Enzyme Replacement Therapy Clears Gb3 Deposits from a Podocyte Cell Culture Model of Fabry Disease but Fails to Restore Altered Cellular Signaling

Fabian Braun a,b,c,d, Linda Blomberg a,b,c, Susanne Brodesser b, Max C. Liebau a,b,c,e, Bernhard Schermer a,b,c,f, Thomas Benzing a,b,c,f, Christine E. Kurschat a,b,c

© 2019 The Author(s). Published by Cellular Physiology and Biochemistry Press GmbH&Co. KG

Key Words
Fabry • Podocyte • Enzyme Replacement Therapy

Abstract

Background/Aims: Fabry disease (FD) is a lysosomal storage disorder characterized by impaired alpha-galactosidase A (α-Gal A) enzyme activity due to mutations in the GLA gene. While virtually all tissues are affected, renal damage is particularly critical for the patients’ outcome. Currently, powerful diagnostic tools and in vivo research models to study FD in the kidney are lacking, which is a major obstacle for further improvements in diagnosis and therapy. The present study focuses on the effects of enzyme replacement therapy on a previously established podocyte cell culture model of Fabry disease. Methods: We investigated the effect of in vitro application of α-Gal A on Fabry podocytes for 3 days, mimicking enzyme replacement therapy. We studied reduction of Gb3 levels and dysregulated molecular pathways such as autophagy, mTOR/AKT signaling and pro-fibrotic signaling by employing immunofluorescence, electron microscopy, tandem mass spectrometry and western blot. Results: We detected complete resolution of Gb3 accumulation in Fabry podocytes upon α-Gal A treatment. Despite robust Gb3 clearance, dysregulation of the signaling pathways investigated was not reversed. Conclusion: This study presents first evidence for Gb3-independent effects regarding dysregulation of signal transduction mechanisms in FD not recovering upon α-Gal A treatment. We assume that intracellular alterations observed in FD

F. Braun and L. Blomberg contributed equally to this work.

Prof. Dr. Christine Kurschat, MD
Department II of Internal Medicine, University Hospital Cologne
Kerpener Str. 62, 50937 Cologne (Germany)
Tel. +49 221 478 4480, Fax +49 221 479 5969, E-Mail christine.kurschat@uk-koeln.de
may have a point of no return after which a reversal of dysregulated cellular signal transduction by α-Gal A treatment is not effective, despite Gb3 clearance. Our observations suggest further research on signal transduction mechanisms altered in Fabry podocytes and on determining the appropriate time for initiation of Fabry therapy.

Introduction

Fabry disease (FD) [OMIM: 301500] is an X-linked hereditary disorder caused by mutations in the alpha-galactosidase A (GLA) gene resulting in an impairment of glycosphingolipid metabolism. Due to loss or defective function of the enzyme alpha-galactosidase A (α-Gal A) the terminal α-glycosidic bond of galactose cannot be cleaved from the sphingolipid globotriaosylceramide (Gb3), thus resulting in accumulation of Gb3 within the lysosome. This defect and the subsequent lysosomal dysfunction can be detected in virtually every cell in every tissue [1]. The clinical presentation encompasses a broad spectrum of symptoms ranging from gastrointestinal dysfunction, hypohidrosis, heat sensitivity and fever to severe neuropathic pain, ischemic stroke at very young age, cardiomyopathy and renal insufficiency [2]. While hemizygous male patients typically display severe symptoms, the clinical manifestations seen in heterozygous female patients can comprise the whole range of clinical features from the absence of symptoms to very severe pathologies as seen in male patients with classical FD. Premature death mostly occurs due to cardiac complications and end-stage renal disease (ESRD) [2].

Renal insufficiency is caused in part by progressive Gb3 depositions primarily affecting podocytes, terminally differentiated cells with limited potential of recovery and regeneration [3, 4]. First renal symptoms, consequently, are (micro-) albuminuria and combined proteinuria at later stages, rarely coinciding with the development of hypertension. Subsequently, the histological phenotype is characterized by progressive glomerulosclerosis and nephron loss leading to tubular atrophy and interstitial fibrosis [2]. Due to the multitude of unspecific symptoms the majority of Fabry patients are diagnosed late in the course of their disease after organ damage has already occurred.

To date, treatment of Fabry disease relies on either intravenous enzyme replacement therapy or, in the case of amenable mutations, oral chaperone administration [5, 6]. Both approaches have been shown to ameliorate clinical symptoms and disease progression, with limited potential to reverse already established organ damage.

Since the current murine models for Fabry disease differ significantly from the glomerular phenotype seen in human disease [7-10], our group established an immortalized human podocyte cell culture model with robust α-Gal A activity reduction and subsequent Gb3 accumulation [11]. Molecular characterization demonstrated dysregulated autophagy as a probable result of decreased mTOR and AKT phosphorylation and protein abundance.

In this study, we describe the successful use of a compound for enzyme replacement therapy, agalsidase alfa, to clear Gb3 deposits in our established podocyte cell culture model for Fabry disease. Interestingly, despite Gb3 clearance, we did not detect any effect of enzyme replacement therapy on dysregulated autophagy, increased mTOR activation or other, already published molecular alterations such as increased Notch1 activation or dysregulated TGF beta signaling in Fabry podocytes. These findings suggest yet unknown disease mechanisms independent of Gb3 accumulation and warrant further research for additional therapeutic targets of renal dysfunction in Fabry disease.

Materials and Methods

Human podocyte cell culture

Conditional immortalized human podocytes and podocytes with reduced α-Gal activity were cultured as previously described [11]. Briefly, immortalized podocytes were stably transduced with co-shRNA
(control) and shRNA 894 (reducing α-galactosidase A activity to 2-4%, Fig. S1 – for all supplemental material see www.cellphysiolbiochem.com), cultured in RPMI media (Sigma-Aldrich, Taukirchen, Germany) with insulin-transferrin-sodium selenite as supplement (ThermoFisher Scientific, Waltham, USA) containing 10% fetal bovine serum (Biochrom, Berlin, Germany). Cells proliferated at 33°C until they reached a confluence of 60-70%. Subsequently, cells were differentiated at 37°C for 14 days before harvesting them for further analyses. Cells were regularly tested for mycoplasma infection using the mycoplasma detection kit from Minerva biolabs (Minerva Biolabs, Berlin, Germany).

**Treatment with α-Gal A**

Agalsidase alfa (Shire, Berlin, Germany) was used at a final concentration of 250 nM. A portion of the provided agalsidase alfa was heat-inactivated at 95°C for 30 minutes to serve as a negative control. Chemical chaperone deoxygalactonojirimycin hydrochlorid (DGJ, Sigma-Aldrich, Taukirchen, Germany) was added at a final concentration of 10 µM to stabilize the enzyme under cell culture conditions. Cells were treated with agalsidase alfa at Day 8, 10 and 12 of differentiation.

**Stress Induction via Interleukin 1β**

For western blot analysis of mTOR and AKT phosphorylation, cells were treated with interleukin 1β at a concentration of 10 ng/ml for one hour prior to western lysate preparation.

**Immunofluorescence**

After 7 days of differentiation podocytes were seeded onto coverslips to a confluency of 80%. Cells were treated with α-Gal A according to the above mentioned protocol for the following 7 days at 37°C. After fixation with 4% PFA and blocking with normal donkey serum in PBS containing 0.1% Triton-X100, coverslips were incubated with rat anti-Gb3 (CD77) antibody (Abcam, ab19795) o/n at 4°C. Secondary antibody staining was performed for 1 hour at room temperature with Cy3-conjugated goat anti-rat IgM μ chain specific (Jackson, 112-165-020). Coverslips were mounted in ProLong Diamond with DAPI (ThermoFischer, P36962) and imaged with a ZEISS LSM710 confocal microscope (Zen software).

**RT-qPCR**

Cells were lysed and total RNA was isolated (TRI Reagent, Sigma Aldrich). RNA concentration was measured (Nanodrop 1000 spectrophotometer, Peqlab) and cDNA was synthesized from 1000 ng RNA template using the High Capacity cDNA RT Kit (Applied Biosystems) according to the supplier’s protocol. Dilutions were made for RT-qPCR analysis to determine mRNA expression levels which were normalized against a reference gene. The Power SYBR Green PCR Master Mix (Applied Biosystems) was used to multiply and measure the cDNA with a 7900HT Fast Real-time PCR System (Applied Biosystems). The following PCR program was used: 95°C for 10 min, followed by 40 cycles of 15s at 95°C, 60s at 60°C, then 15s at 95°C followed by a melt of the product from 60°C-95°C. The primer sequences (IDT) used are: hACTB forward 5'-GGACTTCGAGCAAGAGATGG, hACTB reverse 5'-AGCACTGTGTTGGCGTACAG, hHPRT1 forward 5'-TGACATGCGAAACAAATGCA, hHPRT1 reverse 5'-GGTCTCTTCTACCAAGCAAGCT, hGLA forward 5'-TGGAAGGATGCAGGTTATGAG, hGLA reverse 5'-CCCTAGCTTCAGTCCTTTGCT, hHES1 forward 5'-ACCAAAGACAGCATCTGAGCA, hHES1 reverse 5'-GCCCGGAGCTATCTTTCTTCA, hCOL4 forward 5'-CTCCTGGTTCTCCACAGTCAG, hCOL4 reverse 5'-AAGACCTGCGACAGACGGAAG, hFN1 forward 5'-GGTCCGGGACTCAATCCAAAT, hFN1 reverse 5'-ACCCGGAAGCAGAACTTTCT, hTGFβ1 forward 5'-CCCAGCATCTGCAAAGCTC, hTGFβ1 reverse 5'-GTCAATGTACAGCTGCCGCA, hHPRT1 forward 5'-TGACACTGCGAAACAAATGCA, hHPRT reverse 5'-GGTCTCTTCTACCAAGCAAGCT. The ΔΔCT method was used for statistical analysis to determine relative gene expression levels.

**Enzyme assay**

Cultured human podocytes were collected in cold PBS. Cell lysis was performed by freezing and thawing over five cycles. Enzyme activity for α-Galactosidase A (α-Gal) and β-Galactosidase (β-Gal) was measured after incubation of lysed cells with 4-methylumbelliferyl a-D-galactopyranoside for α-Gal or 4-methylumbelliferyl b-D-galactopyranoside for β-Gal (Sigma, Taufkirchen, Germany) as described previously [12]. α-Gal A activity was normalized to β-Gal activity, functioning as an internal control.
Western blot

Size separation was done using SDS-PAGE. Samples were blotted onto polyvinylidene difluoride membranes and visualized with enhanced chemiluminescence after incubation of the blots with corresponding antibodies anti-phospho AKT (S473), anti-AKT, anti-phospho-mTOR (S2448), anti-mTOR, anti-phospho-p38 (Thr180/Tyr182), anti-p38, anti-phospho-p42/44 (Thr202/Tyr204), anti-p42/44 (Cell Signaling, Danvers, MA, USA); anti-β-catenin (BD, Heidelberg, Germany); anti-β-tubulin (Developmental Studies Hybridoma Bank, University of Iowa, USA), Notch1 cleaved Val (Abcam, Cambridge, UK). For quantification of band intensity, individual bands of at least three independent experiments were quantified using ImageStudio lite software (Li-Cor, Lincoln, USA). The intensity of every band is expressed as ratio to the corresponding loading control.

Gb3 quantification

Gb3 levels in podocytes were determined by Liquid Chromatography coupled to Electrospray Ionization Tandem Mass Spectrometry (LC-ESI-MS/MS): Cells were harvested in ice cold PBS and centrifuged at 3,000g, 4°C for 5min. Supernatants were removed and cell pellets frozen at -20°C. After thawing cells were homogenized in water (10³ cells / 100 µl) using the Precellys 24 Homogenisator (Peqlab, Erlangen, Germany). The protein content of the homogenate was routinely determined using biocinchoninic acid. To 100 µl of homogenate 750 µl of methanol/chloroform 2:1 (v/v) and 200 pmol of the deuterated internal standard D3-Gb3 C18:0 (Matreya, State College, PA, USA) were added. Lipids were extracted overnight at 48 °C. Interfering glycerolipids were degraded by alkaline hydrolysis adding 75 µl of 1 M potassium hydroxide in methanol and incubating for 2h at 37 °C. After neutralization with glacial acetic acid, the lipid extract was desalted by reversed phase C18 (RP-18) chromatography. Dried lipid extracts were resolved in 200 µl of the LC mobile phase solvent (see below) and sonicated for 5min. After centrifugation (12, 000 × g, 5min, 4°C), 40 µl of the clear supernatants were transferred to auto-injector vials. LC-MS/MS analysis was performed using a Core Shell Kinetex C8 column, 100 x 2.1 mm ID, 2.6 µm particle size, 100 Å pore size (Phenomenex, Aschaffenburg, Germany) and detection using a QTRAP 6500 triple quadrupole/linear ion trap mass spectrometer (SCIEX, Darmstadt, Germany). The LC (1260 Infinity Binary LC System; Agilent, Waldbronn, Germany) was operated isocratically at a flow rate of 500 µl/min with a mobile phase of 0.1 % formic acid solvent (see below) and sonicated for 5min. After centrifugation (12, 000 × g, 5min, 4°C), 4 µl of the clear supernatants were transferred to auto-injector vials. LC-MS/MS analysis was performed using a Core Shell Kinetex C8 column, 100 x 2.1 mm ID, 2.6 µm particle size, 100 Å pore size (Phenomenex, Aschaffenburg, Germany) and detection using a QTRAP 6500 triple quadrupole/linear ion trap mass spectrometer (SCIEX, Darmstadt, Germany). The LC (1260 Infinity Binary LC System; Agilent, Waldbronn, Germany) was operated isocratically at a flow rate of 500 µl/min with a mobile phase of 0.1 % formic acid and 2 mM ammonium acetate in methanol. 20 µl of sample were injected. The total run time was 3 min. Gb3 species were monitored in the positive ion mode using their specific Multiple Reaction Monitoring (MRM) transitions of the [M+H]+ precursor ion to the product ion m/z 264.4 (quantifiers) and to the product ion generated by the loss of one molecule of water and the three hexose residues (qualifiers). The instrument settings for nebulizer gas (Gas 1), turbogas (Gas 2), curtain gas, and collision gas were 60 psi, 65 psi, 35 psi, and medium, respectively. The Turbo V ESI source temperature was 350 °C, and the ionspray voltage was 5500 V. For all MRM transitions the values for declustering potential, entrance potential, and cell exit potential were 210 V, 10 V, and 8 V, respectively. The collision energies ranged from 36 to 68 V. The Gb3 quantifier peaks were integrated using the Analyst 1.6.2 software (AB SCIEX). Endogenous Gb3 species were quantified on the basis of external calibration curves which were calculated from LC-MS/MS measurements of serially diluted synthetic Gb3 standard solutions prepared in the mobile phase solvent within the range of 0.0 to 190 pmol on column. To each dilution a fixed amount of the internal standard D3-Gb3 C18:0 was added. The standard calibration curves were plotted based on molar concentration versus peak area ratio of Gb3 standards to D3-Gb3 C18:0. Linearity and correlation coefficients (R2) of the calibration curves were obtained via linear regression analysis. R2 of the calibration curves were >0.99. The calculated amounts of endogenous Gb species were normalized to the protein content of the sample.

Electron microscopy

Cells were grown on collagen-coated melamine foil before start of α-Gal A treatment. After 7 days of treatment, cells were washed in PBS and fixed in 2% Paraformaldehyde, 2% Glutaraldehyde in 0, 1M sodium cacodylate buffer for 30 minutes at room temperature and 30 minutes at 4°C. After washing and incubation in 1% osmium tetroxide cells were dehydrated in rising ethanol and embedded in epoxy resin. Ultra thin slices of 70nm were done at the Ultracut (Leica Systems, Wetzlar, Germany). Contrast enhancement was achieved through incubation in 1, 5% uranylacetate and leadnitrate. Analysis and images were done at the JEM-2100 PlusTransmission Electron Microscope (JEOL, Toko, Japan).
Results

α-Gal A treatment for three days clears Gb3 deposits in human podocytes with reduced expression of alpha-galactosidase

We used the human podocyte cell culture model for Fabry disease published previously [11]. This cell culture model is characterized by reduced levels of GLA mRNA and functional enzyme, as well as Gb3 accumulation (Fig. S1a-c). As previously described [11], this phenotype results in dysregulated autophagy detected by increased LC3-II expression and lower mTOR and AKT phosphorylation in podocytes with reduced α-Gal A activity (Fig. S1d-e). We started treatment of control and α-Gal A deficient podocytes after seven days of differentiation. Active or inactivated α-Gal A, both including 1-deoxygalactonojirimycin (DGJ), was administered every other day for a total amount of 3 treatments. DGJ is an iminosugar also known as migalastat. We used this compound to stabilize α-Gal A enzyme activity in vitro. Migalastat has recently been licensed for the treatment of Fabry disease in patients with GLA mutations leading to α-Gal A folding defects amenable to migalastat treatment. Immunofluorescence staining (IF) for Gb3 deposition depicted a marked reduction in fluorescent signal in Fabry podocytes treated with active α-Gal A (Fig. 1A). This effect was abolished entirely upon heat inactivation of the enzyme (Fig. 1A). To confirm the results in IF and to quantify Gb3 reduction by α-Gal A treatment we quantified Gb3 levels by Liquid Chromatography coupled to Tandem Mass Spectrometry (LC-MS/MS). Likewise, we detected a significant decrease of total Gb3 content down to control levels after α-Gal A treatment, while Gb3 accumulation prevailed when cells were treated with inactivated enzyme (Fig. 1B). These findings were confirmed by LC-MS/MS analysis of Gb3 subspecies differing in the fatty acyl chain lengths (Fig. 1C).

Gb3 clearance upon α-Gal A treatment does not ameliorate dysregulated autophagy

After having established a successful protocol for in vitro enzyme replacement therapy resulting in almost complete clearance of Gb3 in our cell culture model, we analyzed previously described Fabry-associated alterations in autophagy regulation. We expected to reverse dysregulated autophagy by α-Gal A treatment. Surprisingly, despite almost complete clearance of Gb3 from α-Gal A-deficient podocytes by treatment with α-Gal A, we did not observe a decrease in LC3-II expression when compared to untreated Fabry podocytes or podocytes treated with heat-inactivated enzyme (Fig. 2A). To investigate cellular ultrastructure we performed electron microscopy (Fig. 2B). Zebra bodies, a correlate of accumulated storage material, were present in untreated Fabry podocytes. We did not detect any zebra bodies in actively treated α-Gal A-deficient podocytes, indicating clearance of Gb3 by treatment. Despite α-Gal A administration, overrepresentation of autophagosomes was still present (Fig. 2B). Examining the total abundance and phosphorylation of mTOR and AKT proteins under stress conditions (IL 1β treatment), we detected significant effects of reduced Gb3 accumulation only on AKT abundance (Fig. 2C, D), and merely a tendency to augmented mTOR abundance and mTOR/AKT phosphorylation. Interestingly, treatment with DGJ and inactivated enzyme resulted in a significant increase of both AKT/mTOR abundance and phosphorylation even though this condition depicts high levels of Gb3. These results might point to a chaperone-mediated effect on AKT/mTOR. To determine whether the dysregulated autophagy in α-Gal A-deficient podocytes is due to transcriptional changes, we investigated the expression of two genes involved in autophagy initiation, Beclin 1 (BCN1) and GABA Type A Receptor-Associated Protein (GABARAP) [13]. As expected we detected increased expression in Fabry podocytes compared to control cells. This effect remained present despite enzyme replacement therapy and subsequent Gb3 depletion (Fig. 2E, F).

α-Gal A impairment results in activation of the Notch1 and TGF beta cascades

Sanchez-Niño et al. described activation of Notch1 and TGF beta pathways in cultured human podocytes by increased concentrations of circulating lyso-Gb3 [14, 15]. We were
Fig. 1. Gb3 detection in α-Gal A-deficient podocytes at baseline and under ERT. (A) Immunofluorescence of Gb3 (gray), counterstained with DAPI (blue) of control (coshRNA) and α-Gal A-deficient podocytes (shRNA894) (scale bar 50 µm). (B) Quantification of total Gb3 content measured by LC-MS/MS normalized to total protein levels in control (coshRNA), α-Gal A-deficient podocytes (shRNA894) and α-Gal A-deficient podocytes treated with active (+α-Gal A) or inactivated (i-α-Gal A). Bars depict mean ± SD, **** = p<0.0001. (C) Quantification of Gb3 subspecies measured by LC-MS/MS normalized to total protein levels in control (coshRNA), α-Gal A-deficient podocytes (shRNA894) and α-Gal A-deficient podocytes treated with active (+α-Gal A) or inactivated (i-α-Gal A). Bars depict mean ± SD, significance levels not indicated.
Fig. 2. Markers of autophagy in α-Gal A-deficient podocytes at baseline and under ERT. (A) Representative LC3-II and β-tubulin immunoblot and densitometric quantification of LC3-II/β-tubulin immunoblot. Band intensity of LC3-II was normalized to β-tubulin band intensity (mean ± SD, n=5, 1 Way ANOVA, p=0.8128). (B) Representative electron microscopy of control (coshRNA) and α-Gal A-deficient podocytes (shRNA894) as well as α-Gal A-deficient podocytes treated with active (+α-Gal A) or inactivated (i-α-Gal A) enzyme with marked autophagosomal / lysosomal structures (white arrowheads: autophagosomes, black arrowheads: zebrafish bodies, n=4) (C) Representative phospho-mTOR, mTOR and β-Catenin immunoblot and densitometric quantification of immunoblots. Band intensity of phospho-mTOR/mTOR was normalized to β-Catenin band intensity (mean ± SD, n=3 (shRNA894), n=5 (+α-Gal A, i-α-Gal A), 1 Way ANOVA, * = p<0.05. (D) Representative phospho-AKT/AKT and β-Catenin immunoblot and densitometric quantification of immunoblots. Band intensity of phospho-AKT/AKT was normalized to β-Catenin band intensity (mean ± SD, n=3 (shRNA894), n=5 (+α-Gal A, i-α-Gal A), 1-Way ANOVA, * = p<0.05). (E) RT-qPCR of BECN1, GABARAP expression, normalized against ACTB expression in control (coshRNA) and α-Gal A-deficient podocytes (shRNA894) (mean ± SD, n=5, Students T-Test, * = p<0.05). (F) RT-qPCR of BECN1, GABARAP expression, normalized against ACTB expression in α-Gal A-deficient podocytes (shRNA894) and α-Gal A-deficient podocytes treated with active (+α-Gal A) or inactivated (i-α-Gal A) enzyme (mean ± SD, n=5, 2-Way ANOVA).
able to confirm their results in our model by detecting enhanced cleavage of Notch1 in α-Gal A-deficient podocytes (Fig. 3A) and, subsequently, higher expression of target genes fibronectin (FN1) and Hes Family BHLH Transcription Factor 1 (HES1) as well as augmented transcription of TGF beta target Collagen 4A4 (Col4A4) (Fig. 3B). When investigating the influence of Gb3 depletion upon α-Gal A treatment on Notch1, Col4A4, FN1 and HES1, again, we did not find an amelioration of the phenotype in α-Gal A-treated podocytes (Fig. 3C, D).

Discussion

This study presents the first report of successful and rapid Gb3 clearance by enzyme replacement therapy (ERT) in a human podocyte cell culture model of Fabry disease. The employed protocol resulted in complete removal of accumulated Gb3 down to control cell levels (Fig. 1) after only three treatment cycles of α-Gal A and DGJ. DGJ was used to stabilize α-Gal A activity in vitro. The α-Gal A concentration used in our cell culture experiments represents about four times the amount of α-Gal A present in a patient’s circulation with five liters of blood volume directly after agalsidase alfa infusion at a concentration of 0.2 mg/kg body weight. We used a higher α-Gal A concentration in vitro since most cell culture models exhibit high levels of resilience to pharmacological compounds and, subsequently, higher...
concentrations of compounds are usually needed to cause treatment effects in vitro [16]. In our Fabry podocyte model α-Gal A treatment demonstrates a high efficacy in clearance of Gb3 deposits in vitro. We observed no Gb3 clearance after α-Gal A heat inactivation, even though DGJ was still present in this experimental setup. DGJ, also known as migalastat, is capable of enhancing enzyme activity by stabilization of the enzyme, especially in GLA mutations leading to folding defects of the protein structure. Our cell culture model is characterized by a reduction of GLA gene expression by small hairpin RNAs. Therefore, the remaining cellular enzyme activity (2-4%, Fig. S1) is due to the residual expression of a conventionally folded and normally functioning protein. The remaining activity is not enhanced by DGJ in our model, consistent with only a mild increase in normal α-Gal A activity upon DGJ administration [17].

A clear effect of migalastat on Gb3 accumulation in podocytes in the context of amenable mutations, analyzed by EM of patient renal biopsies, has previously been reported [18].

Strikingly, Gb3 clearance from our podocyte cell culture model did not result in the normalization of the autophagy machinery (Fig. 2A). Electron microscopy depicted numerous autophagosomes still present in α-Gal A-treated Fabry podocytes. The absence of α-Gal A treatment effects on dysregulated autophagy in our model might be due to the shorter period of therapy we employed in this study. A decrease of LC3 staining in kidney biopsies after three years of enzyme replacement therapy was reported in 2010, presenting immunofluorescence images of glomeruli [19]. Clear evidence that decreased LC3 immunofluorescence mirrored amelioration of autophagic flux in podocytes was not provided. Our data suggest that mere replacement of active enzyme and subsequent rapid decrease of intracellular Gb3 deposits is not sufficient to directly improve dysregulated autophagy in our podocyte model.

We observed a tendency to increased mTOR and AKT expression as well as mTOR and AKT phosphorylation upon treatment with α-Gal A and DGJ (Fig. 2C, D). In the context of heat-inactivated α-Gal A employment +DGJ this difference was significant. This observation may point towards a chaperone-dependent pleiotropic effect, since DGJ was present while α-Gal A was inactive. We hypothesize that decreased binding of DGJ to the denatured protein may lead to this more pronounced effect. Still, DGJ treatment alone did not result in restoration of dysregulated autophagy activity. Increased transcription of critical genes involved in autophagy initiation was present in Fabry podocytes (Fig. 2E). BECN1 and GABARAP represent two members of the Coordinated Lysosomal Expression and Regulation (CLEAR) network [20] with Transcription Factor EB (TFEB) being its most prominent transcription factor. TFEB, a master regulator of lysosome integrity and autophagy, is known to play a role in multiple lysosomal storage diseases such as Gaucher disease and Niemann-Pick disease, and is currently investigated as a potential therapeutic target in Pompe disease [21-23]. TFEB function also affects podocyte biology [24]. A recent study by our group investigating primary urine cells of Fabry patients identified additional proteins of the CLEAR network to be more abundant in Fabry patients compared to healthy individuals [25].

Profibrotic signaling has been investigated in different models of Fabry disease in the past. We were able to recapitulate enhanced cleavage of Notch1 in our Fabry podocyte model exhibiting endogenous accumulation of Gb3, previously published in a podocyte cell culture system upon administration of exogenous lyso-Gb3 (Fig. 3A, B) [15]. However, treatment with active enzyme and subsequent Gb3 clearance could not abrogate enhanced Notch1 cleavage and expression of TGF-beta targets in our model (Fig. 3C, D). The activation of profibrotic signaling through α-Gal A deficiency and Gb3 accumulation could very well underlie a self-supplying loop in which an initial stimulus triggers a cascade of events that cannot be stopped by simply abolishing the stimulus. Additionally, accumulation of Gb3 might not be the only pathophysiological stimulus leading to enhanced profibrotic signaling in Fabry disease.

The presented study is limited by its in vitro setup and the experimental way α-Gal A deficiency is achieved. In our model, α-Gal A activity is reduced by inhibition of protein transcription due to small hairpin RNAs. Additional intracellular effects resulting from translation of a defective enzyme and a dysregulated unfolded protein response with ER stress in Fabry patient podocytes cannot be mirrored in our model. Further research should focus on studying reported classical and non-classical GLA gene mutations to investigate additional
or differing effects of ERT and migalastat therapy. Likewise, the analysis of different cell types, preferably the ones that can be kept in culture longer than immortalized differentiated human podocytes [26], would add valuable information to whether the presented results are cell type-dependent or independent and whether they can be influenced by prolonging overall time of treatment with ERT or migalastat.

Our study provides the first in vitro results that supplementation of α-Gal A enzyme activity and subsequent Gb3 clearance are not sufficient to rescue profibrotic signaling and dysregulated autophagy in a podocyte cell culture model of Fabry disease. Long-term studies on the effect of ERT on organ dysfunction have shown that current therapeutic approaches in Fabry disease patients can slow down disease progression but do not significantly reverse damage that has already occurred [27, 28]. Short-term results of chaperone therapy also depict some but not complete improvement of organ function after 1.5 years of treatment [29]. Therefore, distinct but thus far incompletely characterized disease mechanisms may not be reversed with currently available therapy. Since our data points to additional, yet unknown disease mechanisms and a point of no return despite Gb3 clearance, these findings may underscore that early treatment initiation is crucial in Fabry disease patients to prevent irreversible cellular damage [30]. New diagnostic strategies employing the advances in mass spectrometry [31-33] as well as initiating kidney biopsies in Fabry disease patients earlier in the course of the disease have already been suggested [34-37].

The use of (lyso)-Gb3 as a biomarker for Fabry disease activity and for monitoring therapy needs to be critically investigated since additional mechanisms besides Gb3 accumulation-induced cellular damage may be present in Fabry disease [18, 31]. Regular assessments of eGFR, proteinuria and cardiac fibrosis have been reported to be superior prognostic markers in a recent study [38].

**Conclusion**

In conclusion, we provide the first evidence for enzyme replacement therapy-induced and Gb3 clearance in a podocyte cell culture model in vitro. Surprisingly, Gb3 removal failed to improve dysregulated autophagy and a profibrotic phenotype providing first molecular evidence for Gb3-independent disease mechanisms warranting intensified research and identification of additional molecular pathways leading to Fabry disease pathology.

**Acknowledgements**

We acknowledge excellent technical support provided by Beatrix Martiny. The authors have no ethical conflicts to disclose. This work was supported by a Gerok-Position [to F. B.] and an MD scholarship [to L. B.] provided by the Medical Faculty of the University of Cologne. B. S. was supported by the German Research Foundation (SCH1562/6). ERT was kindly provided by Shire, Berlin, Germany. The funding organizations as well as Shire had no influence on the design, outcome or discussed conclusions of the presented study.

The research was designed by F. B. and C. E. K. F. B., L. B., and S. B. conducted the experiments and acquired the data. F. B., L. B. and C. E. K. analyzed the data. F. B., L. B., M. C. L., B. S., T. B. and C. E. K. wrote the manuscript. All authors approved the final version of the paper. F. B. and L. B. contributed equally to this paper.

**Disclosure Statement**

F. B. received travel support of Amicus Therapeutics. C. K. received travel support and speaker honoraria of Amicus Therapeutics, Sanofi-Genzyme and Shire and is an advisory board member of Amicus Therapeutics, Sanofi-Genzyme and Shire.
References


