Modulating ryanodine receptors with dantrolene attenuates neuronopathic phenotype in Gaucher disease mice

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Abstract

Neuronopathic Gaucher disease (nGD) manifests as severe neurological symptoms in patients with no effective treatment available. Ryanodine receptors (Ryrs) are a family of calcium release channels on intracellular stores. The goal of this study is to determine if Ryrs are potential targets for nGD treatment. A nGD cell model (CBE-N2a) was created by inhibiting acid β-glucosidase (GCase) in N2a cells with conduritol B epoxide (CBE). Enhanced cytosolic calcium in CBE-N2a cells was blocked by either ryanodine or dantrolene, antagonists of Ryrs and by Genz-161, a glucosylceramide synthase inhibitor, suggesting substrate-mediated ER-calcium efflux occurs through ryanodine receptors. In the brain of a nGD (4L;C*) mouse model, expression of Ryrs was normal at 13 days of age, but significantly decreased below the wild type level in end-stage 4L;C* brains at 40 days. Treatment with dantrolene in 4L;C* mice starting at postnatal day 5 delayed neurological pathology and prolonged survival. Compared to untreated 4L;C* mice, dantrolene treatment significantly improved gait, reduced LC3-II levels, improved mitochondrial ATP production and reduced inflammation in the brain. Dantrolene treatment partially normalized Ryr expression and its potential regulators, CAMK IV and calmodulin. Furthermore, dantrolene treatment increased residual mutant GCase activity in 4L;C* brains. These data demonstrate that modulating Ryrs has neuroprotective effects in nGD through mechanisms that protect the mitochondria, autophagy, Ryr expression and enhance GCase activity. This study suggests that calcium signalling stabilization, e.g. with dantrolene, could be a potential disease modifying therapy for nGD.
Introduction

Gaucher disease is caused by mutations in GBA1 that encodes lysosomal acid β-glucosidase (GCase) that has glucosylceramide (GC) and its un-acylated form, glucosylsphingosine (GS) as substrates (1–3). Gaucher disease is a common lysosomal storage disease with a frequency of 1/57,000 live births (1). Based on neuronalopathic involvement, Gaucher disease is classified as type 1 (non-neuronalopathic variant) and types 2 and 3 (neuronalopathic variants) (1). Type 2 patients present with acute neurological and pathology within the first 3 to 6 months of life and with death before 2 years of age (1,4). Type 3 patients exhibit subacute neurological signs with a later onset and survival to the 2nd to 4th decade (1,5,6). Two therapeutic strategies have shown clinical efficacy in treating non-neuronalopathic Type 1 Gaucher disease and include: 1) enzyme replacement therapy (ERT) and 2) substrate reduction therapy (SRT). However, the enzyme in ERT cannot cross the blood brain barrier and the FDA approved SRT compounds, miglustat and eliglustat, do not show effective central nervous system (CNS) rescue (7–9). Thus, NgDs are not amenable to current ERT and SRT. More recently, pharmaceutical chaperones and newly developed small molecule substrate reduction agents have been shown to penetrate into the brain. However, these have limited efficacy in slowing disease progression and they do not alter the disease course or prevent death in animal models (10–15). New therapeutic approaches are needed to protect neuronal function as a crucial goal for nGD intervention as has been a recent focus to manage the CNS disease progression.

Accumulated substrates due to defective GCase function cause pathology in the CNS of Gaucher disease. Studies from human patients, animal models and cell models show involvement of multiple pathological pathways in nGD pathogenesis including, inflammation, mitochondrial dysfunction, disrupted calcium homeostasis, altered autophagy/protease function and necrosis (16–25). Disrupted calcium homeostasis, in particular, is a major pathological factor contributing to many neurodegenerative diseases and may lead to neurological deterioration in GD (18,19,25).

Dantrolene is an antagonist of rydnoine receptors (Ryrs) and clinically used for the treatment of malignant hyperthermia and neuroleptic malignant syndrome (26). Ryrs are a class of intracellular calcium channels, expressed in muscles, neurons and other cell types that mediate the release of calcium ions from intracellular organelles, sarcoplasmic reticulum and endoplasmic reticulum (ER). These are essential to a variety of signalling pathways (27). The unique mechanism of dantrolene in blocking intracellular calcium release through Ryrs makes it an attractive potential approach to prevent neuronal dysfunction. Indeed, modulating calcium with dantrolene improves neuronal function in several neurodegenerative diseases including Huntington disease, Alzheimer diseases and kinate-injury model (28–32), suggesting potential clinical utility for nGD.

Here, nGD cell (CBE-N2a) and mouse (4L;C*) models were used to determine the biochemical, histological, and behavioural effects of dantrolene in nGD. The 4L;C* model is a viable analog of human nGD that develops progressive accumulation of substrates and CNS pathology and symptoms (4,18,33,34). 4L;C* mice have been used to investigate pathological mechanisms and test potential therapeutics for nGD (14,18,35). The present study shows dantrolene treatment improves mitochondrial function and protects Ryrs expression in nGD cell and mouse models. Furthermore, dantrolene treatment improved gait, reduced inflammation and prolonged survival in 4L;C* mice, indicating a compelling therapeutic potential for dantrolene in nGD.

Results

Ryrs expression in brain of the nGD mouse model

The nGD mouse model (4L;C*) has the homozygous Gba1 mutation V394L/V394L and a lack of saposin C (33). The 4L;C* mice have substrate (GC and GS) accumulation in the brain that increases with age and is associated with progressive gait impairment, brain pathology, and survival limited to approximately 45 days (14,18,33). Transcriptome analysis by RNAseq of cortex, brain stem, midbrain and cerebellum of 4L;C* mice brains showed a reduction of Ryr1, Ryr2 and Ryr3 mRNAs at 44 days of age (Fig. 1A) (18). Although all three Ryrs are expressed in the brain, Ryr2 is predominantly seen in cardiac muscle (27,36).

Levels of Ryrs protein in neuronal cells and mouse brains were determined for Ryr3 or Ryrin. Immunoblot analysis showed Ryr3 protein in 4L;C* brains was maintained at wild-type (WT) levels at 13 days of age, but was significantly reduced to 27% of WT level at 44 days of age, the end-stage of the disease (Fig. 1B). Immunofluorescence staining with anti-Ryr1 and anti-Ryr3 antibodies showed that Ryrs signal at 44-days in 4L;C* cortex, but was significantly reduced to 27% of WT level at 44 days of age, the end-stage of the disease (Fig. 1C). Neural cells expressing Ryrs were characterized by co-staining of anti-Ryr3 antibody with either anti-NeuN (neuron), anti-GFAP (astrocyte), or anti-O4 (oligodendrocyte) antibodies, respectively. The results showed that Ryr3 was expressed in neurons, astrocytes and oligodendrocytes (Fig 1D). 4L;C* brains had reduced Ryr3 levels in all three cell types (Fig. 1D). These data demonstrate a reduction of Ryrs expression in 4L;C* brains at the disease end-stage implicating Ryrs in nGD pathogenesis.

Dantrolene blocks substrate-mediated calcium efflux from Ryrs in nGD cells

GC mediates ER calcium release (25). To determine if Ryrs are involved in substrate-mediated ER calcium release, a new nGD cell model (CBE-N2a) was generated using N2a cells. N2a is a mouse neuroblastoma cell line that can be differentiated into mature neurons by cAMP and retinoic acid (Supplementary Material, Fig. S1)(37). Differentiated N2a cells were treated with conduritol B epoxide (CBE), an irreversible covalently bound GCase inhibitor. This resulted in significant GC and GS accumulation in CBE-N2a cells, thereby creating an nGD model (Fig. 2A). The effects of Genz-161 (G6), an inhibitor of glucosylceramide synthase to reduce the production of the substrates GC and GS, were evaluated in CBE-N2a cells (14). Treating N2a and CBE-N2a cells with G6 gave GC reductions to 7% and 26% of untreated levels, respectively (Fig. 2A). G6 treated CBE-N2a also had significantly decreased GS (Fig. 2A). Dantrolene is an antagonist of Ryrs and blocks Ry-rmediated ER calcium release (17). Dantrolene alone did not affect GC and GS levels in CBE-N2a cells (Fig. 2A). Co-treatment of G6 and dantrolene of CBE-N2a cells led to a similar degree of GC and GS reduction as G6 alone, indicating a specific effect of G6 on inhibition of substrate accumulation (Fig. 2A). These results demonstrate that G6 inhibits GC production and reduces substrate accumulation in CBE-N2a cells.

Using CBE-N2a cells as an nGD cell model, the effects of dantrolene, ryanodine and G6 on substrate-mediated ER-calcium release were evaluated (Fig. 2B). After CBE-N2a cells were treated with those agents, baseline calcium levels were recorded prior caffeine addition. CBE-N2a cells have higher calcium levels than in N2a cells without caffeine at baseline (Supplementary Material, Fig. S2). Caffeine is a stimulant or
releaser of calcium from ER stores for monitoring GC/GS-mediated ER-calcium levels and the Ryr response (38, 39). After adding caffeine, cytosolic calcium levels were measured every 30s over the duration of 300s using Fura-2 (Supplementary Material, Fig. S2). The magnitude of cytosolic calcium above baseline in the cells with each treatment is shown in Fig. 2B. CBE-N2a cells showed a significant increase in cytosolic calcium levels compared to N2a cells. When CBE-N2a cells were treated

Figure 1. Ryrs expression in 4L;C* mouse brains. (A) Decreased mRNA expression (fold change of WT level) of three Ryrs in 4L;C* brain regions determined by RNASeq analysis (n = 3 mice/group). (B) Immunoblot of brain lysate showed Ryr1 was expressed at normal levels at 13 days of age and significantly decreased by 44 days of age in 4L;C* brain (27% of WT level) compared to WT mice. *P-values were from Student’s t-test, n = 3 mice/group, 2-4 replicates of the experiment. (C) Immunofluorescence staining of brain sections by anti-Ryr1 and anti-Ryr3 antibodies (green) showed decreased Ryr signals in 4L;C* brain including cortex and midbrain regions. Scale bar is 20 µm for all the images. (D) Co-staining of Ryr3 (green) with neural cell markers. NeuN (red) is for neurons. GFAP (red) is for astrocytes. O4 (red) is for oligodendrocytes. Ryr3 signals were detected in neurons, astrocytes and oligodendrocytes in WT midbrains. 4L;C* midbrain had low level of Ryr 3 signals in those cells compared to WT. Scale bar is 20 µm for all the images. DAPI (blue) stained cell nuclei. Images are representative from n = 3 mice for each genotype.
Figure 2. Substrates, calcium levels and mitochondrial function in nGD (CBE-N2a) cells. (A) Substrates levels. GC and GS concentration was significantly increased in CBE-N2a cells compared to N2a cells. Treatment with G6 led to a significant reduction of GC and GS in N2a and CBE-N2a cells, respectively. Dantrolene alone did not significantly affect GC and GS levels in CBE-N2a cells. CBE-N2a cells treated with both G6 and dantrolene showed similar level of GC and GS reduction to G6 alone. (B) Cytosolic calcium levels. Differentiated N2a cells were treated with CBE (2 mM) for 5 days. CBE-N2a cells were further incubated for 5 days with ryanodine (Ryan, 10 μM), dantrolene (Dan, 12.5 μM) or G6 (0.8 μM), or combinations of Dan/G6 or Ryan/G6. Cytosolic calcium level was measured using Fura-2 as reporter after adding caffeine. Relative cytosolic calcium level [Fura-2 ΔF/F (340/380-510 nm)] normalized to mg protein in cells shows the magnitude of cytosolic calcium above baseline (see
with G6, the calcium levels were significantly lower than that in untreated CBE-N2a cells (Fig. 2B), suggesting substrate-mediated calcium release. In dantrolene-treated CBE-N2a cells, calcium levels were significantly reduced compared to untreated CBE-N2a and approached levels observed in control N2a cells (Fig. 2B). Consistent with dantrolene, CBE-N2a cells treated with ryanodine also showed a significant reduction in calcium levels (Fig. 2B). Co-treatment of G6 and dantrolene or G6 and ryanodine also showed a significant decrease in calcium levels. These results demonstrate that substrate-mediated calcium release occurs through Rys.

**Dantrolene protects mitochondrial function in nGD cells**

To determine if substrate-mediated ER calcium efflux promotes mitochondrial dysfunction, the effects of dantrolene and G6 on mitochondrial function were evaluated in CBE-N2a cells. Treated cells were plated on a Seahorse plate to assay mitochondrial function reported as Oxygen Consumption Rate (OCR) that includes the following parameters: ATP production, basal respiration, and maximal respiration. CBE-N2a cells showed a significant reduction in OCR as evidenced by approximately 50% in all the parameters, including rate of ATP production, basal respiration, and maximal respiration, compared to N2a cells, indicating reduced mitochondrial function in this nGD cell model (Fig. 2C). Dantrolene treatment significantly improved OCR in CBE-N2a cells (Fig. 2C). G6 treatment also resulted in significant increases in OCR in CBE-N2a cells (Fig. 2C). The greatest increase in OCR was achieved when CBE-N2a cells were treated with both dantrolene and G6 (Fig. 2C). These results demonstrate that both reduction of substrate levels by G6 or antagonizing Rys by dantrolene have protective effects on mitochondrial function in nGD cells. Combining G6 and dantrolene resulted in more improvement than either dantrolene or G6 alone.

**Dantrolene treatment mitigates the neuropathic phenotype in the nGD mouse model**

The in vivo efficacy of dantrolene on nGD was evaluated in the 4L;C* mouse model. 4L;C* mice were treated with dantrolene by intraperitoneal (IP) injection at 10 mg/kg on each of three days per week, starting from postnatal day 5. Because dantrolene formulation contains mannitol, two control groups were included in the study: untreated 4L;C* mice and 4L;C* mice injected with mannitol at 30 mg/kg (same level of mannitol in dantrolene formulation) on each of three days per week. Dantrolene treated littermates (4L;WT) were also included as controls. Littermates do not show neurological impairments and live beyond 80 days.

Gait was measured at 30 and 40 days of age. 4L;C* mice typically develop a duck-like walking with paralysis of the hind limbs (18,33). Untreated 4L;C* mice made significantly shorter strides and had a wider base width at 30 and 40 days of age compared to 4L;WT littermates (Fig. 3A). Dantrolene treatment significantly reduced base stride width at 30 and 40 days of age compared to age-matched untreated 4L;C* mice (Fig. 3A). Dantrolene treated 4L;C* mice made significantly longer strides, a 30% increase compared to untreated 4L;C* mice at 40 days of age (Fig. 3A). However, the gait in dantrolene treated 4L;C* mice did begin to deteriorate after 50 days of age (data not shown). The life span of dantrolene treated 4L;C* mice was also significantly extended compared to untreated 4L;C* mice (P = 0.0012) as determined by Kaplan-Meier analyses and the Mantel-Cox test (Fig. 3B). The treatment prolonged survival in 4L;C* mice by 12.7% compared to untreated 4L;C* mice (Fig. 3B). To determine if mannitol would affect the phenotype, one group of 4L;C* mice was injected with mannitol and gait and survival were evaluated. Mannitol-4L;C* mice showed no changes on base width, but they had a 10% increase (P < 0.05) in stride length compared to untreated 4L;C* mice at 40 days of age. However, this improvement is significantly less than the 30% improvement in dantrolene treated 4L;C* mice (Supplementary Material, Fig. S3). No differences were observed in life span between untreated and mannitol-treated 4L;C* mice (data not shown), indicating a specific effect of dantrolene on survival. In the following biochemical and histological studies, untreated 4L;C* brains were used as a control for dantrolene treatment effect. Dantrolene treatment did not affect body or organ weights in treated mice. These results demonstrate that dantrolene significantly improved gait, delayed motor function decline, and extended the life span of 4L;C* mice.

CNS inflammation was determined by staining tissue sections with an anti-CD68 antibody. CD68 is a marker for activated microglia and macrophages. Positive CD68 signals (brown) were detected in most brain regions of untreated 4L;C* mice (Fig. 3C). With dantrolene treatment, the CD68 signal showed a significant decrease of 63% of that in untreated mice, indicating attenuation of CNS inflammation by dantrolene (Fig. 3C).

Brain mitochondrial function was determined by Seahorse assay using isolated mitochondria from WT, untreated and dantrolene treated 4L;C* brains at 40 days of age. ATP production rate in 4L;C* brain mitochondria was significantly reduced compared to WT (Fig. 3D). Dantrolene treated 4L;C* mice showed significantly increased mitochondrial ATP production rate in the brain, 77% of WT level compared to 37% in the untreated 4L;C* brain (Fig. 3D) indicating a protective effect on mitochondrial function in nGD by dantrolene.

Altered autophagy is also evident in nGD and 4L;C* mouse brains (20,33,34). LC3-II is a marker for autophagy activity and was evaluated in dantrolene treated 4L;C* brains compared to untreated 4L;C* and WT brains to determine the effect of the treatment on autophagy function. LC3-II is a membrane bound form of LC3. Elevated LC3-II levels indicate altered autophagy. By immunoblot analysis, LC3-II was undetectable in WT brain. Increased LC3-II levels were observed in untreated 4L;C* brains whereas dantrolene treated 4L;C* brains had significantly reduced LC3-II levels at 50% of untreated level, but still more than WT level (Fig. 3E). This result indicated that dantrolene treatment partially prevented alterations in autophagy in 4L;C* brains.

Neurodegeneration in 4L;C* brains were evaluated by counting NeuN positive mature neurons in multiple brain regions (cortex, brain stem, midbrain and cerebellum) from whole
Figure 3. Dantrolene treatment in 4L;C* mice. (A) Gait analysis. Left stride (Left panel), right stride (Middle panel), base width (Right panel). The mice were subjected to two to three tests at 30 and 40 days of age. Dantrolene (Dan) treatment significantly increased left and right strides in 4L;C* mice at 40 days of age, and reduced base width at both 30 and 40 days of age compared to untreated 4L;C* mice. Littermate (4L;WT) mice that have no phenotype were used as normal controls in the analysis. Data were analysed by Student’s t-test. (B) Life span. The survival rate of dantrolene treated 4L;C* mice (blue) was significantly increased compared to untreated 4L;C* mice (orange). Median survival days is 50 days or 44 days for treated or untreated 4L;C* mice, respectively. The life span of dantrolene treated 4L;C* mice was prolonged by 12.3%. Littermate (4L;WT) control mice (black) had normal life span. Data are presented as Kaplan-Meier curve analysed by Mantel-Cox test. (C) CNS-inflammation. Positive CD68 staining (brown) in microglial cells indicate inflammation in 4L;C* brain. Compared to untreated 4L;C*, the CD68 signal was significantly reduced in dantrolene treated 4L;C* brains. The representative image for each group is shown. CD68 signal intensity in brain sections was quantitated by NIH image J and presented as % of untreated 4L;C* level. P-value was from Student’s t-test (n = 2–3 mice/group). (D) Mitochondrial ATP production rate. 4L;C* brain had 37% of ATP production rate (pmol/min/mg mitochondrial protein) compared to WT brains. Dantrolene treatment on 4L;C* mice improved ATP production to 77% of WT level. One-way ANOVA with post-hoc Tukey test (P < 0.05), n = 3 mice/group, 6 replicates/sample/assay, duplicate assays. (E) Immunoblot of LC3. LC3-II is barely detectable in WT brain. LC3-II levels were increased in 4L;C* cerebrum compared to WT. Dantrolene treated 4L;C* cerebrum showed significantly reduced level of LC3-II compared to untreated 4L;C*. One-way ANOVA with post-hoc Tukey test (P < 0.05), n = 3 mice/group, duplicate experiments.
sagittal sections (Fig. 4 and supplementary Material, Fig. S4). 4L;C’ brains had significantly fewer NeuN positive cells (60–80% of WT NeuN+ cells) than WT brains. Treatment with dantrolene resulted in an increase in NeuN positive cells to >80% of WT NeuN+ cells. Decreased NeuN positive cells were observed in cortex, cerebellum, midbrain and brain stem of 4L;C’ mice. All of those regions showed an increase in neurons with dantrolene treatment compared to the untreated 4L;C’ mice (Fig. 4). These results indicated dantrolene treatment reduced neurodegeneration in 4L;C’ mice.

**Dantrolene protects RyR, calmodulin and CAMK IV expression**

The effect of dantrolene on RyR protein expression was determined in nGD cells and 4L;C’ mouse brains (Fig. 5). Immunoblot analysis showed that Ryr3 protein was reduced to 47% of the WT level in CBE-N2a cells. In dantrolene-treated CBE-N2a cells, levels of Ryr3 were increased to 76% of WT level (Fig. 5A). In 4L;C’ mice, Ryr3 was profoundly decreased to just 9% of WT levels in the cerebrum at 44 days of age (Fig. S5B). Dantrolene treatment in 4L;C’ mice significantly increased Ryr3 levels to 77% of WT in the cerebrum (Fig. S5B). Immunofluorescence using an anti-Ryr3 antibody confirmed the increase in Ryr signals. In dantrolene treated 4L;C’ midbrain (94% of WT) and brain stem (79% of WT) regions (Fig. 5C). Ryr 3 signals were increased in neurons, astrocytes and oligodendrocytes in dantrolene treated 4L;C’ mouse brains (Supplementary Material, Table S1).

In the calcium/calmodulin-dependent protein kinase pathway, calmodulin, a calcium sensor protein, binds to calcium and promotes a calcium-dependent kinase activity including CAMK IV (Calcium/calmodulin-dependent kinase type IV) for gene expression and biological functions (40). Calmodulin and CAMK IV are potential modulators for Ryr expression. Decreased CAMK IV mRNA and increased calmodulin 2 mRNA have been shown in 4L;C’ brains using RNAseq analyses (18). Expression levels of calmodulin and CAMK IV in 4L;C’ brains were determined by immunoblot. CAMK IV protein level was reduced to 61% of WT level in untreated 4L;C’ cerebrum. With dantrolene treatment, the expression of CAMK IV was increased by 28% (Fig. S5D). In this study, calmodulin was analysed using an anti-Cam antibody that reacts to a class of calmodulins including calmodulin 2. Calmodulin levels were 2 fold above WT levels in 4L;C’ brains and were normalized to WT levels with dantrolene treatment (Fig. 5D). These data suggest that modulating cytosolic calcium levels with dantrolene has protective effects on Ryrs expression as well as proteins involved in the calcium-calmodulin dependent signalling pathway.

**Dantrolene treatment effects on GCase function**

Modulating ER calcium release promotes chaperone enhancement of GCase stability and trafficking to the lysosome, thereby improving GCase activity (17,41,42). The effect of dantrolene on GCase activity was evaluated in mouse fibroblast cells from Gba1 mutants homozygous for V493L/V394L (4L;4L), D409H/D409H (9H/9H), and D409V/D409V (9V/9V), 4L;C’ and WT mice. Dantrolene-treated 4L;4L, 4L;C’, 9H/9H and WT fibroblasts showed significantly increased GCase activity compared to the untreated cells (Supplementary Material, Fig. S5A). Dantrolene treated 9V/9V cells did not show an increase in GCase activity. This result showed that dantrolene promoted GCase activity in select mutants, e.g. V394L and D409H, and WT enzyme.

In vivo effects of dantrolene on GCase function (activity and substrate levels) were analysed in dantrolene treated mouse brains at 44 days of age. The cerebrum was used for GCase activity and cerebellum was used for substrates analysis. Compared to untreated 4L;C’, dantrolene-treated 4L;C’ cerebrum showed a 1.2 fold increase in GCase activity which is significantly higher than untreated 4L;C’ (Supplementary Material, Fig. S5B). However, substrate (GC and GS) concentrations were not reduced in the treated 4L;C’ cerebellum (Supplementary Material, Fig. S5C). Co-staining for GCase and Lamp1 of mouse brain sections showed a 1.7-fold increased colocalization of GCase within the lysosomes of dantrolene-treated 4L;C’ mice compared to untreated 4L;C’ brains. However, this level is < 40% of Pearson coefficient for WT GCase (Supplementary Material, Fig. S5D), indicating >40% mature mutant enzyme trafficked to the lysosomes in dantrolene-treated brains. These results suggest that the effect of dantrolene on GCase activity with the regimen used in this study was not sufficient to translate into significant hydrolytic function for clearance of excess substrates.

**Discussion**

Currently, there are no effective treatments available for nGD. The goal of the present study was to test the therapeutic potential of a ryanodine receptor antagonist in cell and animal models of nGD. Our results show that dantrolene treatment improves mitochondrial function, protects basal autophagy, delays abnormal gait, and most importantly prolongs survival in nGD mice. Dantrolene prevented the decreased expression of Ryrs in nGD cells and mice, and normalized expression of calmodulin and CAMK IV, key molecules in the calcium/calmodulin-dependent regulation pathway, in mice (40,43,44). Furthermore, dantrolene reduced inflammation and the loss of NeuN+ neurons in mice indicating a compelling neuroprotective effect of treatment. This study shows that modulating ER calcium efflux through Ryrs by dantrolene has therapeutic value for nGD. Neurprotection by dantrolene is likely through several mechanisms including protection of mitochondrial function, Ryrs expression and autophagy, and enhancement of GCase activity.

Abnormal calcium homeostasis has been implicated in GD (25,45,46). Accumulation of GC causes excess calcium efflux from ER, specifically through Ryrs in neurons in GD (38). Sustained increase in cytosolic calcium accounts for the abnormal cellular function in many lysosomal storage diseases including GD (47), although these changes are not fully understood. Using CBE-N2a cells treated with G6 to inhibit glucosylceramide production, or dantrolene to inhibit calcium efflux through Ryrs, we confirmed that increased cytosolic calcium efflux through Ryrs is mediated by excess substrates, supporting previous findings (25). Most importantly, this new nGD cell model allows further investigation of altered cellular function resulting from disrupted calcium homeostasis.

Ryr protein down-regulation in nGD cells and mouse brains is a novel finding. All three Ryrs are expressed in brain. Ryr1 is predominantly in the skeletal muscle and Ryr2 is mainly in cardiac muscle, whereas Ryr3 is expressed at a low level in a variety of tissues including brain (27,36,48). Decreased expression of Ryrs has been reported in a mouse model of Alzheimer disease that also shows aberrant cellular calcium homeostasis (49). Ryrs protein levels in this study were determined primarily on Ryr3 with a high quality antibody. Ryr3 was shown to be ubiquitously
expressed in the mouse brain. Reduced staining for Ryr3 was found in neurons, astrocytes and oligodendrocytes in 4L;C* mice at end-stage. Normal levels of Ryr3 protein were expressed in 4L;C* brains at 13 days of age, but were reduced to 27% of WT level by the end stage (44 days), suggesting Ryr down-regulation is associated with disease progression and increased substrate accumulation. Thus, Ryr down-regulation is likely caused by substrate accumulation. In 4L;C* mouse brain, substrate levels were higher than that in WT even prior to 13 days, and continued to increase with age (Supplementary Material, Fig. SSE). Excess substrate can cause ER-calcium efflux and increases in cytosolic calcium (25). The sustained increase of cytosolic calcium could lead to a feed-back regulation, via calcium dependent transcription regulation pathways, and reduce the expression of Ryrs thereby preventing further calcium efflux from the ER. Our results show that blocking ER-calcium release through Ryrs with dantrolene protects Ryr expression, supporting this feed-back regulation mechanism. Early intervention

Figure 4. NeuN positive neurons in brain regions. WT, dantrolene (Dan) treated 4L;C* and untreated 4L;C* brain sagittal sections from 44 day old mice were stained with anti-NeuN antibody. (A) Compared to WT cortex, cerebellum, midbrain and brain stem, 4L;C* mice had reduced NeuN positive cells (green) in those regions. Representative images from each group are shown. (B) Dantrolene treated 4L;C* mice had significantly more NeuN positive cells than untreated 4L;C* in each region. In the graph, NeuN positive cell counts in each group are shown as a percentage of WT for each brain region. Data were analysed by One-way ANOVA with post-hoc Tukey test (P < 0.05), n = 4 images/section, 2 sections/mouse, 3 mice/group.
Figure 5. Ryr expression in dantrolene treated 4L;C* brain. (A) Immunoblot of Ryr3 in CBE-N2a cells. Ryr3 protein level was lower in CBE-N2a than N2a cells, and increased in dantrolene treated CBE-N2a cells. (B) 4L;C* cerebrum showed significantly reduced Ryr3 protein at 9% of WT level. In dantrolene treated 4L;C* cerebrum, Ryr3 protein level was significantly increased compared to untreated 4L;C*. (C) Immunofluorescence staining of Ryr3. 4L;C* midbrain and brain stem showed reduced Ryr3 (green) signal at 49% or 34% of WT level, respectively. In dantrolene treated 4L;C* brain, Ryr3 signal was increased to 94% in midbrain and 79% in brain stem of WT level. DAPI (blue) stained cell nuclei. Scale bar is 20 μm for all the images. (D and E) CAMK IV and calmodulin (CAM). 4L;C* cerebrum showed decreased level of CAMK IV (D) and increased level of CAM (E) compared to WT. Dantrolene treatment normalized expression of CAMK IV and CAM to nearly WT level. One-way ANOVA with post-hoc Tukey test (P < 0.05), n = 2-3 mice, 2-4 replicates of the experiment.
before Ryrs level reduction would be critical to achieve maximal protection.

Expression of Ryrs can be regulated by the calcium-signalling pathway. In this pathway, calcium and calmodulin binding leads to basal CAMK IV activity which can regulate the expression of Ryrs. Calmodulin bound to calcium is required for the initiation of CAMK-CREB cascade for gene regulation (43,44,50,51). Like many calcium binding proteins, Ryrs have CREB binding elements (52), therefore, their expression is likely regulated by CAMK IV-CREB signalling (53,54). Enhanced expression of calmodulin in 4L;C* brains may reflect increased cytosolic calcium. Reduced CAMK IV mRNA and protein levels in 4L;C* brains suggest that less CAMK IV is available for gene regulation, or decreased transcriptional activity (18,52–54). Excess substrate-mediated calcium release in nGD brains could hinder calcium/calmodulin kinase signalling dependent CREB phosphorylation and lead to reduced Ryrs expression. When nGD mice are treated at an earlier stage with dantrolene (starting at postnatal 5 days) expression of CAMK IV and calmodulin were normalized, highlighting their involvement in the modulating Ryrs expression in nGD. Therefore, protection of Ryrs expression with dantrolene should benefit nGD.

Ryrs down-regulation was concomitant with disease progression. Importantly, the expression of Ryrs was protected by dantrolene. This suggests a role for Ryrs in nGD pathogenesis and an attractive target for therapy. Ryrs on ER are key for the regulation of intracellular calcium and an important calcium modulator for neuronal function (49,55). Both ryanodine and dantrolene are antagonists for Ryrs. Ryanodine is toxic to cells making it unsuitable for in vivo studies. In contrast, dantrolene is FDA approved for treatment of malignant hyperthermia (56), readily available, and is distributed in tissues e.g. muscle and liver. It also penetrates the blood-brain barrier to allow for access into the brain (57). Dantrolene is specific to Ryr1 and Ryr3 (58). Blocking Ryrs with dantrolene is associated with improvement in learning and memory performance and protection of synaptic function in neurons in Alzheimer and Huntington disease models (28,32). The significant attenuation of neuromaptic phenotype in 4L;C* mice with dantrolene also suggests a clinical benefit for nGD. Impaired calcium homeostasis is a common pathologic factor in glycosphingolipid storage diseases. Specific substrate accumulation causes reduced lysosomal calcium in Niemann-Pick type C disease and decreased ER calcium in Sandhoff and Niemann-Pick A diseases (59–61). Thus, modulating Ryrs in order to normalize calcium homeostasis could have therapeutic potential for glycosphingolipid storage diseases.

Targeting Ryrs to modulate calcium homeostasis likely acts through multiple mechanisms to achieve neuroprotection in nGD, including protection of mitochondrial function and autophagy, normalizing gene expression/regulation (Ryr), and rescuing mutant GCase activity. Mitochondrial dysfunction is a well-documented pathological feature in nGD (18,62). Inhibition of GCase function by CBE leads to defective mitochondrial function in a human dopaminergic cell line (23). In chronic nGD mouse brains decreased mitochondrial ATP production and oxygen consumption are associated with protein aggregation (e.g. synuclein and amyloid precursor protein) on mitochondria (21). Impaired respiratory chain and mitochondrial membrane potential has been reported in primary neurons and astrocytes of acute nGD mice (23,62). Reduced mitochondrial function is also a significant feature in 4L;C* mice and in CBE-N2a cells (18). ER and mitochondria are physically and functionally linked with altered calcium homeostasis promoting mitochondrial dysfunction and disruption of the mitochondrial membrane potential (63–65). Furthermore, the role of Ryr in ER calcium and mitochondrial function is supported by a study of the conditional knockout of cardiac Ryr2 in mice (66). Here, depletion of Ryr2 leads to reduced mitochondrial calcium and oxidative metabolism (66). Therefore, modulating calcium homeostasis through Ryrs would have benefits for mitochondrial function. Indeed, in both CBE-N2a cells and 4L;C* mice, dantrolene treatment significantly improved the rate of mitochondrial oxygen consumption and ATP production. These data support the notion that Ryr antagonism may have protective effects on mitochondrial function in nGD.

In many lysosomal storage diseases, defective lysosomal enzyme/protein function interrupts the fusion of the lysosome with the autophagosome for proper degradation of macromolecules (67–70). Abnormal autophagy, either increased or decreased, has been demonstrated in nGD cell and animal models, and patient brain samples (5,20,22). Little is known, however, about the mechanisms regulating autophagy in nGD, although it has been shown that autophagy in general is regulated by cytosolic calcium (71,72). Increased calcium load induced by starvation is known to induce autophagy (71). In this study, we used LC3-II to monitor autophagy. LC3 (microtubule-associated protein 1 light chain 3) has two forms, LC3-I is distributed within the cytoplasm and nucleus and LC3-II conjugates to phosphatidylethanolamine on the autophagosomal membrane. Increased LC3-II reflects an induction of autophagy in response to cellular stress. In the nGD mice, dantrolene treatment significantly reduced LC3-II to WT levels suggesting that autophagy in nGD is regulated by the ER-calcium flux through Ryrs. Thus, modulating cytosolic calcium levels could be a target for maintaining basal autophagy and normal cell function.

Modulation of ER-calcium can regulate calcium-dependent chaperones for GCase protein folding control and proper degradation (17,41,42,73). Inhibition of Ryrs reduces intraluminal calcium levels and enhances the expression of ER chaperone proteins that promote protein folding. Specifically, for mutated GCase L444P in fibroblasts, this leads to increased folding and trafficking to the lysosome (17,74). Modulating ER proteostasis by antagonizing Ryr with dantrolene has shown improvement in Niemann-Pick disease by enhanced type C1 protein level and attenuated the cholesterol and sphingolipids in Niemann-Pick type C disease fibroblasts (73). In addition to Ryrs, blocking L type calcium channels (LTCC) also induces chaperone activity on mutant GCase (75). Although diltiazem, an LTCC blocker, improves expression of cellular chaperones and folding of the mutated enzymes in mouse and human fibroblasts (75,76), it fails to translate to similar changes in animal models of GD (76). In the present study, dantrolene was able to enhance GCase activity in select homozygous Gba1 mutations (V493L/V394L, D409H/D409H), 4L;C* and WT mice fibroblasts. In 4L;C* mouse brains dantrolene enhanced mutant GCase activity and lysosomal localization, however, the level was not sufficient to prevent substrate accumulation. Increasing administration to a daily regimen may achieve better efficacy on GCase hydrolytic function.

This study highlights a profound pathogenic role for Ryrs in nGD and supports a novel strategy to preserve calcium homeostasis in the early stages of nGD and thereby slow disease progression. Our results demonstrate that targeting Ryrs to modulate calcium homeostasis has neuroprotective potential in nGD by protecting mitochondrial function, normalizing expression of Ryrs, protecting autophagy, enhancing mutant GCase activity and reducing inflammation and neuron loss.
Materials and Methods

Materials

The following were from commercial sources: N2a (ATCC®-CCL™-131, VA); CBE (EMD Millipore, Bedford, MA); Dantrolene, retinoic acid, dbcAMP and caffeine (Sigma, St. Louis, MO); M-Per Mammalian Protein Extraction Reagent, SuperSignaL™ West Dura ECL detection kit, Fierce BCA protein assay kit (ThermoFisher Sci, Waltham, MA). Antibodies: Ryr1, Ryr3, Map2 and Nestin (EMD Millipore, Bedford, MA); CAM (calmodulin) (Santa Cruz Biotechnology, Dallas, Texas); LC3 (Novus, Littleton, CO); CAMK IV (calcium/calmodulin-dependent kinase IV) (Abcam, San Francisco, CA); CD68 (BD Biosciences, Franklin Lakes, NJ); NuPAGE Gels (3-8%, 4-12%, 10%, 16%) and Bis-Tris Buffer, Sodium Acetate Buffer, SeaBlue Protein Standard Plus, PVDF membrane, iBlot transfer kit, and iBind Cards (Life Technologies. Carlsbad, CA); AP colour reagent and AP Conjugate Substrate Kit (Bio-Red, Hercules, CA); VECTASHIELD mounting medium containing DAPI (Vector, Burlingame, CA); Glass chamber slides (LAB-TEK, Rochester, NY). Genz-161(G6) (GenScript ToxinSenor). The cells were from same passage and treated at the same time for all the experiments.

Cell culture and treatment

N2a cells were maintained in DMEM/10% FBS (Gibco) medium. Neuronal differentiation of N2a was carried out in the differentiation medium containing 0.5% FBS, 10 μM retinoic acid, 50 ng/ml GDNF (Alomore lab) and 1 mM dbcAMP in DMEM for 3 days (37). The differentiated N2a cells were treated with 2 mM CBE for 5 days in the differentiation medium. Dantrolene (12.5 μM), ryanodine (10 μM) or G6 (0.8 μM) was added to the CBE-N2a cells and cultured for 5 days. The medium with the reagents was changed every 2 days. The differentiation status of the neuronal cells was confirmed by anti-Map2 (mature neuron) or anti-Nestin (undifferentiated neuronal cells) antibody staining. Cell viability after each drug treatment was measured by CellTiter-Fluor™ Cell Viability assay (Promega). The dose for each compound was determined from CellTiter-FluorTM Cell Viability assay with >95% viability and endotoxin test of -0.05 EU/ml (GenScript ToxinSenor). The cells were from same passage and treated at the same time for all the experiments.

Calcium level measurement

Intracellular calcium levels were determined using the Fura-2 QBT Calcium Kit (Molecular Devices, Sunnyvale, CA) on 96 well plates by M5 SpectraMax plate reader (Molecular Devices). N2a cells were seeded in each well (20,000/well) with differentiation medium (0.5% FBS, DMEM, 10 μM retinoic acid, 1 mM dbcAMP) for 72 h (37). The differentiated N2a cells were cultured with DMEM medium with 10% FBS containing CBE (2 mM) for 5 days, followed by treatment with different combinations of dantrolene (12.5 μM), ryanodine (10 μM) and G6 (0.8 μM) for a further 5 days. The cell medium was replaced every two days. The treated cells were incubated with Fura-2 QBT dye at 37 C for 1 h for Fura-2 QBT uptake (Fura2 QBT® calcium kit protocol, Molecular Devices). Caffeine (10 mM) was used as an ER calcium efflux stimulant in the assay (38,39). The plate reader setting was 340/380 nm (excitation) and 510 nm (emission). Baseline calcium levels were measured prior to caffeine addition (~30 s). Cytosolic calcium levels were recorded every 30 s for duration of 300 s after addition of caffeine. Relative cytosolic calcium levels were determined with the values at 300 s endpoint minus 0 s baseline. Relative cytosolic calcium level [Fura-2 ΔF/F (340/380-3510 nm) for control or treated cells was calculated by Fura-2 fluorescence unit (FU) measured at 340nm/510nm (ΔF = RFU/340 at 300 s end point minus baseline RFU/340 at 0 s) divided by FU at 380nm/510nm (F = RFU/380 at 300 s - baseline RFU/380 at 0 s) (Fura2 QBT® calcium kit protocol, Molecular Devices) and normalized to protein level in cells. Data were collected and analysed by SoftMax Pro software (Molecular Devices) and graphical charts were generated by Graphpad Prism 6.0.

Mice and treatment

4L,C* mice were generated by backcrossing of V394L/V394L Gba1 (4L) and saposin C-/- (C*) homozygosity as described previously (33). The strain background of 4L,C* mice was C57BL/6J:129SvEv. Strain- and age-matched WT mice and non-4L,C* littermates (no neurological phenotype) were included as controls. All mice were housed under pathogen-free conditions in an animal facility according to IACUC approved protocols at Cincinnati Children’s Hospital Medical Center (CCHMC).

The 4L,C* mice and non-4L,C* littermates were treated with Dantrolene sodium (Reventon®) US WorldMeds, Louisville, KY starting at postnatal day 5 by IP injection at 10 mg/kg, three days per week. Mouse body weight was recorded before each injection. Control mice received vehicle (mannitol, APP Pharmaceuticals, Schaumburg, IL) injections or no injection at all. The mice were monitored for survival and gait impairments during the treatment.

Gait analysis

Progression of the neurobehavioural phenotype was assessed by gait analysis to determine sensorimotor function (77,78). Mice were trained to walk through a narrow alley leading into their home-cage. Once trained, paper was placed along the alley floor and the hind paws of each mouse were brushed with non-toxic paint. The mice were then placed at the beginning of the alley. As they walked into their home-cage they left their paw prints on the paper underneath. Stride length and width were determined by measuring the distance between hind paw prints. The 4L,C* and control mice, untreated and non-4L,C* littermates were tested for gait at 30 and 40 days of age.

RNAseq analyses

Expression of Ryr mRNAs in 4L,C* brain regions (cortex, cerebellum, midbrain and brain stem) was analysed by RNAseq as described previously (18).

Immunohistochemistry and immunofluorescence

Mouse brains were collected after transcardial perfusion with saline. Half of the brain (sagittal cut) was fixed in 4% paraformaldehyde (PFA) for processing as frozen blocks. CD68 monoclonal antibody staining was performed as previously described (33) using The BenchMark XT IHC/ISH Staining Module (Ventana Medical System, Tucson, AZ) at CCHMC Pathology Research Core. Fixed sections were counterstained with hematoxylin. The sections were then scanned by AperioImageScope v2. The CD68 signal was quantified using Image J FIJI (79).

Immunofluorescence staining was performed on PFA fixed brains. The brain sections were incubated in 0.3% Triton X-100 for 30 min, and treated with 50 mM NH4Cl in 1xPBS for 15 min followed by 1xPBS wash. The sections were blocked for 1 hr at RT in
Blocking buffer (10% goat serum and 0.4% Triton X-100 in PBS). Rabbit anti-Ryr 1 antibody (1:100) was diluted in the blocking solution and incubated overnight at 4°C. After washing in PBS (3x10 min), the secondary antibody goat anti-rabbit conjugated with Alexa Fluor® 488 (1:1500) in blocking buffer was applied to the sections and incubated for 2 hrs at RT. Mouse anti-Ryr 3 antibody (1:100) and goat anti-mouse conjugated with Alexa Fluor® 488 (1:1500) were used for Ryr 3 detection. For co-staining of neuronal cells with Ryr3, mouse anti-NeuN antibody (1:500, Millipore), mouse anti-GFAP (1:100, Millipore) or mouse anti-O4 (1:100, Millipore) with goat anti-mouse conjugated Alexa Fluor® 594 (1:1000) were used for NeuN, GFAP, or O4 detection, respectively. Rabbit anti-Ryr 3 (1:250, Millipore) with goat anti-rabbit conjugated Alexa Fluor® 488 (1:1000) were used for Ryr3. Lysosomal localization of GCase was determined using rabbit anti-mouse GCase (1:50) antibody generated in our lab (15) with goat anti-rabbit conjugated Alexa Fluor® 488 (1:1000) for mouse GCase detection. Rat anti-Lamp1 (1:100, D78) and rabbit anti-lgG conjugated Texas Red (1:1000, Abcam) were used for Lamp1 detection. Sections were mounted with VECTASHIELD mounting medium containing DAPI (Vector H1200). Fluorescence signals were visualized and captured by Zeiss Axiovert 200 M microscopy equipped with an Apotome. Pearson correlation coefficient software is incorporated in Apotome microscope and used to analyse the colocalization of two signals (GCase and Lamp1) in cells. The average Pearson correlation coefficient number (from 0 to 1) was derived from the cells in multiple images (21). For counting NeuN+ neurons, mouse anti-NeuN antibody (1:500, Millipore) was applied to brain sections followed by secondary antibody conjugated with FITC. The images of NeuN+ cells in brain sections were acquired by Nikon C2 plus confocal microscope with NIS image stitching function for large image capture in ND acquisition. All NIS images in 32 bit were converted into 16 bit format and analysed by ImageJ FIJI. NeuN+ cells were counted from whole brain images and from multiple images (3.5 × 2.0 mm) in each brain region (cortex, midbrain, cerebellum and brain stem).

For cell culture immunostaining differentiated N2a cells were seeded on chamber slides and stained with anti-Map2 antibody (1:200) followed by the protocol described above except for using 3% BSA in PBS as blocking buffer. Fluorescent images were processed by Zeiss Apotome 200M.

Immunoblot
Half of mice brain (sagittal cut) tissues were homogenized in M-PER Mammalian Protein Extraction Reagent and subjected to electrophoresis. Ryr proteins (~500 kD) in tissues lysate were separated on 3–8% NuPAGE gel running at 90V for 12 h at 4°C in acetate buffer. β-actin was resolved on 4–12% Bis-Tris gel. The proteins were transferred to PDVF membrane using iBlot 2 gel transfer device following manufacturer’s instructions. The blots were incubated with anti-mouse Ryr3 monoclonal antibody (1:250) or anti-β-actin (1:500) overnight at 4°C in 1.5% BSA/1.5% milk/PBS buffer. LC3, CAMK IV and calmodulin were separated on 16% Tris-glycine gel (Novex) and detected by anti-LC3 (1:1000), anti-CAMK IV (1:500) or anti-CAM (1:1000), respectively. The signals were detected with either ECL detection reagent or AP Conjugate Substrate Kit according to manufacturer’s instructions. Band intensities on immunoblots were quantitated by Image J (NIH, Baltimore, VA).

Seahorse mitochondrial function assay
Cultured N2a cells treated with or without CBE, dantrolene or G6 were transferred to a XF96 assay plate at 10,000 cells/well and allowed to grow overnight in DMEM medium containing 10% FBS with each compound as described above in Cell culture and treatment section. Cell numbers were determined using a haemocytometer. Prior to mitochondrial function assay on XF96 Extracellular Flux Analyzers (Seahorse Biosciences), the cells in the wells of XF96 assay plates were washed by gently adding warm assay medium to the side of each well. The plates were transferred to a 37°C incubator without CO2 for 30–60 min before the addition of XF assay medium (Seahorse Bioscience) containing 25 mM glucose and 1 mM pyruvate. After calibration of the Analyzer, the plate was sequentially injected with A: 25 μl of 8 μM oligomycin, B: 25 μl of 27 μM cyanide-p-trifluoromethoxyphenylhydrazone (FCCP), and C: 25 μl of 50 μM antimycin A. Oxygen consumption rate (OCR) was measured, and the data were analysed using the XFe Wave software. The respiration parameters were calculated by subtracting the average respiration rates before and after the addition of the electron transport inhibitors: oligomycin, FCCP and antimycin A. The parameters include basal respiration (baseline respiration minus antimycin A post injection respiration), ATP production (baseline respiration minus oligomycin post injection respiration), and maximal respiratory (FCCP stimulated respiration minus antimycin A post injection respiration). Cell mitochondrial function presented as respiration parameters that were normalized by cell numbers (29).

Brain mitochondria were isolated from 4L;C*, WT, and dantrolene treated 4L;C* mice as described with slight modifications (21,30). Mouse brain tissues were digested with trypsin for 30 min on ice followed by homogenization on the gentle MACS Dissociator using Program m_mito_tissues_01 (MACS, Miltenyi Biotec). Homogenates were suspended in 4 ml ice-cold buffer (1.0 mM KCl, 1.0 mM Tris-HCl, and 0.1 mM EDTA, pH 8.0) and mixed with 0.67 ml of 2 M sucrose. The suspension was centrifuged at 1300 x g for 5 min to remove nuclei, un-broken cells and large membrane fragments. The supernatant containing mitochondria were pelleted after centrifugation at 9600 x g for 10 min at 4°C. The mitochondrial pellet was resuspended in the Storage Buffer (MACS, Miltenyi Biotec). Total mitochondrial protein was determined using Bradford assay reagent (Bio-Rad). ATP production rates were determined with isolated mitochondria using XF96 Extracellular Flux Analyzers. The mitochondria were diluted in cold 1X MAS (Mitochondrial Assay Solution) and substrate (pyruvate/malate) (Seahorse Biosciences). The mitochondrial suspension (20 μg mitochondrial proteins in 25 μl) was aliquoted into each well while the plate was on ice. The plate was then centrifuged using a swinging bucket microplate adaptor at 2000 × g for 20 min at 4°C. To start the assay, 155 μl of pre-warmed (37°C) 1X MAS and substrate were added to each well containing isolated mitochondria and incubated at 37°C with no CO2 for 10 min. After calibration of the Analyzer, the plate containing mitochondria was sequentially injected with A: port A, 20 μl of 40 mM ADP (4 mM, final); port B, 22 μl of 25 μg/ml oligomycin (2.5 μg/ml, final); port C, 24 μl of 40 μM FCCP (4 μM, final); and port D, 26 μl of 40 μM antimycin A (4 μM, final). OCR was measured and the data were analysed using the XFe Wave software as described above. ATP production rate in brain mitochondria was normalized to mg of mitochondrial protein (18).

GCase activity analyses
Cells were homogenized in 1% sodium taurocholate/1% Triton X-100. GCase activity was determined fluorometrically using 4MU-Glucose as the substrate in 0.25% sodium taurocholate,
Glycosphingolipid analyses

Glycosphingolipids in mouse brains and N2a cells were extracted with chloroform and methanol as described (83). GC and GS content in the extracts was analysed by ESI-MS/MS using a Waters Quattro Micro API triple quadrupole mass spectrometer (Milford, MA) interfaced with Acquity UPLC system as described (35). The concentration of GC and GS in the brain was normalized to mg tissue weight and in the cells was normalized by mg protein in the cell lysate.

Statistical analyses

The data are presented as mean ± SEM and were analysed by Student’s t-test or one-way ANOVA with post-hoc Tukey test using GraphPad Prism 6. The level of significance was set at P < 0.05. Survival analysis was performed using Kaplan-Meier and the Mantel-Cox tests.

Supplementary Material

Supplementary Material is available at HMG online.

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References


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