In all four cases, integration was closest to the Rian locus. A recent paper from Chandler et al. further examines the role of AAV integration in the formation of HCC in mice. Similar to previous studies, it was demonstrated that AAV most frequently integrates into highly expressed genes, including albumin and α-fetoprotein. Integration of AAV into the Rian locus was the only integration site associated with increased development of hepatocellular carcinoma, and only under a restricted set of conditions. First, HCC formation was dose dependent. A statistically significant increase in HCC formation was observed at $1 \times 10^{14}$ GC/kg (84%), but not at doses $\leq 7 \times 10^{12}$ GC/kg (12%). Second, formation of HCC was highly dependent on the age at inoculation. Neonatally inoculated mice developed increased HCC, but mice inoculated at $\geq 6$ weeks of age did not develop tumors above background. This is consistent with previous studies, which found increased risk of HCC in neonatally inoculated mice but not in older mice. Third, development of tumors appeared to be related to specific promoter/enhancer combinations. There was a statistically significant increase in tumorigenesis with AAV vectors encoding the TBG or chicken B-actin enhancer and promoter combination when compared with other promoters, including α-1 antitrypsin (AAT). Similar to previous studies, the majority of integration sites within Rian were located in MIR341, a gene lacking an orthologue in humans. No microhomology between any AAV vector genome and the integration sites in MIR341 could be detected, making it difficult to translate this finding to the human genome. In this study, there was no statistically significant difference in the incidence of HCC between AAV8 containing the transgene of interest (MUT) when compared to AAV8 encoding GFP. These results, along with similar data from other publications, indicate that the development of HCC is likely not associated with a particular transgene, although integration specificity due to transgene sequence cannot be ruled out.

Translation of data from mice to humans can often be difficult. A recent publication demonstrated an association between wild-type (wt) AAV2 insertional mutagenesis, the dysregulation of neighboring genes, including cancer driver genes CCNA2, TERT, CCNE1, TNFSF10, and KMT2B, and development of HCC in humans. The authors demonstrated the presence of wtAAV2 sequences within 11/193 human HCC biopsies. The relationship of the presence of AAV DNA to the development of carcinoma is not clear, however, as 5/11 biopsies were from patients with other well-known HCC driving conditions, including alcoholism and hepatitis B and C infections. An additional four biopsies presented with AAV-unrelated mutations that have previously been linked to HCC formation. Therefore, only two biopsies contained fragments of wtAAV2 sequences with no other clear driver for mutagenesis. In addition, more non-tumor samples were positive for AAV2 DNA than tumor samples in this study. However, the number of recovered reads in tumor tissue was higher. Unlike in the murine studies, no integration was noted at or near any gene within the Rian locus. This suggests that the specific locations for integration of AAV into the human genome likely vary from the locations within the mouse genome. The author’s conclusion that AAV is associated with an increased risk of HCC is hotly disputed, with a number of rebuttals quickly following publication, arguing that the presence of this ubiquitous virus is simply association and not causation.

The relevance of this study to the safety of AAV-mediated gene therapy is not clear, as 30–50% of the human population is seropositive for AAV2 and yet AAV2-induced hepatocellular carcinoma is not widespread, and only a small fraction (5.7%) of the examined HCCs were positive for AAV2 in this study. When comparing data between wtAAV and rAAV vectors, two things must be considered. The first is that rAAV vectors lack wild-type rep proteins, and although it has been demonstrated that integration can still occur, it is at lower frequency compared with wild-type viruses. The second is that conversion of wtAAV to rAAV vectors involves the deletion and rearrangement of portions of the viral genome to make room for insertion of transgenes. Recent unpublished work from the lab of Ian Alexander demonstrates the presence of strong promoters within wtAAV2 ITRs, in regions that are largely absent from many AAV gene therapy vectors.

Unfortunately, the total risk of development of hepatocellular carcinoma secondary to AAV gene therapy in humans is still unknown, but, to date, no genotoxic events have been reported in the >130 rAAV clinical trials conducted so far. Kaeppel et al. performed integration site analysis in the muscle of patients from their lipoprotein lipase trial and discovered very low integration frequency and random nuclear integration. Additionally, a recent publication by Gil-Farina et al. examining liver samples from nonhuman primates in nonclinical studies and human patients in a clinical trial for acute intermittent porphyria demonstrates
that although rAAV did integrate into the human genome, it did so with extremely low frequency, and no integration clusters were found in genes previously reported to link AAV and hepatocellular carcinoma development. Further assessment of risk will likely be an evolving theme as clinical trial data accumulates from next-generation AAV gene therapy vectors.

SUMMARY

AAV gene therapy for the liver has taken some time to come of age, with gene transfer research in hemophilia B as the forerunner of liver-directed gene therapy more broadly. Animal models for hemophilia B existed or were created, providing valid test subjects for therapeutic benefit. Assays existed for quantitative measurement of FIX, and the correlations of FIX level with therapeutic benefit in humans were well-established. The FIX gene is small, removing barriers for AAV vector engineering, and the field of mouse transgenics had provided basic information regarding control of gene transcription within hepatocytes. The stage was set for hemophilia B gene therapy proof-of-concept in animals, setting in motion the approach to human proof-of-concept for hemophilia B gene therapy. The final barriers that slowed progress to the clinic were cGMP vector manufacturing and the parallel regulatory requirements. With diligent work across the field, these barriers too were overcome.

Early clinical successes were limited, but importantly shed light on areas that required subsequent translational research, especially in the realm of immune responses to vector capsid proteins. While detailed hypotheses have been proposed as to mechanisms of the observed limitations of vector “expression,” no iron-clad proof has been elucidated. This is not all that surprising, since the phenomena have not been recapitulated in animal models and the number of clinical examples is limited. One could conclude that advances in AAV vector capsid technology have likely played an important role in clinical successes. There is an enormous foundation of vectorology, animal science, and now clinical successes that potentially point to a brighter future for patients.

We should not be surprised nor sheepish about the duration of this journey to date. Consider the profound influence on medicine that could be wrought if gene therapy success was brought to an ever-increasing number of diseases. Today, there are multiple clinical trials underway in hemophilia and other liver-based disorders (see Table 1), as

| Trial Sponsor          | Indication                      | AAV serotype | Stage          | NCT Identifier |
|------------------------|--------------------------------|--------------|----------------|----------------|
| Audentes Therapeutics  | Crigler Najjar                 | AAV8         | IND-enabling   | NCT02576795    |
| BioMarin               | Hemophilia A                   | AAV5         | Phase I/II     | NCT02618915    |
| CIMA/Aligen            | Wilson disease                 | AAV8         | ND             | NA             |
| Dimension Therapeutics | Hemophilia B                   | AAVrh10      | Phase I/II     | NCT02618915    |
|                        | OTC deficiency                 | AAV8         | IND-enabling   | NA             |
|                        | GSDla                          | AAV8         | IND-enabling   | NA             |
|                        | Wilson disease                 | ND           | DC             | NA             |
|                        | PKU                            | ND           | DC             | NA             |
|                        | Citrullinemia type 1           | ND           | DC             | NA             |
| Dimension/Bayer        | Hemophilia A                   | ND           | IND-enabling   | NA             |
| Genethon               | Crigler Najjar                 | ND           | Preclinical    | NA             |
| REGENXBIO/PENN         | HoFH                           | AAV8         | Phase I/II     | NCT02651675    |
| Sangamo                | Hemophilia A                   | AAV6         | IND-enabling   | NA             |
| Selecta Biosciences    | OTC deficiency                 | AAV          | Preclinical    | NA             |
|                        | Methylmalonic acidemia         | Anc80        | Preclinical    | NA             |
| Shire                  | Hemophilia B                   | AAV8         | Preclinical    | NA             |
|                        | Hemophilia B                   | AAV8         | Discontinued   | NCT0187608    |
|                        | Hemophilia A                   | AAV8         | IND-enabling   | NA             |
| Spark Therapeutics/Pfizer | Hemophilia B                | ND           | Phase I/II     | NCT02484092    |
| Spark                  | Hemophilia A                   | ND           | IND-enabling   | NA             |
| Spark Therapeutics/CHOP | Hemophilia B                  | AAV8         | Phase I/II     | NCT01620801    |
| UCL/SJCRH              | Hemophilia B                   | AAV8         | Phase I/II     | NCT00978238    |
| uniQure/Chiesi         | Hemophilia B                   | AAV5         | Phase I/II     | NCT02398342    |
| uniQure                | Hemophilia A                   | AAV5         | Preclinical    | NA             |
| uniQure/Digna          | AIP                            | AAV5         | Complete       | NCT02082860    |

ND, not disclosed; CIMA, Center for Applied Medical Research University of Navarra; HoFH, homozygous familial hypercholesterolemia; CHOP, Children’s Hospital of Philadelphia; UCL/SJCRH, University College London/St. Jude Children’s Research Hospital; OTC, ornithine transcarbamylase deficiency; GSDla, glycogen storage disease type Ia; PKU, phenylketonuria; AIP, acute intermittent porphyria; IND, investigational new drug; DC, development candidate; NA, not applicable.

Sources: company websites, clinicaltrials.gov, Thomson Cortellis 2016.
well as other target organs. Definitive assessments of the results of ongoing clinical trials will have to await publication of results in peer-reviewed publications, but as a field we should expect to witness successes.

As we expand beyond the realm of secreted protein therapeutics characterized by hemophilia, additional aspects of liver gene transfer become important. For example, the importance of which hepatocytes undergo gene transfer becomes paramount. Due to metabolic zonation of the liver, hepatocytes proximal to the portal triad are active in metabolic processes such as the urea cycle, while those hepatocytes adjacent to the central vein are specialized for other functions. Thus, gene transfer to hepatocytes involved in metabolic syndromes is likely more critical than gene transfer across the liver. Endpoints for assessing success of gene transfer also become more challenging than the simple measurement of FIX in the blood. Fortunately, there are AAV vector capsids that do efficiently transduce metabolically active hepatocytes, and for many metabolic diseases, blood-based biomarkers and endpoints are already available.

We should not be surprised or overwhelmed by the opportunities for improvements across the field: vector design, manufacturing operations, adjunct therapies. This scenario would be expected and familiar given the field of hybridomas leading to therapeutic antibodies, where discoveries are still being made at a fast pace. The field is ripe for application of today’s technology for tomorrow’s therapies while pursuing technological improvements that may fuel expansion of gene therapies to new and more complex therapeutic indications. It is probably an apt paraphrase to say that we are now at “the end of the beginning” of this amazing and important enterprise.

AUTHOR DISCLOSURE

The authors declare that they have no potential competing financial interests. K.C. and S.W. are shareholders in Dimension Therapeutics. L.K., C.T., L.S., D.G., T.W., K.C., and S.W. are employees of Dimension Therapeutics.

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