Systemic Treatment of Fabry Disease Using a Novel AAV9 Vector Expressing α-Galactosidase A

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Fabry disease is a rare X-linked disorder affecting α-galactosidase A, a rate-limiting enzyme in lysosomal catabolism of glycosphingolipids. Current treatments present important limitations, such as short half-life and limited distribution, which gene therapy can overcome. The aim of this work was to test a novel adeno-associated viral vector, serotype 9 (AAV9), ubiquitously expressing human α-galactosidase A to treat Fabry disease (scAAV9-PGK-GLA). The vector was preliminary tested in newborns of a Fabry disease mouse model. 5 months after treatment, α-galactosidase A activity was detectable in the analyzed tissues, including the central nervous system. Moreover, we tested the vector in adult animals of both sexes at two doses and disease stages (presymptomatic and symptomatic) by single intravenous injection. We found that the exogenous α-galactosidase A was active in peripheral tissues as well as the central nervous system and prevented glycosphingolipid accumulation in treated animals up to 5 months following injection. Antibodies against α-galactosidase A were produced in 9 out of 32 treated animals, although enzyme activity in tissues was not significantly affected. These results demonstrate that scAAV9-PGK-GLA can drive widespread and sustained expression of α-galactosidase A, cross the blood brain barrier after systemic delivery, and reduce pathological signs of the Fabry disease mouse model.

INTRODUCTION

Fabry disease (FD; MIM: 301500) is one of the most frequent lysosomal storage disorders (LSDs), which affects around 1:7,000 individuals, according to several newborn screening studies.1,2 FD is caused by mutations in α-galactosidase (α-GaL; BRENDA: EC3.2.1.22), a rate-limiting enzyme in the lysosomal metabolism of glycosphingolipids. Lack of α-GaL leads to the progressive accumulation of glycosphingolipids, mainly globotriaosylceramide (Gb3), and its decacetylated form LysoGb3. Progressive accumulation of glycosphingolipids within lysosomes of FD individuals occurs in a variety of cell types, including endothelial, smooth muscle, and renal cells (podocytes, tubular cells, glomerular endothelial, mesangial, and interstitial cells), as well as cardiac (cardiomyocytes and fibroblasts) and nerve cells. These events cause a progressive multiorgan disorder that manifests with a painful small fiber neuropathy, cardiac disease, chronic renal insufficiency, and a high predisposition for cerebrovascular strokes.3 FD equally affects males and females because random inactivation of one of the two X chromosomes in females may be sufficient to develop severe manifestations.4 Up-to-date FD is treated by enzyme replacement therapy (ERT), which consists of biweekly intravenous (i.v.) injections of recombinant human α-GaL (agalasidase alpha or agalsidase beta). This therapeutic approach slows down organ damage, stabilizes renal or cardiac parameters, and reduces neuropathic pain crisis in FD patients.5 Nonetheless, ERT presents significant limitations for long-term treatment of FD, such as low half-life and biodistribution, activation of the immune system, the inability to cross the blood brain barrier (BBB), and the mode of administration. Recently, a novel orally active chaperone, migalastat HCl, has been approved for FD.6 Although this drug can achieve therapeutic concentrations in the central nervous system (CNS), its use is only indicated for a fraction of FD patients with amenable mutations in GLA (~70%). Different strategies are currently being developed to increase the efficacy of ERT, including gene therapy and small molecules.7,8 These therapeutic approaches are based on the evidence that even a modest increase in α-GaL activity could prevent clinical manifestations. Indeed, in several LSDs, substrate accumulation occurs when residual enzyme...
activity decays below a threshold (usually activity <1%). The classical form of FD is related to residual α-GalA activity <1% in men, whereas a residual activity of 5%–10% may be sufficient to prevent clinically significant Gb3 accumulation.

In comparison with ERT, adeno-associated viral vector (AAV)-based gene therapy ensures an increased half-life and bioavailability of the enzyme and could be easily directed to specific tissues or even cell types. AAVs are a group of DNA viruses of the Parvoviridae family and the Dependoparvovirus genus, which are incapable of self-replication and can be easily manipulated to produce recombinant proteins. For these advantages, they are currently, extensively used in gene therapy clinical trials.

AAV1, AAV2, and AAV8 serotypes have been used to express α-GalA in murine models of FD, where they successfully cleared glycosphingolipid storage from peripheral organs.

Ogawa et al. used an AAV1 to drive the expression of α-GalA in newborns and adult males of a FD mouse model. AAV1 achieved α-GalA expression in liver, heart, and plasma; however, no effects were observed in adult females.

Ziegler et al. designed hepatospecific targeting to treat FD animal models by combining the AAV8 serotype (with high transduction affinity for the liver) and a liver-restricted promoter (DG190). The local administration of the vector in the liver afforded successful levels of α-GalA even in peripheral organs; in addition, the elevated hepatic concentration of the enzyme induced immunotolerance to the transgene, a favorable feature for possible clinical applications. Site-specific genome editing, providing long-term, stable therapeutic expression from the endogenous promoter in the serum albumin locus, has also been tested. Lastly, a lentiviral vector-based ex vivo gene therapy approach for FD is under investigation in clinical trials.

Overall, the major limitation of the methods that have been developed thus far is that they do not allow sufficient expression of α-GalA in the CNS. Although no severe cognitive impairment has been associated to FD, α-GalA is expressed in neurons and glia. Neurologic symptoms are substantial in the pathophysiology of the disease, and stroke at early age is one of the main causes of premature death in FD patients.

Therefore, the aim of this work was to develop an efficient vector for the expression of α-GalA in the CNS and peripheral organs of an FD mouse model. Our approach was based on the systemic injection of an AAV9, which demonstrated tropism for nervous tissue and the ability to cross the BBB.

We found that systemic administration of the newly generated vector mediated widespread α-GalA expression and restored its function in the CNS and in multiple tissues of a knockout mouse model of FD at both the presymptomatic (at birth and 1 month of age) and symptomatic stages of the disease (at 3 months of age). Furthermore, we demonstrated that the treatment prevented glycosphingolipid accumulation in the most affected tissues (heart, liver, kidney, CNS) by immunofluorescence, ultrastructural examination, and liquid chromatography-mass spectrometry (LC-MS) analysis of either Gb3 or Lyso-Gb3. Interestingly, the therapeutic effect was observed in both males and females and was not reduced even in the presence of a humoral immune response in mice injected at 3 months of age. Our data indicate that this gene therapy is a very promising approach for the translation into treatment protocols for FD.

**RESULTS**

**Injection of Self-Complementary AAV9-Phosphoglycerate Kinase-α-GalA (scAAV9-PGK-GLA) Induces High and Widespread α-GalA Activity In Vivo**

To efficiently express α-GalA in vivo, we produced a novel scAAV9 vector encoding the human α-GalA cDNA (GenBank: NM_000169.3), under the control of the ubiquitous PGK promoter (scAAV9-PGK-GLA). We also generated a similar AAV9 vector, expressing the green fluorescent protein (GFP), used as control (scAAV9-PGK-GFP).

We initially tested the expression of the two vectors in wild-type FVB/NJ mice. To ensure a widespread distribution of the viral particles and detection of α-GalA in multiple organs, we codelivered a high dose of vector (3.2 × 10^{11} vector genome [vgl/kg]) via combined intracerebroventricular (i.c.v.) and i.v. injections at postnatal day 1 (P1), as we previously described. 2 months after treatment, we analyzed α-GalA activity in the brain, spinal cord, heart, kidney, and liver of injected mice. In all tissues of the animals injected with scAAV9-PGK-GLA, the enzymatic activity was significantly higher (14- to 82-fold) compared to noninjected or GFP-treated mice (Figure S1) except in the kidney (5-fold increase). Notably, α-GalA activity was 20-fold higher in the brain and spinal cord of the mice injected with scAAV9-PGK-GLA compared to noninjected or scAAV9-PGK-GFP-treated mice, indicating that the vector gets efficiently expressed in the CNS. This pilot experiment demonstrated that the scAAV9-PGK-GLA vector was efficient to induce widespread overexpression of a functional α-GalA in vivo.

**Single i.v. Injection of scAAV9-PGK-GLA Induces Sustained α-GalA Activity and Reduces Gb3 Accumulation in FD Mice Treated at Birth**

To test the therapeutic efficacy of the scAAV9-PGK-GLA vector in vivo, we used the α-GalA knockout mouse strain B6;129-Gal<sup>ΔmIKse</sup>/J (FD mice) in which Galα is deleted in the X chromosome. These mice recapitulate FD symptoms, namely the accumulation of glycosphingolipids and abnormal kidney and liver morphology, as well as mild cardiovascular pathology, by 3 months of age. However, they have a mild phenotype and normal lifespan, and do not develop evident movement or behavioral alterations.

The i.v. delivery of scAAV9 vectors has already been approved for gene replacement to treat an infantile form of a rare motor neuron disease. For this reason and to facilitate translation into the clinic,
we chose to test our vector following a single-dose i.v. injection in FD mice. We injected scAAV9-PGK-GLA or control scAAV9-PGK-GFP in the temporal vein of newborn hemizygous FD mice at the dose of 1.8 × 10^{14} vg/kg. To assess short- and long-term effects of α-GalA expression, we sacrificed FD mice at either 3 or 5 months after injection and compared data obtained in mice treated with scAAV9-PGK-GLA to age-matched positive (wild-type male mice) and negative controls (noninjected or scAAV9-PGK-GFP-injected hemizygous). The experimental plan is presented in Table 1.

We measured α-GalA activity in different organs (brain, spinal cord, liver, heart, and kidney). We found that the human enzyme was expressed and functional 3 and 5 months after injection, with maintained or higher levels of α-GalA activity at 5 rather than at 3 months after injection. Moreover, the enzymatic activity was similar in mice injected with the scAAV9-PGK-GLA and wild-type mice in all of the analyzed tissues. In contrast, residual activity was undetected, either in noninjected hemizygous FD mice or in mice treated with the control vector scAAV9-PGK-GFP (Figure 1). Interestingly, this test in FD mice confirmed that the vector was able to drive transgene expression also in the B6129 mouse background.

We also assessed the distribution of the glycosphingolipid deposits by immunofluorescence and electron microscopy. The amount of Gh3 storage was lower in the heart, kidney, liver, and brain of animals injected with the scAAV9-PGK-GLA vector compared to noninjected hemizygous FD mice. Specifically, Gh3 deposits, detected by immunohistochemistry, were reduced in intracardiac fibroblasts, vascular smooth muscle cells, endothelial cells, and cells (possibly Kupffer cells) of the hepatic sinusoidal space (Figure 2). Importantly, the ultrastructure of electron-dense, intralysosomal inclusions and multilamellar bodies typical of FD were reduced in the heart (including cardiomyocytes), kidney, and vessels of an injected mouse compared with a nontreated hemizygous (Figures S2 and S3). Consistent with previously reported data, we could not observe intralysosomal inclusions in neurons and hepatocytes of the mice from all of the experimental groups, including the noninjected FD mice (Figure S4). By immunofluorescence, we detected the Gh3 signal (but not lysosomal storage) in the cells of the CNS (Figure S5; cerebellum), where this glycosphingolipid is expressed in neurons of cerebral cortex, hippocampus, and cerebellum, independently of the genotype. The Gh3 signal appeared more intense in nontreated mice compared to 5-month-old wild-type mice, and a partial decrease of signal intensity was detected in mice treated at birth with scAAV9-PGK-GLA after 3 and 5 months. These results demonstrated that the presymptomatic treatment of FD mice with the scAAV9-PGK-GLA vector restores α-GalA activity in both the CNS and peripheral organs for up to 5 months and induces a marked, long-lasting reduction of Gh3 deposits in multiple organs of FD mice.

**Systemic Injection of scAAV9-PGK-GLA Preserves Body Weight of Adult FD Mice of Both Sexes, Treated at the Presymptomatic and Symptomatic Stages**

The encouraging results observed in the newborn animals prompted us to test the scAAV9-PGK-GLA vector in adult FD mice at two different stages of the disease (1 month of age, presymptomatic; or 3 months of age, presenting clinical symptoms).

We performed a comprehensive analysis of the effects induced by the scAAV9-PGK-GLA systemic injection in adult FD mice.

To evaluate a dose-response effect, we tested two vector doses: 6 × 10^{13} vg/kg and 1.8 × 10^{14} vg/kg. For a complete therapeutic assessment, we also included heterozygous females in which Gla is knocked out in one allele, as shown in the experimental plan presented in Table 2. The use of heterozygous female mice instead of homozygous is clinically relevant because women FD patients are usually heterozygous.

We monitored mice weekly for health conditions and body weight over a period of 5 months after treatment. The body weight analysis did not reveal any significant effect of the treatment on the growth of FD mice (Figure S6), with no difference in body weight in treated or untreated FD mice versus wild-type mice (mixed ANOVA). This indicates that the FD mouse model does not present growth defects when compared to wild-type mice over the first 8 months of age and suggests that the treatment was well tolerated.

Although the hemizygous mice, injected at 1 month of age with the highest dose, had a lower starting body weight compared to the animals of the same age in the other experimental groups, the rate of weight increase in this group was comparable with the one registered in the remaining groups (p = 0.191, ANOVA). This confirms a lack of impact of the gene therapy on FD mouse growth.

Survival was unaffected in all treated mice, and no phenotype alteration was noticed during the study, confirming the lack of a severe phenotype in FD mice. Both treated and untreated FD mice were slightly more aggressive than wild-type animals. This could be explained by the presence of sensorimotor function alteration, similarly to the human pathology. However, this symptom tends to disappear over time in both humans and mice, due to progressive nerve-ending degeneration, caused by Gh3 accumulation. In general anxiety, depression, and cognitive symptoms, which are frequently described in FD patients, have a subordinate relevance in the mouse model.
Systemic Injection of scAAV9-PGK-GLA Induces Expression of a Functional α-GalA in Multiple Tissues of Adult FD Mice, Including the Brain

To assess to what extent the scAAV9-PGK-GLA was able to target different tissues, we titrated vector particles in tissues 5 months after injection, by droplet digital PCR (ddPCR), using specific primers for scAAV9 genome. We detected scAAV9-PGK-GLA in the brain, heart, liver, and kidney of mice injected at 1 month and 3 months of age, at a concentration ranging between 11 and 2,120 vg per diploid genome (dg; Figure 3A). The highest transduced organ was the liver (ranging from 757 to 2,120 vg/dg), whereas the brain was the organ with the lowest absolute quantification of viral particles (ranging from 11 to 75.4 vg/dg). Viral vector concentrations in the different tissues were comparable between mice injected at 1 month or 3 months of age, since differences (nonparametric t test) between groups with the same sex and treatment, at the two different ages, were not statistically significant. This suggests that equal amounts of vector reached the tissues independently of the stage of the disease. Despite that vector dose was adjusted to mice weight, it appeared that the transduction of the BBB was more efficient when mice were injected at a younger age (1 month). Viral vector titer in tissues tends to be higher in animals treated with the higher dose of the vector, although differences among the groups did not reach statistical significance.

To analyze whether the scAAV9-PGK-GLA induced enzyme expression in different organs of FD mice, we assessed α-GalA expression by western blot (WB) in brain, liver, heart, kidney, and plasma samples from treated and nontreated FD mice. The enzyme was expressed in all of the analyzed tissues from treated animals injected at either 1 month or 3 months of age (Figures 3B, 4, S7, and S8) compared to nontreated FD mice. Plasmatic protein levels were variable among animals, and in 12 out of 32 of the analyzed samples, α-GalA was barely detectable (Figure S7).

To evaluate whether the replaced enzyme was functional, we assessed specific α-GalA activity in plasma and tissue samples (brain, spinal cord, liver, heart, and kidney) of all groups at the study end point (Figure 5).

Similarly to what we observed in newborn mice, the α-GalA was functional and in all analyzed tissues of the adult animals that were injected with scAAV9-PGK-GLA vector. Consistently with WB results, activity in plasma was variable.
Importantly, the levels of enzyme activity in the brain and spinal cord of the adult hemizygous-treated animals ranged between 10% and 30% of physiological activity (noninjected wild-type FD males) in the same organs. Heterozygous-treated animals presented enzyme levels in the CNS comparable to those of noninjected wild-type FD females.

We also found that when we injected the mice at 1 month of age (in the presymptomatic phase), the activity levels increased with the dose, whereas this was not always the case when animals were injected at 3 months of age, consistently with the appearance of cell damage at this stage.
To analyze the interdependence of the variables (activity, protein levels, and viral titer), we calculated Spearman’s rank correlation coefficient (Rho; Table S1) using data obtained in plasma and tissue samples derived from organs that were evenly and consistently cut in fragments and lysed with the appropriate protocol.

Plasmatic α-GalA levels and activity tendentially correlated in samples from the treated hemizygous mice. The presence of the endogenous allele likely impacted a potential correlation between the two variables in heterozygous mice (Figure S9).

Viral particles distribution (assessed by ddPCR), exogenous protein expression (WB data), and α-GalA activity, among the different organs, were consistently higher in injected mice compared to noninjected FD mice, but we did not find a significant correlation among these three variables. However, we observed that in several of the assessed conditions, Rho value was higher than 0.7 (correlated variables), and in the case of heterozygous mice injected with the lower dose, correlation between virus titer and activity in the liver was close to significance (p = 0.051).

Overall, we demonstrated that scAAV9-PGK-GLA induces the expression of functional α-GalA in adult FD mice, and the efficacy of the treatment is higher when this is applied before the appearance of the symptoms.

**scAAV9-PGK-GLA Mediates Expression of α-GalA-Reduced Glycosphingolipid Deposits in Adult FD Mice**

To assess whether FD symptoms were reversed by scAAV9-PGK-GLA injection, we also analyzed levels of the major Gb3 circulating metabolite, Lyso-Gb3, in plasma (before injection, as well as 4 months and 5 months after treatment) and in tissues (brain, liver, heart, and kidney) by LC-MS.

Although Lyso-Gb3 levels in plasma tended to increase over time in nontreated mice, they were dramatically reduced in the mice injected with scAAV9-PGK-GLA (Figure 6). Lyso-Gb3 concentration was significantly decreased in treated hemizygous mice injected at 1 month (p < 0.0001) and at 3 months (p < 0.001) for each time and dose, as well as in heterozygous mice injected at 1 month (p < 0.05 for each time and dose) compared to noninjected genotype-matched mice. In heterozygous FD mice injected at 3 months, the p value of Lyso-Gb3 decrease is <0.05, 4 months after injection, and p < 0.09 (6 × 10^{13} vg/kg) or p < 0.06 (1.8 × 10^{14} vg/kg), 5 months after injection, compared to non-treated FD heterozygous. Plasma Lyso-Gb3 concentration in noninjected FD heterozygous was about 10-fold lower compared to the one in noninjected hemizygous. This may explain why the difference in Lyso-Gb3 concentration between treated heterozygous and nontreated animals is smaller and therefore not statistically significant in all of the assessed conditions. This is not surprising, since the low concentration of Lyso-Gb3 in plasma of female human patients is one of the main limitations in which to consider Lyso-Gb3 as an optimal biomarker for FD diagnosis.

Similarly, Lyso-Gb3 concentration in the tissues of the treated FD mice was significantly lower compared to the levels found in nontreated genotype-matching animals.

This was in line with what we observed in the immunohistochemistry experiments. Gb3 deposits were detectable in the same cell types described for the newborn-injected mice (i.e., intracardiac fibroblasts, renal epithelial tubular cells, vascular smooth muscle cells, endothelial cells, Kupffer cells; Figures 7A and S10). Moreover, we observed an intense Gb3-positive signal in the bone (osteoblasts and bone marrow cells) and the neurons of the dorsal root ganglia in spinal cord sections from noninjected FD hemizygous mice, which was considerably decreased in the same cells of FD mice treated at 3 months of age with both doses of the virus (Figures 7B and S11). In the brain, deposits are mainly detectable in blood vessels of nontreated FD mice and are partially cleared in the same mice after injection with scAAV9-PGK-GLA (Figure S12; showing hippocampus and cortex).

These results denote that the exogenous α-GalA is able to reverse the pathological signs of the FD model.

**The Detected Humoral Immune Response to α-GalA Does Not Affect Enzyme Activity in Tissues of Injected FD Mice**

Lastly, we analyzed whether the injection with the scAAV9-PGK-GLA induced the production of immunoglobulin G (IgG) antibodies against α-GalA, which may neutralize the action of the enzyme. We found that 9 out of 32 injected animals (28.1%) produced IgG antibodies against human α-GalA in relevant concentrations (Figure 8).

Six of these mice were injected at 3 months of age during the symptomatic phase. Moreover, the activity in plasma was lower than average in samples from all of the animals injected at 3 months and

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**Table 2. Experimental Plan of Treatments in Adult FD Mice**

| Age of Injection | Injected with scAAV9-PGK-GLA | Noninjected Controls |
|------------------|-----------------------------|---------------------|
|                  | Dose (6 × 10^{13} vg/kg)    | Dose (1.8 × 10^{14} vg/kg) | Hemizygous | Heterozygous | Wild Type |
| 1 month          | 4 hemizygous, 4 heterozygous | 4 hemizygous, 4 heterozygous | 4 males | 4 females | 4 males, 4 females |
| 3 months         | 4 hemizygous, 4 heterozygous | 4 hemizygous, 4 heterozygous | 4 males | 4 females | 4 males, 4 females |

Mice were injected in the tail vein with a solution of scAAV9-PGK-GLA at the concentration of 3.13 × 10^{12} vg/mL in PBS, adjusting the injected volume to the weight of the mice, to obtain a final dose of either 6 × 10^{13} vg/kg or 1.8 × 10^{14} vg/kg.
Figure 3. Viral Vector Is Distributed in Tissues of Presymptomatic and Symptomatic FD Mice and Drove the Expression of α-GalA after Systemic Injection

(A) Histograms of viral titer (vg/dg) assessed by ddPCR absolute quantification in different tissues (brain, heart, liver, and kidney) from mice of both sexes injected with scAAV9-PGK-GLA at either 1 month or 3 months of age and sacrificed 5 months later. Measurements were carried out in duplicate samples for each mouse of the group (N = 4). Error is expressed as SEM. Data were compared using nonparametric t test. (B) Histograms of western blot densitometric analysis of α-GalA, normalized to the corresponding total protein per lane (Figure S8) and reported to the intensity value obtained for the corresponding noninjected control. Error is expressed as SEM. Significance of the data, comparing each value with the value of the corresponding noninjected control, was assessed by nonparametric t test (*p < 0.05; **p < 0.01; ***p < 0.001).
two of the animals injected at 1 month (presymptomatic) that produced anti-$\alpha$-GalA antibodies (Table S2). This suggests higher immunological tolerance in presymptomatic mice (81% of the animals) compared to symptomatic mice (44%). Although low enzymatic activity in plasma of treated FD mice tended to correlate with low levels of the protein in plasma, a neutralizing effect of the activity due to the presence of anti-$\alpha$-GalA antibodies cannot be excluded.

Nevertheless, $\alpha$-GalA activity was not significantly affected in the analyzed tissues from mice that produced IgG antibodies. Indeed, activity in the liver, brain, heart, and kidney from mice that produced IgG antibodies against $\alpha$-GalA was comparable or higher than the average activity of the whole group in each tested condition. Only in the spleen, which is directly involved in IgG production, did we detect decreased enzymatic activity (34%–86% of average) in anti-$\alpha$-GalA IgG-producing mice (Table S2). Thus, these results suggest that despite the possible presence of neutralizing antibodies, the therapeutic effect of the treatment is maintained, since levels of $\alpha$-GalA activity in tissues are similar to the ones found in mice that do not produce anti-$\alpha$-GalA IgGs.

Further studies will be necessary to fully evaluate the extent of the immune response and the significance for the clinical translation of this vector.

**DISCUSSION**

In this proof-of-concept study, we generated and fully characterized a novel AAV vector for in vivo delivery of $\alpha$-GalA as an efficacious treatment for FD. Through a single i.v. injection of an $\alpha$-GalA-expressing scAAV9, we were able to observe therapeutic effects in a well-established mouse model of FD. We demonstrated that scAAV9-PGK-GLA was able to efficiently express human $\alpha$-GalA in both the CNS and peripheral organs of FD newborn and adult mice. The human enzyme was functional and reached sufficient levels to mediate a beneficial effect in either hemizygous or heterozygous FD mice at both presymptomatic and symptomatic stages. Indeed, after treatment, Lyso-Gb3 levels, which inversely correlate with $\alpha$-GalA, were almost null in plasma and multiple tissues (including the CNS) of FD mice treated with scAAV9-PGK-GLA compared to nontreated FD mice.

CNS targeting represents a major advancement for FD treatment, since currently developed therapies fail to rescue pathological signs in the brain. Among the treatments in use, ERT is not able to reach the CNS, and migalastat, which can diffuse through the BBB, is not indicated for all patients. Although FD is mostly considered a pathology with peripheral and cardiovascular implications, we believe that the expression of $\alpha$-GalA in the CNS of FD patients has a therapeutic value, especially when this is reached using noninvasive methods. $\alpha$-GalA is normally present in different cell types of the brain, whereas glycosphingolipid inclusions have been observed in neurons, Schwann cells, perineural cells, dorsal root ganglia, and autonomic ganglia in FD patients, leading to CNS and peripheral nervous system pathology. Neurological symptoms are important for patients who very frequently experience neuropathic pain, depression, anxiety, and stroke at an early age, as well as white matter damage. Moreover, a recent study also suggests a possible link between FD and Parkinson’s disease.
Other gene therapy-based treatments, which target mainly peripheral organs or the liver, are in development for FD; however, none of them is focused on the neurological effects of the protein.

Ex vivo gene therapy approaches with lentiviral vector-mediated correction of a GLA genetic defect in CD34+ hematopoietic stem cells and endogenous cell transplantation are currently being tested in clinical trials. However, preclinical data did not assess the expression of α-GalA in the CNS. In addition, ex vivo gene therapy has some constraints that are generally related to the percentage of engrafted cells and sustainability of the transgene expression.

In vivo gene therapy, based on direct injection of viral vectors, allows sustained expression of the α-GalA and represents an advantage over ERT. For instance, Ziegler and coworkers tested a hepatocyte-specific AAV8 vector achieving long-lasting α-GalA expression in FD mice and nonhuman primates. The main advantage of liver-specific gene therapy relies on its low immunogenicity, since this approach facilitates the development of tolerance. Liver-specific expression of GLA is also achieved by the injection of ST-920, an AAV2/6-based vector that reaches supraphysiological levels of α-GalA in plasma and peripheral tissues without safety issues at the preclinical stage. Nonetheless, these strategies behave like an improved ERT, since the recombinant protein is specifically produced in the liver and distributed to other tissues, but it cannot be expressed in the CNS.

Similar to our vector, the construct tested by Ogawa et al. also drives the expression of α-GalA through a ubiquitous promoter (CAG). However, unlike scAAV9-PGK-GLA, their vector was not able to reach a significant increase of α-GalA in FD females’ organs nor the CNS. Indeed, Ogawa et al.’s vector preferentially drives α-GalA expression in the liver and to a lesser extent, in the heart of hemizygous adult mice. On the contrary, we demonstrated that our novel vector scAAV9-PGK-GLA is able to transduce tissues and brain of the adult FD mice, independently of sex, leading to increased α-GalA activity in peripheral organs (liver, heart, and kidney), as well as the CNS of FD-treated mice versus untreated mice of the same strain. Therefore, the expressed α-GalA is functional and is able to cross the BBB, with highest levels measured in FD mice injected at the presymptomatic stage.

AAV9 was successfully used to develop gene therapy vectors in other lysosomal disorders, such as the mucopolysaccharidosis I, II, and III. Specifically, intrathecal cervical AAV9 gene transfer of

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Figure 5. Activity of α-GalA Is Restored in FD Mice Treated with scAAV9-PGK-GLA at Presymptomatic and Symptomatic Stages

Activity of α-GalA (nmol/hour x mg) was measured 5 months after injection in brain, spinal cord, liver, heart, kidney, and plasma samples from FD mice. Activity values from mice injected with scAAV9-PGK-GLA, at either 1 month or 3 months of age, were compared with values obtained in noninjected (negative controls) or wild-type (positive controls) mice. Values are expressed as the mean ± SEM. Statistical significance of the differences between treated mice and noninjected controls was analyzed by nonparametric t test ( * p < 0.05; ** p < 0.01; *** p < 0.001; **** p < 0.0001).
human α-L-iduronidase administered to 1-month-old rhesus monkeys supports the safety and efficiency of the treatment in these animals, even when they were tolerized to the human transgene at birth via systemic administration of an AAV8 vector. The AAV9 expressing α-L-iduronidase efficiently transduced pyramidal neurons in the cortex and hippocampus, Purkinje cells in the cerebellum, lower motor neurons, and dorsal root ganglia neurons.

Systemic injection of scAAV9-PGK-GLA in FD mice produced levels of α-GalA activity in the CNS of about 30% of the physiologic activity (wild-type controls); this finding is clinically relevant because 10% residual enzymatic activity is known to be sufficient to prevent glycosphingolipid accumulation and the development of symptoms in FD patients. Moreover, the vector demonstrated a very high capacity to transduce CNS cells (α-GalA activity increased up to 20-fold) when we administered it through combined i.v. and i.c.v. injection in FVB/NJ newborn mice.

Overall, to the best of our knowledge, our newly developed scAAV9-PGK-GLA is the first vector to enable sustained therapeutic levels of α-GalA in the CNS, through a systemic injection. Therefore, it could be intended as a viable alternative or as a complement for other successfully developed gene therapy approaches when elevated levels of gene expression are needed in the CNS.

Importantly, multiple complementary approaches (LC-MS, immunofluorescence, and ultrastructural examination) demonstrated that glycosphingolipid accumulation was decreased after scAAV9-PGK-GLA treatment, and deposits were mostly cleared in a variety of cell types of injected FD mice. Interestingly, the distribution of Gb3 deposits in tissues from FD mouse does not completely resemble the pattern found in FD classical patients, whereas it mimics mainly the histological features of late-onset FD patients. Indeed, podocytes are minimally or nonaffected by the presence of Gb3 storage, which accumulates in epithelial cells of collecting ducts and distal tubules or in vascular smooth muscles and interstitial fibroblasts of the heart, as confirmed by our results (Figures 2, 7A, S2, S3, and S10). However, unlike Bangari et al., we were also able to detect electron-dense, intralysosomal inclusions in a small percentage (~5%–10%) of cardiomyocytes and endothelial cells of the noninjected hemizygous mice, in which formation was prevented by treatment with scAAV9-PGK-GLA (Figures S2E and S2F).
Furthermore, in contrast with human pathology, classical multilamellar body accumulation was not described in hepatocytes and brain neurons of hemizygous FD mice, whereas affected neurons were only found in the sensory region of dorsal root ganglia. In accordance with these findings, glycosphingolipid deposits were considerably decreased by the scAAV9-PGK-GLA treatment in neurons of the dorsal root ganglia of hemizygous FD mice (Figure 7B), but we did not observe electron-dense, intralysosomal inclusions, characteristic of FD, in brain neurons and hepatocytes of nontreated FD hemizygous mice (Figure S4). However, we could detect a Gb3-positive signal in the brain by immunohistochemistry (Figures S5, S10, and S12), consistently with the fact that these glycosphingolipids are normally present in the brain of C57BL/6J mice, and the intensity of the signal is partially reduced by the treatment with scAAV9-PGK-GLA in the FD model.

These data were also supported by the LC-MS detection of increasing concentrations of Lyso-Gb3 in the brain of nontreated hemizygous and heterozygous FD mice over time, which are prevented by the treatment that we applied (Figure 6).

Notably, scAAV9-PGK-GLA was effective in reducing Gb3 storage in osteoblasts and bone marrow cells of FD mice (Figure 7B). This finding is important for the translational application of the vector, as it strongly supports the potential of AVV9-based gene therapy for the treatment of LSDs other than FD, such as Gaucher disease and mucopolysaccharidoses, in which bone involvement is a major problem.

Our data also demonstrate that the systemic injection of a scAAV9 vector mediates a sustained expression of α-GalA over time, rescuing the FD pathological signs up to 3 months after treatment in mice. This is not surprising, because AAV9 vectors are known to mediate long-lasting expression of transgene under different settings, including mouse models and patients. However, here, we report evidence of the prolonged and widespread effect of such an approach for FD, including the CNS, thereby opening a concrete perspective for the translational application of this vector. Specific experiments will have to be performed to assess safety issues related to α-GalA overexpression, although the general health condition of the animals and the survival data suggest that the enzyme levels are well tolerated.

α-GalA activity data were in accordance with the distribution of the viral particles and the expression of the protein among the different tissues, even though we did not find a significant correlation among the three variables. In plasma, there is a clear tendency of the activity to correlate with the levels of α-GalA in samples from the treated hemizygous mice, which is also evident in mice that produce anti-bodies against the enzyme. Indeed, IgG antibodies against the α-GalA were produced in 9 out of 32 injected mice with plasmatic activity lower than the average of the group. Although reduced activity correlates with low levels of plasmatic protein, we cannot exclude that the anti-α-GalA antibodies neutralize the activity of the enzyme in plasma, since the enzyme-antibody complexes could have been removed from the circulation by macrophage action. However, the production of IgGs against α-GalA did not seem to affect the functionality of the enzyme in tissues, since activity values in organs from IgG-producing mice were not significantly different from the average value of the group. In particular, animals that were treated at the earlier stage of the disease (1 month, presymptomatic) were less prone to produce antibodies against the enzyme and also showed a more robust dose-response effect. On the contrary, mice treated at 3 months, when Gb3 was already substantially accumulated in the organs, more frequently produced IgG antibodies against the enzyme and did not show a correlation between the enzymatic activity and the administered vector dose. We believe that the poor dose-response correlation in the scAAV9-PGK-GLA-treated FD mice at 3 months of age could be due to a combination of factors, including both pre-existing tissue damage and immune response.

In contrast, the treatment with scAAV9-PGK-GLA in pups with the highest dose of the virus (1.8 × 10^{14} vg/kg) resulted in high production of functional α-GalA in all analyzed organs. In these initial studies, we did not assess the immune response due to the immaturity of the newborn immune system and the presence of maternal antibodies that inhibit newborn antibody production. During this phase, the immune system is not fully developed, and foreign elements can be more easily tolerated. Thus, the AAV administration to newborns could facilitate an immune tolerance to the transgene products.

Importantly, functional α-GalA was expressed in the brain of all the AAV-treated pups, as the transduction across the BBB was more efficient in newborns than in the adult-injected mice, most likely related to the higher permeability of the barrier at this age. Overall, our findings support the conclusion that an early administration of scAAV9-PGK-GLA will result in higher therapeutic efficacy, similar to currently used treatments in FD (ERT or chaperones).

Further studies will be necessary to reduce immunogenicity of the vector, other than applying the treatment at an early age in the presymptomatic stage. For example, local intraventricular injection in the brain could be used to reduce adaptive immune response and widespread expression of the transgene. This delivery route could be used in combination with a treatment that is excluded by the CNS. However, systemic application will most likely lead to a better compliance of the patients, and therefore, early stage treatment...
should be prioritized to reach immunotolerance in a pathology like FD with limited damage in the CNS.

When envisioning systemic delivery of AAV9 vectors as treatment for human diseases, several aspects need to be considered to avoid immune-mediated toxicity that would lead to deaths in clinical trials, as recently reported. Attention must be taken in preclinical studies regarding dose, immune response, and toxicity.

In conclusion, our work demonstrates that the scAAV9-PGK-GLA vector could be a novel potential therapy to treat systemic as well as neurological manifestations of FD. This vector was able to successfully rescue the pathological phenotype of a mouse model of FD when it was injected in the blood circulation of newborns and presymptomatic and symptomatic animals. Given that the standard of care, ERT, does not reach the CNS, and chaperone therapy is indicated for a restricted group of patients, the proposed gene therapy vector represents a promising preclinical therapeutic candidate for FD, which overcomes most of the limitations of currently used approaches.

**MATERIALS AND METHODS**

**Vector Cloning**

Human α-GalA cDNA (GenBank: NM_000169.3) was amplified using Phusion High-Fidelity DNA polymerase (F532; Thermo Scientific) from the pR-M10-α-GalA plasmid, and GFP was amplified using the same method but different primers from the scAAV9-cytomegalovirus (CMV)-GFP vector plasmid, previously described in Tanguy et al. PCR fragments were cloned in the recombinant plasmid (pscAAV9)-PGK-survival of motor neuron gene (SMN) described in Besse et al., following digestion with Hind III and Not I (unique restriction sites) to eliminate the chimeric intron and the cDNA. Ligation with PCR-amplified GLA cDNA was performed with Quick Ligation (NEB; #M2200S). The ligated vector was transformed in XL10-Gold ultracompetent cells (Agilent Technologies), and DNA was extracted to identify positive colonies by enzymatic restriction. Correctly oriented clones were selected by sequencing with primers aligned to the vector.

**Virus Production**

The viral particles were produced following the protocol described in Biferi et al. Briefly, HEK293T cells were cotransfected, using polyethylenimine, with the adenovirus helper plasmid (pXX6Helper), the rep2 cap9-encoding plasmid (AAV9 capsid), and either pscAAV9-PGK-GLA or pscAAV9-PGK-GFP plasmids. Following 72 h
incubation at 37°C, cells were harvested, and viral particles were purified through an iodixanol gradient. Virus titer, expressed as vg per milliliter, was measured by qPCR using specific primers and virus standard curve with known concentrations.

**Animals, Injections, and Tissue Harvesting**

Wild-type FVB/NJ mice (Janvier Labs) were used to perform a pilot study to assess vector activity. Animals were maintained in the Centre of Functional Evaluation-UMS28 (Sorbonne University, Paris, France) under controlled conditions (22 ± 2°C, 50 ± 10% relative humidity, 12 h/12 h light/dark cycle, food and water *ad libitum*). All animal procedures followed the European guidelines for the care and use of experimental animals.

* B6:129-Gla^tm1Kul/J (FD mice; The Jackson Laboratory; #3535) were maintained in the animal facility at Vigo University (Servicio de Bioexperimentación SB-Uvi REGA; ES60570215601), in accordance with external and internal biosafety and bioethics guidelines. The experimental procedures (ES60570215601/17/INVMED02/OUTROSO4/SO01 and SO02) were approved by Vigo University Committee (0001-2017SO and 00003-2017SO) and authorized by the competent authority (Xunta de Galicia, Conselleria do Medio Rural, Pontevedra, Spain).

Mice were identified at P1 with ink tattoos in the fingers and genotyped following the protocol indicated by The Jackson Laboratory for this strain.

For the first therapeutic test in newborn FD mice, groups of 4 hemizygous newborn mice, between P1 and P3, were injected with either scAAV9-PGK-GLA or scAAV9-PGK-GFP through the temporal vein (40 μL of viral solution at 1.3 × 10^{13} vg/mL). These treated animals and a control group of noninjected hemizygous or wild-type animals were maintained in the experiment for either 3 or 5 months. At the end points, organs (brain, spinal cord, heart, liver, kidney, and spleen) were removed and divided in two tubes: one snap frozen in liquid nitrogen for biochemical assays and one containing paraformaldehyde (PFA; 4% v/v). In subgroups of mice, tissue fragments were embedded in glutaraldehyde for ultrastructural analysis.

In adult FD mice, the scAAV9-PGK-GLA was injected in groups of 4 hemizygous and heterozygous animals at 1 month or 3 months of age to assess presymptomatic and symptomatic treatment, respectively. Viral suspension (3.13 × 10^{13} vg/mL) was injected in the tail vein at two different doses (6 × 10^{13} vg/kg or 1.8 × 10^{14} vg/kg) in each group of animals; volume of vector solution was adjusted to compensate for weight differences.

As control groups, noninjected wild-type or knockout mice of both sexes were used. Total number of animals N = 64, as described in Table 2. Submandibular vein blood withdrawal was performed at baseline (before injection), at 4 months following injection, and at the end of the protocol. 5 months after the administration of the vector, as described for newborn mice, animals were sacrificed by lethal injection (200 mg/kg ketamine and 30 mg/kg xylazine) to collect brain, spinal cord, heart, liver, kidney, spleen, thymus, and skeletal muscle. An oriented fragment of the organs was fixed in PFA and used for immunohistochemistry. The remaining tissues were snap frozen and then lysed for biochemical assessments. Brains were divided in hemispheres; one was embedded in PFA, and the other was consistently cut, as evenly as possible, in three fragments that were used for the biochemical analyses with the appropriate lysis protocol.

**AAV9 Particle Distribution in Tissues**

DNA was automatically extracted from frozen tissue sections on the QIAcube platform, using the DNeasy kit (#69504; QIAGEN) and QIAamp Mini Rotor Adaptor (#1064532; QIAGEN). Absolute quantification was determined by ddPCR testing DNA samples (2.5 ng/μL) in duplicates. Primers/probe mix was specific for the recombinant AAV9 genome (primers: 5’-TTTATAGGCTTCCTTG-3’ and 5’-TGGATTAGTGGCCTG-3’, 900 nM; probe: Fluorescein-AMester-5’-TAGTAATGATTACCC-3’-Quencher, 250 nM). Negative (no template control [NTC]) and positive (scAAV9-PGK-GLA particles 10^3 vg/μL) controls were used in each experiment. Droplets were automatically generated with a QX200 Automated Droplet Generator (Bio-Rad). ddPCR was performed using the following cycling conditions: 10 min at 95°C, 40 cycles of two-step thermal profile: 95°C at 30 s and 57.5°C at 60 s with a ramp rate of 2°C/s. Plates were finally transferred to a QX200 Droplet Reader (Bio-Rad), and data analysis was performed using QuantaSoft software (Bio-Rad). The threshold separating negative and positive droplets was set manually just above the cluster of negative droplets. Data from wells with number of droplets below 10,000 were excluded from analysis. Results were expressed as vg/μL (vector copies per ng of extracted DNA).

**Expression of Human α-GalA in Tissue Lysates**

Relative expression levels of α-GalA in tissues were determined by WB. Briefly, tissues were lysed with lysis buffer (#B9803; Cell Signaling Technology) containing protease inhibitor cocktail (#P8340; Sigma-Aldrich), homogenized with Ultra-Turrax, sonicated, and centrifuged 20 min at 10,000 x g (4°C) to collect cytosolic fractions. Proteins were separated through SDS-PAGE in 4%–15% polyacrylamide stain-free Mini-Protein or Criterion TGX gels (#4568083, #5678081; Bio-Rad) and transferred to 0.2 μm polyvinylidene fluoride (PVDF) membranes (Transblot Turbo, #1704157; Bio-Rad). The membranes were blotted with a monoclonal anti-human-α-GalA antibody (#ab129173; Abcam). Total load was estimated using Image Lab software (Bio-Rad), by imaging the activated stain-free gels and following the recommendations of the manufacturer. Each antigen band was normalized to its total protein lane (Figure S8) using the total protein normalization tool (Image Lab Software) and following the manufacturer’s protocol. Band intensity values were divided for the intensity value obtained in noninjected hemizygous or heterozygous control sample (background).
**α-GalA Enzymatic Activity Assay**

Activity of α-GalA was measured using the method of Chamoles et al. Briefly, tissues were embedded in deionized water (300 μL) and subjected to three freeze-thaw cycles before homogenization with the Ultra-Turrax homogenizer (IKA-Werke). The preparations were sonicated and centrifuged at 10,000 × g to collect cytosolic fractions. The activity of cell lysates was measured in 0.15 M phosphate-citrate buffer (pH 4.2) in the presence of the substrate, 4 mM 4-methylumbelliferyl-α-D-galactopyranoside (Glycosynth) and 50 mM N-acetyl-D-galactosamine (#A2795; Sigma-Aldrich). The reactions were incubated for 2 h at 37°C, and at this point, the stopping solution (0.1 M ethylenediamine, pH 11.4) was added to halt the reaction. Then, the fluorescence from samples was read using a Twinkle LB 970 fluorometer (Berthold) at 360 nm excitation and 450 nm emission wavelengths. Specific activity was expressed as nanomoles of hydrolyzed substrate per hour and milligram of total protein, extrapolating nanomoles of hydrolyzed substrate from a standard curve (fluorescence versus known concentrations of 4-methylumbelliferone; Merck; #M1381). Total protein concentration was measured with the Pierce Bicinchoninic Acid (BCA) Protein Assay Kit (Thermo Fisher Scientific), following the manufacturer’s protocol.

**Gb3 and Lyso-Gb3 Deposits Quantification and Distribution**

Lyso-Gb3 was quantified by LC-MS in tissue and plasma samples using the protocol described by Nowak et al. This analysis was performed at ARCHIMED Life Science (Vienna, Austria). Plasma samples were collected at pre-dose at 4 months and at 5 months following injection. Tissue lysates were prepared as described for enzymatic activity measurements, and lipid extraction was performed at ARCHIMED Life Science. Lyso-Gb3 concentration in tissue was expressed as nanograms per milligrams of protein and in plasma as nanograms per milliliter of sample.

Distribution of affected lysosomes was examined by Gb3 immunostaining with anti-CD77 antibody (clone BGR23; #A2506; AMS Biotechnology) and costaining with 4′,6-diamidino-2-phenylindole (DAPI) and/or rhodamine phalloidin to detect nuclei and membranes (polymerized actin), respectively. Immunohistochemistry was performed on frozen tissues sections of 10 μm. For spinal cord samples, organs were decalcified overnight before organs were decalcified or 1% citrate phosphate buffer (pH 6; 5°C) for antigenic retrieval. Images were taken with a conventional fluorescence microscope DM6 equipped with a DFC550 camera (Leica; Figures 2, 7A, S5, and S10) or the inverted microscope DMi6000B with a DFC365EX camera (Leica; Figures 7B and S12). Quantitative assessment of deposits was performed using the quantification tool of LAS AF software (Leica) and expressed as mean fluorescence intensity (pixel gray levels).

Ultrastructural analysis was performed by transmission-electron microscopy in selected samples. Rectangular pieces for electron microscopy were fixed in 2.5% glutaraldehyde, post-fixed in osmium tetroxide, and embedded in Epon after routine dehydration. Semi-thin sections were stained with toluidine blue, and ultra-thin sections were contrasted with uranyl acetate and lead citrate and mounted in copper grids. Ultra-thin sections were examined with a Philips CM100 transmission electron microscope.

**Antibody Production against Human α-GalA**

To assess the immunoresponse of α-GalA, an ELISA test in plasma samples was set up to detect anti-human α-GalA IgG antibodies. Briefly, clear 96-well plates (Nunc-ImmuNo MicroWell 96-well solid plate; Merck) were coated with 1 μg/mL α-GalA (#6164-GH; R&D Systems) and incubated overnight at 4°C. Plates were blocked with 2% BSA in PBS at 4°C. Serum samples were diluted (1:100 in blocking solution) and incubated for 90 min at 37°C. Antibodies binding α-GalA were detected using horseradish peroxidase (HRP)-conjugated goat anti-mouse IgG antibodies (1:10,000) (#ab6789; Abcam). Samples were incubated 1 h at room temperature and then visualized using the BioFX TMB/Stop solution (#ab171522; Abcam). The reaction was halted with 1 vol of HCl 0.1 N. Absorbance was measured at 450 nm. IgG concentrations were calculated referring to a standard curve of commercial anti-human GLA antibody (#ab129173; Abcam) and donkey anti-rabbit IgG polyclonal secondary antibodies (#NA934; GE Healthcare).

**Statistical Analysis**

Power analysis to determine the number of animals to be included in each group was performed with the software InVivoStat, assuming that α-GalA activity would increase at least 2 mmol/h × mg in hemizygous animals injected with the scAAV9-PGK-GLA vector (power 80% and p > 0.05). For ddPCR, WB, activity, and Lyso-Gb3 data, significance was assessed by nonparametric t tests (2 tails) using Graph Prism software.

For each tissue, time point, and doses, the correlation among variables (virus titer, α-GalA expression, and activity) was assessed using Rho.

Statistical differences in the body weight were assessed by mixed ANOVA, combining Mauchly’s test of sphericity and Greenhouse-Geisser correction.

**SUPPLEMENTAL INFORMATION**

Supplemental Information can be found online at https://doi.org/10.1016/j.omtm.2020.10.016.

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AUTHOR CONTRIBUTIONS

M.G.B. designed the vectors and participated in the analysis of the results and in the writing of the manuscript. M.C.-T. prepared the viral particles for injections. A.G.-S. and S.T.-B. performed western blot analysis of α-GalA in the different tissues and enzymatic activity assays. A.G.-S. also prepared the figures of the paper and analyzed the data. O.S.-R., B.S.-M.-T., and S.B. performed immunohistochemistry and ultrastructural analysis of tissues and the interpretation of the results. I.V.-G. and T.M. performed DNA isolation from tissues and the determination of viral titration and distribution in tissues. T.M., A.F.-C., and V.D. took care of animal maintenance, monitoring, and dosing. A.G.-F. helped to design the animal experimental project and develop the serum antibody detection assay. She also revised the manuscript. M.B. supervised the study. S.O. designed the project, and develop the serum antibody detection assay. She also revised the manuscript. T.M., and ultrastructural analysis of tissues and the interpretation of the results. I.V.-G. and T.M. performed DNA isolation from tissues and the determination of viral titration and distribution in tissues. T.M., A.F.-C., and V.D. took care of animal maintenance, monitoring, and dosing. A.G.-F. helped to design the animal experimental project and develop the serum antibody detection assay. She also revised the manuscript. M.B. supervised the study. S.O. designed the project, and develop the serum antibody detection assay. She also revised the manuscript.

DECLARATION OF INTERESTS

The authors declare no competing interests.

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