Sustained high-level expression of human factor IX (hFIX) after liver-targeted delivery of recombinant adeno-associated virus encoding the hFIX gene in rhesus macaques


The feasibility, safety, and efficacy of liver-directed gene transfer was evaluated in 5 male macaques (aged 2.5 to 6.5 years) by using a recombinant adeno-associated viral (rAAV) vector (rAAV-2 CAGG-hFIX) that had previously mediated persistent therapeutic expression of human factor IX (hFIX; 6-10% of physiologic levels) in murine models. A dose of 4 x 10^12 vector genomes (vgs)/kg of body weight was administered through the hepatic artery or portal vein. Persistence of the rAAV vgs as circular monomers and dimers and high-molecular-weight concatamers was documented in liver tissue by Southern blot analysis for periods of up to 1 year. Vector particles were present in plasma, urine, or saliva for several days after infusion (as shown by polymerase chain reaction analysis), and the vgs were detected in spleen tissue at low copy numbers. An enzyme-linked immunosorption assay capable of detecting between 1% and 25% of normal levels of hFIX in rhesus plasma was developed by using hyperimmune serum from a rhesus monkey that had received an adenooviral vector encoding hFIX. Two macaques having 3 and 40 rAAV genome equivalents/cell, respectively, in liver tissue had 4% and 8% of normal physiologic plasma levels of hFIX, respectively. A level of hFIX that was 3% of normal levels was transiently detected in one other macaque, which had a genome copy number of 25 before abrogation by a neutralizing antibody (inhibitor) to hFIX. This nonhuman-primate model will be useful in further evaluation and development of rAAV vectors for gene therapy of hemophilia B.

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Introduction

Recombinant adeno-associated viral vectors (rAAVs) hold great promise for the treatment of hemophilia B. Preclinical studies in murine and canine models of hemophilia B have demonstrated persistent therapeutic expression of factor IX (FIX) leading to correction of the bleeding phenotype after intramuscular or portal vein administration of rAAV.1-3 Liver-targeted delivery of rAAV resulted in significantly higher levels of FIX than observed after intramuscular administration of an equivalent dose of vector in immunodeficient mice.4 More important, the development of neutralizing antibodies (NABs) to human FIX (hFIX) is more common after intramuscular injection of rAAV than after intraportal administration of the same dose of vector.5-11 Mice with persistent hFIX expression after liver-targeted delivery of rAAV could not mount a substantial antibody (Ab) response when challenged repeatedly with hFIX antigen in Freund adjuvant, a result suggesting that the liver has a unique ability to induce immune tolerance to neoantigens.12 Endothelial cells of the liver, which can efficiently process and cross-present exogenous soluble antigens to CD8+ T cells, are intimately involved in antigen-specific immune tolerance after liver-targeted gene transfer with rAAV vectors.13

The findings of sustained expression of hFIX at therapeutic levels after a single bolus injection of rAAV vector into the portal vein of mice1-6,8 and the absence of toxicity after liver-directed delivery of high titers of rAAV vector in murine and canine models of hemophilia led to initiation of a phase I-2 clinical study. This study, however, has raised concerns about germline transmission because infusion of 2 x 10^11 vector genomes (vgs)/kg of body weight into the hepatic artery resulted in detection of rAAV vgs in the semen of the first adult volunteer with hemophilia B.14 Although there was no evidence of transduction of spermatocytes with this low dose of vector particles, therapeutically relevant doses of AAV may lead to germline transmission. The absence of vgs in the semen of hemophilic dogs given an intrahepatic bolus of a 10- to 30-fold higher dose of rAAV than that received by the first adult volunteer15 calls into question the value of preclinical safety data derived from murine and canine models. The biodistribution of the vector in these laboratory animals may differ markedly from that in humans, a natural host for the virus from which the vector was derived.

Although hemophilia has not been observed in rhesus macaques, these monkeys, like humans, are a natural host for AAV and...
macaque models thus offer a unique opportunity for evaluating the biodistribution, gene-transfer efficiency, and toxicity of rAAV in the context of preceding natural sensitization to AAV. Results of studies in macaques have consistently predicted clinical outcomes in humans after gene transfer with adenoviral and retroviral vectors. In addition, macaques have a considerably longer lifespan than the commonly used laboratory animals, an important asset for evaluating long-term safety and durability of expression after rAAV-mediated gene transfer. However, difficulties in distinguishing native rhesus FIX from the highly homologous human cognate have impeded use of macaques for preclinical evaluation of rAAV vector–mediated expression of FIX. We overcame this difficulty to demonstrate stable, high-level transduction of rhesus liver tissue leading to expression of hFIX at therapeutic levels in rhesus plasma for more than a year. Our studies also provided data on the biodistribution and shedding of vector particles in biologic fluids and long-term transduction of distant organs in a context that is relevant to humans.

Methods

Production and purification of rAAV-hFIX vector

The pAAV CAGG-FIX plasmid encodes hFIX complementary DNA (cDNA) under the control of the cytomegalovirus enhancer, β-actin promoter complex that was used to generate the rAAV CAGG-FIX vector. AAV vector preparations for the studies in rhesus macaques were made by scaling up our previously described adenosine-free transient transfection method. Briefly, subconfluent 293T cells either in cell blocks (Nunc, Roskilde, Denmark) or plated on 15-cm plates were cotransfected with pAAV CAGG-FIX vector plasmid, an adenosive helper plasmid (80-XX6), and a packaging plasmid (XX2) by using the calcium phosphate precipitation method. Cells were harvested 50 to 60 hours after transfection, lysed, and incubated with nuclease before purification by heparin affinity-chromatography. Standard slot-blot analysis was used to determine the vector particle titer. Contamination with wild-type AAV was determined by a polymerase chain reaction (PCR) assay. Aliquots of rAAV were periodically subjected to polyacrylamide gel electrophoresis (PAGE). The vector stocks were consistently free of contamination with wild-type AAV or adenoviral and cellular proteins.

Studies in animals

All animals were treated according to the standards described in the Guide for Care and Use of Laboratory Animals (National Institutes of Health publication 85-23). All procedures were performed in accordance with institutional guidelines under protocols approved by the Institutional Biosafety Committee and the Institutional Animal Care and Use Committee at St Jude Children’s Research Hospital and the University of Tennessee, Memphis. Six captive-bred male Macaca mulatta aged 2.5 to 6.6 years and weighing 1.6 to 7.6 kg were purchased from Covance Research Products (Alice, TX) and housed in the dedicated primate facility at the University of Tennessee Health Science Center. The same total dose of rAAV CAGG-FIX vector was administered into the portal venous circulation of 2 other macaques either as a bolus infusion (monkey 4) or in 2 fractionated doses of 2 × 10^12 vgs/kg (monkey 5). A limited laparotomy through a midline abdominal incision was performed in macaques anesthetized with isoflurane to gain access to the superior mesenteric vein, into which vector was infused slowly through a 27-gauge needle. The fractionated infusion of rAAV in monkey 5 required 2 separate laparotomy procedures through the same midline abdominal incision. All animals recovered well after abdominal surgery.

To provide quality assurance of the high-titer rAAV CAGG-FIX stock prepared for each macaque, 4 × 10^12 vgs derived from each of these vector stocks was injected into the portal vein of C57Bl/6j severe combined immunodeficient mice and shown to have equivalent biologic activity on the basis of the level of hFIX expression, as observed previously.

Enzyme-linked immunosorbent assay (ELISA) for quantitating hFIX in rhesus plasma

We initially used an ELISA method described by Lozier et al. to detect hFIX in rhesus plasma. This method, which relies on commercially available polyclonal Abs against hFIX, was unable to reliably detect levels of hFIX in rhesus plasma below 10% of physiologic levels because of high background cross-reactivity with endogenous rhesus FIX. We therefore developed an alternative assay using antiserum from a macaque (RQ1305) in which high titers of anti-hFIX Abs developed after adenovirus-mediated gene transfer of a vector encoding the hFIX gene. This antiserum specifically neutralized hFIX activity (> 30 Bethesda inhibitor assay units [BAUs]) without significantly impairing the function of rhesus FIX.

Flat-bottomed, 96-well plates (Dynatech, Chantilly, VA) were coated with 50 μL of a 1:50 dilution of the rhesus antiserum, left overnight at 4°C, washed with PBS containing 0.05% Tween 20 (PBST), and blocked with 200 μL/well 5% bovine serum albumin (BSA; Sigma, St Louis, MO) in PBST during a 1-hour incubation at 37°C. Standards were made by using serial dilutions of rhesus plasma spiked with affinity-purified hFIX (starting concentration, 5000 ng/mL; Sigma). Rhesus samples and standards were diluted 1:5 in PBST containing 2% BSA, and 50 μL was loaded in duplicates. After a 2-hour incubation at room temperature, the plates were washed and incubated for an additional hour with 100 μL horseradish peroxidase (HRP)–conjugated goat anti-hFIX polyclonal secondary Ab at a final concentration of 1 μg/mL (Affinity Biologicals, Ancaster, ON, Canada). After a final wash step, plates were developed with o-phenylenediamine dihydrochloride peroxidase substrate (Sigma) and the optical density was assessed spectrophotometrically at 492 nm. Background cross-reactivity from endogenous rhesus FIX was eliminated by coating the plate with a 1.5 dilution of naive rhesus plasma.

Western blot analysis to detect hFIX in rhesus plasma

Rhesus plasma from macaques in the experimental and control groups was subjected to the barium citrate precipitation method described previously to isolate the vitamin K–dependent proteins. Approximately 5 μg precipitated protein was loaded on a 10% sodium dodecyl sulfate–PAGE gel and transferred to a nitrocellulose membrane after electrophoresis. The blots were probed with a 1:50 dilution of rhesus antiserum (from macaque RQ1305) in Tris-(tris(hydroxymethyl)aminomethane)–buffered saline with 0.05% Tween 20 containing 2% BSA. After washing, the blots were exposed to an HRP-conjugated rabbit antirhesus IgG (1:1,000 dilution; Sigma) secondary Ab. The signal was developed with an enhanced chemiluminescence kit (Amersham Biosciences, Arlington Heights, IL). Dilutions of affinity-purified hFIX (Sigma) in rhesus plasma served as a positive control.
Molecular studies to evaluate rAAV gene-transfer efficiency in macaques

Wedge-section liver biopsy specimens were obtained from both lobes of the liver between 3 and 12 months after liver-targeted delivery of vector in macaques. Biopsy specimens were also obtained from the spleen, kidney, and testis of monkey 3 at 11 months after fractionated-dose infusion of vector into the hepatic artery. All biopsies were performed with the macaques under isoflurane anesthesia. A portion of the tissue sample was frozen in liquid nitrogen and then pulverized with a mortar and pestle.

Hematologic and clinical chemistry analyses

Samples for complete blood counts (CBCs), serum chemistry analyses (blood urea nitrogen and liver-function tests), coagulation-profile determinations (prothrombin and activated partial thromboplastin time [APTT]), and serum interleukin 6 (IL-6) assays were obtained every 48 hours for 2 weeks after vector infusion to assess toxicity. Thereafter, these variables were assessed monthly. The CBCs, serum chemistry analyses, and coagulation-profile determinations were performed by Antech Diagnostics (Southaven, MS). Serum IL-6 levels were assessed with an IL-6 immunoassay kit (R&D Systems, Minneapolis, MN) used according to the manufacturer’s instructions.

Detection of anti-hFIX Abs

Plasma samples from the macaques were screened for the presence of Ab against hFIX by using an ELISA described previously. Briefly, plates were coated with affinity-purified hFIX protein diluted to a concentration of 1 µg/mL with 0.1 M sodium bicarbonate buffer, left overnight, washed, and blocked for 1 hour at 37°C with PBST containing 6% BSA. After washing with PBST, 50-µL dilutions of rhesus plasma (1:50, 1:100, and 1:1000) were applied to these wells in duplicate. Abs against hFIX were detected with a 1:40 000 dilution of HRP-conjugated rabbit antirhesus IgG (Sigma). Dilutions of plasma from macaques with anti-hFIX Ab (monkeys RQ1305, RQ1234, and 84268) were included as positive controls. The titer of the Ab was defined as the maximum dilution at which the absorbance at 492 nm exceeded that of preimmune serum by 0.05 absorbance unit. Additionally, positive samples were subjected to the Bethesda assay as described previously. Briefly, citrated rhesus test plasma diluted in Owren buffer was incubated with normal pooled human plasma (Sigma) or normal rhesus plasma at 37°C for 2 hours. The residual hFIX activity (APTT) was then determined by using a 1-step APTT assay. One Bethesda unit was defined as the reciprocal of the dilution of test plasma at which 50% of FIX activity was inhibited. The sensitivity of the assay was 1 BU/mL.

Detection of anti-AAV Abs

An immune-capture assay was used to detect anti-AAV–specific Abs in rhesus plasma. Briefly, wells in a 96-well plate were coated with 100 µL AAV2 antigen (5 × 10^10 rAAV vgs/mL) in 0.1 M sodium bicarbonate buffer, left overnight at 4°C, washed 5 times with PBST, and blocked with 200 µL 6% BSA in PBST for 1 hour at 37°C. After washing, 50 µL appropriately diluted rhesus plasma (1:25, 1:250, and 1:5000 dilutions in PBST) was added to the wells in duplicate. Rhesus IgG Ab against rAAV was detected by using a 1:40 000 dilution of HRP-conjugated rabbit antirhesus IgG (Sigma). Since all values were within the linear range, the titer of the Ab was defined arbitrarily as the maximum absorbance at 492 nm with the 1:5000 dilution of the rhesus serum.

NAb titers were analyzed by determining the ability of rhesus serum to inhibit transduction of 293T cells by a reporter virus in which green fluorescent protein was under the control of the CAGG promoter (rAAV CAGG-GFP). Rhesus samples were diluted in PBS (1:50, 1:500, 1:2000, and 1:5000) and incubated for 60 minutes at 37°C with 1 × 10^9 rAAV CAGG-GFP particles in a total volume of 250 µL in 12-well plates. The mix of virus and serum was then added to 1 × 10^5 293T cells/well in 12-well plates. After overnight incubation, the cells were washed twice with PBS and cultured for an additional 48 hours before evaluation of transfection efficiency by flow cytometry. The NAb titer was calculated arbitrarily as the highest dilution that inhibited transduction of 293T cells by 50%.

Results

Development of an assay for distinguishing hFIX in rhesus plasma

The close homology of the rhesus FIX protein with its human cognate (~97% identity) posed a major challenge to reliable and selective detection of hFIX in rhesus plasma. Previous studies
When antiserum from one of these macaques (RQ1305) was used in a Bethesda assay at 4 months) which did not cross-react with rhesus FIX. Whether the differences in the levels of hFIX among monkeys 1 to 4 were related to technique, the route or schedule of administration, the genetic diversity of these outbred animals, or a combination of these factors is unknown.

Liver biopsy specimens were obtained at various intervals from all macaques and assessed by using Southern blot analysis. The overall copy number of the vgs in liver cells ranged from less than 1 (monkeys 1 and 2) to approximately 40 (monkey 4, 4 × 10^5 copies/μg of genomic DNA) and approximately paralleled the levels of hFIX observed in plasma from these monkeys (Table 1). DNA species corresponding to circular monomers, linear monomers, circular dimers, and HMW concatamers—most of which were present in a head-to-tail configuration, although a small proportion was present in a head-to-head configuration—were detected in monkeys 3 and 4 (Figure 3). Because the monomer form was converted to a linear form on digestion with EcoRI, we assume that it was episomal. The integration status of the HMW concatamers could not be determined from our analyses. RT-PCR analysis confirmed expression of the hFIX transgene at levels roughly reflecting the genome copy numbers in individual monkeys (data not shown).

Dissemination and shedding of virus and humoral response to viral capsid after liver-targeted delivery of vector particles

A PCR assay having a sensitivity of at least 500 genomes (particles)/mL was used to detect vector sequences in body fluids (plasma, urine, and saliva) from monkeys 1 to 4 for 12 days after vector infusion (Figure 4). Varying vg amounts were detected in plasma from all monkeys for 3 days after vector administration. In monkey 3, which received 2 infusions of vector particles separated by 72 hours, vgs were detected in plasma for 6 days. The vgs were detectable at approximately 1% to 25% hFIX in rhesus plasma spiked with hFIX (Figure 1C).

Stable transduction and expression of hFIX after liver-targeted delivery of rAAV particles in rhesus macaques

Vector particles were generated by scale-up of the transient transfection and heparin affinity-column chromatography purification procedures described previously. Before injection in rhesus macaques, each vector preparation was shown to have anticipated bioologic activity in studies in immunodeficient mice (data not shown). The results of liver-targeted administration of rAAV CAGG-FIX in macaques are shown in Table 1. The first 2 macaques received a single bolus infusion into the common hepatic artery with use of interventional radiologic techniques. Both rhesus macaques that specifically neutralized hFIX activity (> 1% to 25% of normal levels). The vitamin K-dependent proteins were then detected in a dose-dependent fashion in rhesus plasma spiked with hFIX (Figure 1B). The specificity of the antisera was further demonstrated by Western blot analysis in which a prominent 70-kDa band, which was detected in a dose-dependent fashion in rhesus plasma spiked with hFIX (Figure 1C).

Table 1. Gene transfer after liver-targeted delivery of rAAV CAGG-FIX in rhesus macaque monkeys

<table>
<thead>
<tr>
<th>Monkey</th>
<th>Weight, kg</th>
<th>Delivery route</th>
<th>Infusion*</th>
<th>Transgene copy no./diploid genome</th>
<th>Maximum hFIX expression (% of normal)</th>
<th>Anti-hFIX Abs</th>
<th>Duration of expression, wks</th>
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<tr>
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<td>&gt; 40</td>
</tr>
<tr>
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<td>7.6</td>
<td>PV</td>
<td>2</td>
<td>25</td>
<td>3</td>
<td>Yes</td>
<td>&lt; 3</td>
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</tbody>
</table>

HA indicates hepatic artery; and PV, portal vein.

*Vector particles (4 × 10^{12} vgs/kg) were given either as 1 infusion or in 2 equal divided doses separated by 3 days.
but in neither urine nor saliva from monkeys 3 and 4. Negative samples were spiked with vector plasmid and subjected to PCR to demonstrate the absence of inhibitors of PCR in the body fluids.

Other than the transient elevations in levels of transaminases in monkeys 1 and 2, all laboratory values, including those for IL-6, remained normal. Low titers of anti-AAV Abs were detected by ELISA in monkeys 1 to 4 before vector infusion at dilutions of 1:25 (monkeys 3 and 4) or 1:100 (monkeys 1 and 2). However, these binding Abs did not function as NAb s, as indicated by their inability to block transduction of 293T cells by a reporter vector rAAV CAGG-EGFP. Vector infusion led to a robust anti-AAV binding IgG response in all 4 macaques after a lag period of 5 and 7 days and coincided with clearance of vector particles from body fluids (Figure 5A). In parallel, there was an increase in the titer of NAb s, which reached peak levels by 3 weeks (Figure 5B). The NAb titers were sustained at maximum levels for an additional 5 weeks before decreasing to undetectable levels by 12 weeks. In contrast, anti-AAV IgG Ab titers persisted at high levels for more than 250 days in all 4 monkeys assessed.

Analysis of spread of rAAV CAGG-FIX to distant organs

There was no evidence of inflammation or dysplasia on histopathological analysis of liver, spleen, kidney, or testes tissue from

Figure 2. Expression of hFIX in rhesus macaques after liver-targeted delivery of rAAV CAGG-FIX. (A) ELISA assay. The hFIX concentration in rhesus plasma was determined at the indicated times after liver-directed administration of rAAV CAGG-FIX in 5 rhesus macaques (M1, M2, M3, M4, and M5). Samples from monkeys 1 to 4 were evaluated independently on 4 occasions; those from monkey 5 were assessed on 2 separate occasions. Results are means ± SEM. (B) Western blot analysis of barium citrate precipitated rhesus serum samples.

Figure 3. Status of rAAV genome in rhesus liver after rAAV-mediated gene transfer. (A) Southern blot analysis of DNA derived from liver biopsy specimens from monkey 2 (M2), monkey 3 (M3), and monkey 4 (M4) at 11, 9, and 6 months after liver-targeted delivery of rAAV CAGG-FIX. Each lane contains 10 µg uncut (UC) DNA or DNA digested with EcoRI, which cuts once within the transgene. The rAAV transgene was not detectable in DNA from the liver of a naive macaque (control). Four different rAAV species were detected in UC genomic DNA from the livers of monkeys 3 and 4, and these were deduced to represent HMW concatamers as well as double-stranded dimer, monomer, and circular forms (arrows). The EcoRI digest indicated that the rAAV transgene persists mainly as head-to-tail tandems with approximately a third maintained in the head-to-head orientation. (B) Schematic representation of the rAAV CAGG-FIX vector showing the position of the probe and the expected fragment size after EcoRI digestion of the vector.

Figure 4. Virus shedding after administration of rAAV in rhesus macaques. A PCR-based assay was used to detect vector sequences in plasma (P), urine (U), and saliva (S) collected from monkeys 1 to 4 on the stipulated days after liver-targeted delivery of rAAV CAGG-FIX. The PCR primers were chosen specifically to amplify a 512-bp hFIX transgene–specific product (arrow labeled FIX). Standards consisting of serial dilutions of rAAV CAGG-FIX in rhesus plasma were used to define the sensitivity of the assay. Twenty percent of the samples were electrophoresed on a 1.5% agarose gel. Negative samples were spiked with vector plasmid and subjected to PCR to ensure that the sample did not inhibit the PCR reaction.

Figure 5. Humoral response to rAAV vector particles. (A) Plasma obtained from monkeys 1 to 4 (same symbols as in Figure 2) was analyzed for the presence of AAV2-specific IgG by ELISA at various times after liver-directed administration of rAAV CAGG-FIX. Results are expressed as the maximum absorbance at 492 nm with the 1:5000 dilution of the rhesus serum. (B) The NAb titer was assessed by defining the highest dilution of rhesus plasma from monkeys 1 to 3 (same symbols as in Figure 2) that inhibited transduction of 293T cells with a reporter virus by 50%.
monkey 3. Genomic DNA from each type of tissue was subjected to PCR analysis using vector-specific primers that amplified a 512-bp region from the 3’ end of the rAAV CAGG-FIX expression cassette. As expected, rAAV CAGG-FIX sequences were found in the liver, with an average of 2 vector copy numbers/diploid genomic equivalent (Figure 6). Vector sequences were also detected in the spleen (~0.005 copies/diploid genome), whereas the kidneys and gonads were consistently negative (detection threshold, <0.001 transgene copies/diploid genome).

**Discussion**

In this study, we examined the safety and efficacy of liver-targeted delivery of $4 \times 10^{12}$ vgs/kg rAAV CAGG-FIX vector particles given to 5 rhesus macaques with use of different routes and schedules of administration. Using an assay developed to specifically detect hFIX in rhesus plasma, we observed stable expression of hFIX at therapeutic levels for more than a year in 2 of 5 macaques. Another macaque initially had therapeutic levels of hFIX, although a specific NAb subsequently developed. We also documented highly efficient transduction of the liver at the genomic level in these 3 macaques, with persistence of the rAAV transgene as HMW concatameric forms as well as smaller episomal forms. Liver-targeted administration of rAAV was well tolerated, even though viral particles were readily detected in plasma and other body fluids for approximately 3 to 6 days after infusion of the vector. An analysis of abdominal organs and the testis found that apart from the liver, the spleen was the only organ with stable persistence of the rAAV transgene.

The copy numbers of the rAAV genome in the 2 macaques that received $4 \times 10^{12}$ vgs/kg vector particles through the portal vein were 40/diploid genome and 25/diploid genome for the single dose and split dose, respectively. These values are more than 10-fold higher than the copy number in mice (0.65 copies/diploid genome) given an equivalent dose of vector particles.8 If these rAAV genomes are distributed to a small subset of hepatocytes, as occurs in mice,32 the copy number in some hepatocytes may be quite high (>500/cell). Additional studies are needed to validate this potentially important biologic advantage for rAAV vectors in primates. If confirmed, this advantage may allow administration of a lower dose of vector particles. Of interest is the fact that the levels of hFIX in the macaque with 40 copies/diploid genome were estimated to be only about 2-fold higher than levels in mice with 0.65 copies/diploid genome, a discordance that also needs further investigation. At high copy numbers, some factor other than genome number may limit hFIX production. On the other hand, because the assays used in the murine and rhesus models employ different Abs for capture of hFIX and different conditions to reduce background, direct comparison of the absolute levels in the 2 species is difficult.

Transduction of the liver after bolus infusion of an equivalent dose of vector into the hepatic artery of monkeys 1 and 2 was poor, leading to subtherapeutic levels of hFIX in rhesus plasma. The vector preparations administered to monkeys 1 to 3 were different, although they were shown to have equivalent biologic activity in mice and therefore differences between them did not produce the difference in transduction efficiency.8 Perhaps noteworthy is the fact that monkeys 1 and 2 had the highest titers of capsid-binding Abs before infusion of the vector particles. These monkeys also had liver toxicity immediately after administration of the vector, which raises the possibility that the transduced hepatocytes were lost during this acute inflammatory insult. Hepatotoxicity has not been observed in mice3-5,8 or dogs4,6 after liver-targeted delivery of rAAV, and the precise cause of this complication in these 2 macaques, which did not occur in the 3 other macaques we studied, remains unknown. There was no evidence of a humoral response to either vector particles or the transgene product during this acute episode, and T-lymphocyte–mediated cytotoxicity is not a feature of rAAV transduction, unlike gene transfer with naked DNA and adenoviral vectors.33-35 Cytotoxic T-lymphocytes (CTLs) to the transgene product or rAAV capsid proteins have not been observed in humans or animal models in the absence of coinfection with adenovirus.35,36 In addition, we readministered rAAV-2 into monkey 2 through the portal vein without producing liver damage or an inflammatory response (A.C.N., A.M.D., A.W.N., unpublished data, January 15, 2002), a finding that suggests that CTLs were probably not responsible for the hepatitis in this macaque.

The development of NAbs against hFIX in monkey 5 represents a serious adverse event that was not anticipated on the basis of results in murine and canine studies. These animal models have provided substantial evidence supporting the ability of the liver to induce immune tolerance to neoantigens.8,9,12,13,37,38 Formation of NAbs (or inhibitors) is a recognized complication of protein-based therapy for hemophilia.39 The incidence after gene therapy may be different than that with protein administration, since the endogenously expressed transgene product is likely to be presented in the context of the major histocompatibility complex (MHC) class I pathway as well as the class II pathway, in contrast to protein concentrates, from which the antigen is presented through the MHC class II pathway. We speculate that the repeated surgical insult primed the rhesus T cells to mediate formation of anti-hFIX Abs in monkey 5. However, the relative contribution of other factors, such as the monkey’s genetic composition, older age, and possible vector extravasation during infusion to the formation of the FIX inhibitor in the context of gene transfer, must be defined. Management of inhibitors involves constant exposure to the offending antigen.40 It is possible that sustained transgene expression after gene therapy will induce tolerance to hFIX in monkey 5, but the inhibitor tier remained elevated (24 BU/mL) at 5 months. A study in a canine hemophilia model also found a neutralizing anticanine FIX Ab in 1 of 4 dogs with a null mutation in the FIX gene.31

Our use of a fractionated-dose schedule in 2 macaques was based on the observations by Miao et al32 that the subset of
hepatocytes susceptible to transduction in mice changes during 2 to 3 days; thus, administration of rAAV at intervals results in persistence of the transgene in distinct populations of hepatocytes. Our goal of augmenting the level of hFIX was achieved when 2 injections were given through the hepatic artery to monkey 3, in which hFIX levels higher than those in monkeys 1 and 2 were observed, but as noted above, the values in the latter 2 monkeys may have been artificially low because of technical difficulties. When the portal route was used, the genome copy number was found to be lower in the monkey that received the split dose (monkey 5) than in the monkey given the single dose (monkey 4). Because of the possible role of repeated surgical trauma in the development of the hFIX inhibitor in monkey 5, we will use the single-dose regimen in future studies.

Southern blot analysis indicated that a marked proportion of the rAAV transgene was maintained episomally, even at 6 to 9 months after administration of the vector. Previous studies in mice suggested that these episomally maintained rAAV genomes make an important contribution to stable expression. These observations raise concerns that rAAV-mediated transgene expression may be lost over time because of a low but constant turnover of hepatocytes, as was observed after AAV-mediated gene transfer into skeletal muscle in nonhuman primates. If expression is lost, readministration of AAV2 may be problematic because macaques have a robust immune response to rAAV particles that persists for more than a year. However, the ability of these anti-AAV Abs to inhibit transduction by rAAV in an in vitro assay appeared to be limited to 3 to 6 months, suggesting that readministration of AAV2 after this period might lead to efficient transduction in vivo.

Our studies demonstrated the feasibility of achieving sustained expression of therapeutic levels of hFIX for more than a year after liver-targeted delivery of rAAV vector in a nonhuman-primate model and identified several critical issues to be addressed in future studies. These include the relation between genome copy number and hFIX levels, the duration of hFIX production at therapeutic levels, the importance of the immune response to vector capsid proteins with respect to the potential of readministration of vector particles for re-establishing or augmenting hFIX production, the relative merits of hepatic artery and portal vein administration, the incidence of inhibitory Abs to hFIX, and the potential for germline transmission of the vg. Although we support initiation of a clinical trial on the basis of current knowledge, concomitant studies in nonhuman-primate models are needed to provide critical information necessary to guide the development of future clinical trials. Such nonhuman-primate studies are likely to be expensive and labor intensive, but the investment is justifiable given the risks involved in gene-therapy approaches and the benefits to be gained from success.

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