Unexpected phenotypic and molecular changes of combined glucocerebrosidase and acid sphingomyelinase deficiency

Marcus Keatinge¹,²,³, *, Matthew E. Gegg⁴, Lisa Watson¹,², Heather Mortiboys², Nan Li¹,², Mark Dunning², Deepak Ailani¹,², Hai Bui⁵, Astrid van Rens⁶, Dirk J. Lefeber⁶,⁷, Anthony H.V. Schapira⁴, Ryan B. MacDonald¹,⁷,⁴,⁹, Oliver Bandmann¹,²,*,‡

¹Bateson Centre, Firth Court, University of Sheffield, Western Bank Sheffield S10 2TN, UK
²Sheffield Institute for Translational Neuroscience (SITraN), University of Sheffield, 385a Glossop Road, Sheffield S10 2HQ, UK
³Centre for Discovery Brain Sciences, Chancellor’s Building, Edinburgh EH16 4TJ, UK
⁴Department of Clinical and Movement Neurosciences, UCL Queen Square Institute of Neurology, London NW3 2PF, UK
⁵Eli Lilly and Company, Drop Code 1940, Indianapolis, IN 46285 USA
⁶Department of Laboratory Medicine, Translational Metabolic Laboratory, Radboud University Medical Center, Nijmegen, The Netherlands
⁷Department of Neurology, Donders Institute for Brain, Cognition and Behaviour, Radboud University Medical Centre, Nijmegen, The Netherlands

*Authors for correspondence: Mkeatinge2@exseed.ed.ac.uk; ryan.macdonald@ucl.ac.uk; o.bandmann@sheffield.ac.uk.
‡Joint senior authors of this work.
§Present address: Institute of Ophthalmology, University College London, 11-43 Bath St, Greater London EC1V 9EL.

Summary Statement
Simultaneous inhibition of gba1 and smpd1 gene function (both implicated in Parkinson’s Disease) unexpectedly rescues neuronal gene expression, mitochondrial defects and extends the lifespan of gba1 mutant zebrafish.

Abstract
Heterozygous variants in GBA1 encoding glucocerebrosidase (GCase) are the most common genetic risk factor for Parkinson’s disease (PD). Moreover, sporadic PD patients also have a substantial reduction of GCase activity. Genetic variants in SMPD1 are also overrepresented in PD cohorts, whilst a reduction of its encoded
enzyme (ASM) activity is linked to an earlier age of PD onset. Despite both converging on the ceramide pathway, how combined deficiencies of both enzymes may interact to modulate PD has yet to be explored. Therefore, we created a double knock out (DKO) zebrafish line for both \textit{gba1} and \textit{smpd1} to test for an interaction \textit{in vivo}, hypothesising an exacerbation of phenotypes in the DKO compared to single mutants. Unexpectedly, DKO s maintained conventional swimming behaviour and had normalised neuronal gene expression signatures when compared to single mutants. We further identified rescue of mitochondrial Complexes I and IV in DKO s. Despite having an unexpected rescue effect, our results confirm ASM as a modifier of GBA1 deficiency \textit{in vivo}. Our study highlights the need for validating how genetic variants and enzymatic deficiencies may interact \textit{in vivo}.

**Keywords:** Parkinson’s disease, glucocerebrosidase 1, acid sphingomyelinase, zebrafish, gene-gene interaction.

**INTRODUCTION**

There is compelling evidence of an excessive burden of lysosomal disease gene variants and lysosomal dysfunction in PD.\textsuperscript{1,2} Bi-allelic mutations in \textit{glucocerebrosidase 1} (\textit{GBA1}) cause the lysosomal storage disorder (LSD) Gaucher’s disease (GD), whilst heterozygous mutations are the most common and strongest genetic risk factor for sporadic PD, with a prevalence of ~ 5-20\% depending on the population investigated.\textsuperscript{3–5} PD patients also exhibit reduced GCase activity in different tissues, including the brain\textsuperscript{6}, regardless of their \textit{GBA1} mutation status.\textsuperscript{6–8} In a similar fashion to \textit{GBA1} mutations, homozygous \textit{SMPD1} mutations (encoding acid sphingomyelinase, ASM) also cause an LSD, in this case Niemann Pick disease while heterozygous \textit{SMPD1} variants are associated with increased risk of sporadic PD.\textsuperscript{9–13} Due to their rarity within the PD population, the functional significance of these \textit{SMPD1} variants is still not completely understood.\textsuperscript{9,14} However, a reduction of ASM activity is correlated with an earlier age of disease onset in PD, as well as in other synucleinopathies including dementia with Lewy bodies and multiple system atrophy.\textsuperscript{9,14} Both \textit{GBA1} and \textit{SMPD1} encode lysosomal enzymes that converge on ceramide metabolism (Figure 1). Therefore, an additive interaction between these two enzymes is biologically plausible and but awaits experimental confirmation. We hypothesised that ASM deficiency could worsen the functional consequences of GCase deficiency, aggravating phenotypes which could potentially lead to, or enhance, neurodegeneration.
Zebrafish (Danio rerio) are an attractive vertebrate model to study the biological effect of both monogenic PD genes and genetic risk factors for PD\textsuperscript{15-17}, including gba1 deficiency.\textsuperscript{18} We had previously characterised a gba1 mutant zebrafish line (gba1\textsuperscript{-/-}),\textsuperscript{19} and demonstrated its usefulness to study gene-gene interactions.\textsuperscript{20} gba1\textsuperscript{-/-} zebrafish faithfully model key features of GCase deficiency/Gaucher’s disease, including Gaucher cell accumulation, marked inflammation with microglial infiltration, mitochondrial dysfunction and neurodegeneration.\textsuperscript{19} gba1\textsuperscript{-/-} larvae develop normally but gba1\textsuperscript{-/-} juvenile zebrafish then rapidly deteriorate from 10-12 weeks onwards and die between 12 and 14 weeks.

As expected, combined GCase and ASM deficiency acted synergistically on key sphingolipid metabolites in the gba1\textsuperscript{-/-};smpd1\textsuperscript{-/-} double-mutant zebrafish. However, instead of a worsening of phenotypes, we unexpectedly observed markedly prolonged survival and conventional swimming behaviour in gba1\textsuperscript{-/-};smpd1\textsuperscript{-/-} mutants compared to the gba1\textsuperscript{-/-} (single) mutant zebrafish. RNAseq-based pathway analysis confirmed restoration of neuronal health in gba1\textsuperscript{-/-};smpd1\textsuperscript{-/-} compared to the gba1\textsuperscript{-/-}. Mechanistic experiments identified a rescue effect of combined GCase and ASM deficiency on mitochondrial Complexes I and IV function in gba1\textsuperscript{-/-};smpd1\textsuperscript{-/-} compared to the marked but distinct mitochondrial dysfunction in gba1\textsuperscript{-/-} or smpd1\textsuperscript{-/-} single mutant zebrafish. The mitochondrial rescue led to an abrogation of oxidative membrane damage, further reflecting the overall restorative effect of ASM deficiency on neuronal health in GCase deficiency. Our study highlights the need of functional, mechanistic validation for the interaction of any putative genetic/enzymatic risk factors for human diseases in suitable model systems, rather than readily assuming an additive effect.

Methods

Zebrafish husbandry

All larval and adult zebrafish were housed at the University of Sheffield; experimental procedures being in accordance UK Home Office Animals (Scientific Procedures) Act 1986 (Project license PPL 70/8437, held by Dr Oliver Bandmann). Adult zebrafish were housed at a density of 20 per tank, on a cycle of 14 h of light, 10 h of dark. Adults and embryos were kept at constant temperature of 28°C.

Mutant line generation and line maintenance

The gba1\textsuperscript{-/-} mutant lines was generated using TALEN technology.\textsuperscript{19} The smpd1\textsuperscript{-/-} mutant line was generated by the Crispr/Cas9 method as previously described.\textsuperscript{21,22} The following ultramer template was used: 5'
The `smpd1` line was genotyped using primers F 5' - AGCCGTGGTGGTTTCTACAG and R 5' - CCTTCTCTCCCTGTTCTCG. The `smpd1` line was crossed to `gba1` to generate double heterozygous individuals. These were subsequently in-crossed to generate double mutants, single mutants and WT controls. At each in-cross larvae were genotyped at 3 days post-fertilisation (dpf) by larval tail biopsy as previously described. Each genotype was raised in genotype-specific tanks at a density of 10-15 fish per tank. All individuals were re-genotyped at 10 weeks post-fertilisation. For survival curves, animals were culled for humane reasons when they could no longer maintain consistent buoyancy.

**Immunohistochemistry**

WT and mutant fish were fixed in 4% paraformaldehyde overnight at 4°C before removal of the eye, which was incubated in 30% sucrose in PBS overnight at 4°C. Eyes were then embedded in OCT and cryosectioned at 20um (Leica). Slides were rehydrated in PBS before blocked for 1 hour at room temperature using 150 µl block (1% sheep serum, 5% BSA, 0.3% Triton-X, 0.1% Tween-20 in PBS). Slides were then incubated with 150 µl primary antibody solution (4.C.4; 1:50 dilution in block solution; Antibody registry ID: AB_10013752) overnight at 4°C. After incubation slides were washed 3 times in PBS for 20 minutes followed by incubation with secondary antibody (Alexa anti-mouse-647; A-21235) for 2 hours. Slides were then washed in PBS 3x20min, before adding Fluoroshield with DAPI (Sigma; F6057-20ML) and applying a glass coverslip. Slides were imaged on a Zeiss LSM 900 confocal microscope using a 40x water immersion objective.

**Biochemical activity assays and mass spectrometry**

ASM activity was determined using homogenates prepared as follows. Tubes containing twenty embryos (5 dpf) were sonicated in 500 µl MilliQ and centrifuged (11000 rpm). 20 µL of supernatant was incubated with substrate HMU-PC (6-hexadecanoylamino-4-methylumbelliferyl-phosphorylcholine 0.66 mM, 20 µL, Moscerdam Substrates, NL) at pH 5.2 and 37 °C for 2 hours. Fluorescence intensity was measured at 415 (ex) and 460 (em) nm using a plate reader (LS55, Perkin Elmer). Lysosomal and mitochondrial enzyme assays as well as mass spectrometry were undertaken as previously described.19
**Lipid peroxidation assay**

We were unable to isolate enough mitochondria from brain tissue to robustly measure lipid peroxidation signals above background levels. However, enough mitochondria could be isolated from 3-month-old adult zebrafish bodies to perform the assay robustly. Bodies were homogenised in ice cold mitochondrial isolation buffer. The Abcam lipid peroxidation kit (ab118970) fluorometric assay was used to measure lipid peroxidation according to manufacturer’s instructions. Results were normalised to WT.

**RNA preparation for gene expression analysis**

RNA was prepared from brain tissue of 12 weeks old zebrafish. A Trizol based protocol was used to extract RNA from the tissue. Briefly, individual brains were homogenized in 250 µl TRI Reagent (Sigma) and incubated at room temperature before adding 50 µl chloroform (Thermo Fisher). The samples were centrifuged at 13,300 x g and the top aqueous phase was collected and transferred to a separate tube. RNA was precipitated from the aqueous phase by mixing with equal volume of isopropanol (Thermo Fisher) and centrifugation at 13,300 x g. Precipitated RNA was resuspended in DEPC-treated water (Thermo Fisher) and its concentration and quality were quantified using the Nanodrop 1000 Spectrophotometer. 750 ng of high quality total RNA, with an RNA integrity number (RIN) of 9 or above, was used in the preparation of sequencing libraries using the NEB Ultra II Directional RNA Library Prep Kit (NEB catalogue number E7760), following the polyA mRNA workflow. Libraries were individually indexed and pooled for sequencing. Single-end 100bp sequencing was performed on the Illumina HiSeq 2500 platform using Rapid Run mode with V2 chemistry.

**RNA-seq Analysis**

Raw sequencing reads were processed using the bcbio workflow system. The quality of the samples was checked using FastQC and multiqc. The salmon tool (v0.9.01) was used to quantify genes from the zebrafish reference transcriptome (Danio rerio.GRCz11.98.gtf from Ensembl.org). Unsupervised clustering and PCA with DESeq2 revealed a batch effect corresponding to sample preparation date. Differential expression was performed using DESeq2 incorporating a batch factor into the model. The contrast tested was between gba1<sup>−/−</sup>, double mutants and WT; a positive log fold-change indicating higher expression in GBA single mutants. The clusterProfiler Bioconductor package was used to identify enriched pathways in
up- (adjusted p-value less than 0.05 and log fold-change > 1) and down-regulated genes (adjusted p-value less than 0.05 and log2 fold-change <-1).27

Gene set enrichment analysis
The analysis was performed with Gene Set Enrichment Analysis (GSEA) software version 4.0.3 (https://www.gsea-msigdb.org/gsea/index.jsp). GSEA preranked analysis was used with default settings except for “Collapse/Remap to gene symbols” set to “No_Collapse”. A ranked list used for the analysis was calculated with each gene assigned a score based on the adjusted p value and the log2 fold change. Zebrafish lysosomal and mitochondrial gene sets were prepared by identifying zebrafish homologues of the genes in human genes sets in Molecular Signatures Database (MSigDB) v7.1.

Statistical analysis
Graphpad prism V.6 software (Graphpad) was used for statistical analysis and all error bars shown denote the mean ± SD of the mean. All experiments were performed in biological triplicate unless otherwise stated. All data were analysed with either T test, two-way ANOVA. Significance in all enzyme activity assays was determined by two way ANOVA with Tukey’s multiple comparison test.

Results
smpd1⁻/⁻ zebrafish display abolished acid sphingomyelinase activity and marked sphingolipid accumulation
We initially hypothesised that deficiency of both GCase and ASM enzymes may synergise, leading to a further aggravation of gba1 deficient phenotypes in vivo. To address this, we identified a single smpd1 orthologue in zebrafish (ENSDARG00000076121) with 59% shared identity to the human SMPD1 gene at both the DNA and the protein level. CRISPR/Cas9 technology was used to generate a smpd1 stable mutant line (smpd1⁻/⁻). The selected mutant allele contained a 5bp deletion and 136bp insertion within exon 3, resulting in a frame-shift and the generation of a premature stop codon (Figure 2A and Supplemental Figure S1). Enzymatic activity of ASM in smpd1⁻/⁻ at 5 dpf was reduced by 93% (p=0.006, Figure 2B). The large reduction in ASM enzymatic activity resulted in a significant increase of key glycolipid substrates in the smpd1⁻/⁻ larvae already at 5 dpf (Figure 2C). The smpd1⁻/⁻ line was crossed with gba1⁻/⁻ to generate gba1⁻/⁻; smpd1⁻/⁻. The latter were subsequently in-crossed to generate double mutants, single mutants and WT controls for all subsequent experiments. At each in-cross larvae were genotyped at 3
days post-fertilisation (dpf) and then raised in genotype-specific tanks. Every genotype was present in its expected mendelian ratio (1/16th) during genotyping at larval stages (Supplemental table 1), but only WT, gba1<sup>−/−</sup>, smpd1<sup>−/−</sup> and gba1<sup>−/−</sup>; smpd1<sup>−/−</sup> were raised for experiments.

**Combined ASM and GCase deficiency synergistically increases sphingolipid metabolites**

We had previously reported marked sphingolipid accumulation in gba1<sup>−/−</sup> zebrafish. We hypothesised that combined (enzymatic) GCase and ASM deficiency would synergistically increase distinct sphingolipid subtypes. Using mass spectrometry, a comprehensive panel of glycolipid substrates was analysed in the brains of gba1<sup>−/−</sup> and smpd1<sup>−/−</sup> single mutant as well as in gba1<sup>−/−</sup>;smpd1<sup>−/−</sup> double mutant zebrafish and WT controls at 12 weeks of age. As expected, a marked additive effect of combined GCase and ASM deficiency was observed for Glucosylceramide levels (the direct substrate of GCase) (Figure 3A). This was likely due to both a block in Glucosylceramide catabolism and also due to metabolic compensation in the flux of sphingolipid generation. Combined GCase and ASM deficiency also resulted in an additive effect on lactosylceramide, ceramide and sphinganine levels (Figure 3B-D). Sphingosine levels were increased in gba1<sup>−/−</sup>;smpd1<sup>−/−</sup> compared to WT, reflecting an increase compared to gba1<sup>−/−</sup> but not compared to smpd1<sup>−/−</sup> (Figure 3E). Unexpectedly, there was no synergistic effect in sphingomyelin levels in the gba1<sup>−/−</sup>;smpd1<sup>−/−</sup> double mutants (Figure 3F).

The inflammation markers chitotriosidase and β-Hexosaminidase are markedly increased in the serum of GD patients and used as biomarkers to monitor disease activity. We previously observed a marked increase in chitotriosidase and β-hexosaminidase activity in gba1<sup>−/−</sup> zebrafish brain tissue at 12 weeks. As key GCase substrates were synergistically increased in gba1<sup>−/−</sup>;smpd1<sup>−/−</sup> double mutant zebrafish, we investigated whether combined GCase and ASM inactivation may also result in a further increase of chitotriosidase and β-hexosaminidase activity. Unexpectedly, gba1<sup>−/−</sup>;smpd1<sup>−/−</sup> double mutant zebrafish displayed a similar increase in chitotriosidase and β-hexosaminidase activity compared to gba1<sup>−/−</sup> (Supplemental Figure 2). Furthermore, we had previously detected Gaucher cell invasion in the CNS of end stage gba1<sup>−/−</sup>. Analyses of the retinas across all genotypes demonstrated similar Gaucher cell invasion in the double mutants (Figure 4) with both gba1<sup>−/−</sup> and gba1<sup>−/−</sup>;smpd1<sup>−/−</sup> double mutants showing a ~50% increase in 4C4 positive cells compared to WT. These 4C4 cells in both gba1<sup>−/−</sup> and gba1<sup>−/−</sup>;smpd1<sup>−/−</sup> double mutants
also tended to be larger and rounder, indicative of the Gaucher cells we had previously described. These data suggest persistent yet unaltered neuroinflammatory states in the double mutants despite a marked synergistic increase in sphingolipid metabolites.

**ASM deficiency unexpectedly prolongs survival in GCase deficiency**

The marked additive effect of combined GCase and ASM deficiency on sphingolipid levels led us to hypothesize that ASM deficiency would further worsen the motor phenotype and shorten survival in \( gba1^{-/-};smpd1^{-/-} \) double mutant zebrafish. Unexpectedly, genetic inactivation of ASM led to a complete rescue of this behaviour in the \( gba1^{-/-};smpd1^{-/-} \) double mutant zebrafish (Supplemental Movies S1 (WT), S2 \( (smpd1^{-/-}) \), S3 \( (gba1^{-/-}) \) and S4 \( (gba1^{-/-};smpd1^{-/-}) \)). Importantly, disease free survival, where animals could consistently maintain buoyancy, was also markedly increased by 22% in \( gba1^{-/-};smpd1^{-/-} \) double mutant zebrafish compared to \( gba1^{-/-} \) (median survival of 102 dpf in \( gba1^{-/-} \) and 125dpf in \( gba1^{-/-};smpd1^{-/-} \), \( p=0.0055 \), Figure 5A). Despite not exhibiting the same barrel rolling phenotype as the \( gba1^{-/-} \), and could also maintain their buoyancy, the \( gba1^{-/-};smpd1^{-/-} \) double mutants would ultimately be found unresponsive at the bottom of the tank and so were culled for humane reasons. We also raised \( smpd1^{-/-} \) to determine its lifespan, but never encountered a decrease in viability compared to WT, even up to the age of 18 months (data not shown).

**RNAseq base pathway analysis confirms restored neuronal health in \( gba1^{-/-};smpd1^{-/-} \)**

We next applied RNAseq-based pathway analysis to further elucidate the underlying mechanisms of the observed rescue effect. The differential gene expression analysis all four genotypes (wild type, \( gba1^{-/-} \) and \( smpd1^{-/-} \) single mutants, \( gba1^{-/-};smpd1^{-/-} \) double mutants) identified a total of 512 genes which were dysregulated in \( gba1^{-/-} \) but rescued in \( gba1^{-/-};smpd1^{-/-} \). Amongst these, there are a downregulation of 219 genes and an upregulation of 293 genes in \( gba1^{-/-} \) compared to wild-type and \( gba1^{-/-};smpd1^{-/-} \) (adjusted \( p \)-value \( \leq 0.05 \), \( \log 2 \text{Fold change} \geq 1 \)). We next employed ClusterProfiler analysis on gene ontology (GO) categories to identify functionally relevant pathways within the rescued gene sets. Key neuronal pathways including the GO terms for synaptic signalling, chemical synaptic transmission and calcium ion regulated exocytosis were markedly downregulated in \( gba1^{-/-} \) but normalised in \( gba1^{-/-};smpd1^{-/-} \) (Figure 5B). This suggests that key aspects of neuronal function were restored in the \( gba1^{-/-};smpd1^{-/-} \) double mutants.
We also observed an enrichment of upregulated genes in \textit{gba1^{−/−}} compared to \textit{gba1^{−/−};smpd1^{−/−}} in a broad range of GO terms, the top 5 of which thought to regulate muscle function. However, since our RNA-seq analysis was carried out on brain tissue, we consider these changes to be of limited relevance only (Supplementary Figure 3). Upregulation of the inflammatory signature in \textit{gba1^{−/−}} was retained in the \textit{gba1^{−/−};smpd1^{−/−}} but not further enhanced (data not shown).

As both GCase and ASM are lysosomal hydrolases, we specifically focused on the effect of isolated GCase deficiency in \textit{gba1^{−/−}} compared to combined GCase and ASM deficiency in \textit{gba1^{−/−};smpd1^{−/−}} on lysosome transcriptomic pathways. Gene set enrichment analysis led to the identification of 27 leading-edge, dysregulated lysosomal genes, which account for the pathway’s enrichment signal. The expression of these 27 lysosomal genes was increased in \textit{gba1^{−/−}} compared to wild-type and \textit{gba1^{−/−};smpd1^{−/−}} (Figure 5C, Supplementary Table 2). Amongst these 27 genes, acid hydrolases contributed the most. Cathepsin L, involved in the initiation of protein degradation, ranked as the top rescued gene. The apparent normalisation of lysosomal gene expression profiles in \textit{gba1^{−/−};smpd1^{−/−}} was in contrast to the observed marked increase in a wide range of sphingolipid levels in \textit{gba1^{−/−};smpd1^{−/−}} compared to \textit{gba1^{−/−}} or \textit{smpd1^{−/−}} single mutants (see above).

We had previously observed marked mitochondrial dysfunction in \textit{gba1^{−/−}}. We therefore also focussed on the analysis of mitochondrial genes involved in the oxidative phosphorylation pathway. This leading-edge mitochondrial gene subset included 16 genes encoding the subunits of the Complex I, II, IV and V in mitochondrial electron transport chain (Supplementary Table 3). Interestingly, gene set enrichment analysis showed an upregulation of this mitochondrial gene subset in \textit{gba1^{−/−}}, presumably as a compensatory mechanism to the impaired function of the mitochondrial respiratory chain, but similar mitochondrial gene expression levels in wild-type and \textit{gba1^{−/−};smpd1^{−/−}} (Figure 5D).

\textbf{Restoration of mitochondrial Complex I and IV function in gba1^{−/−};smpd1^{−/−}}

We next compared the mitochondrial respiratory chain function across all four genotypes to further determine whether the normalised gene expression levels for oxidative phosphorylation-related genes would be reflected in normalised mitochondrial function. Complex I activity was reduced by 65% in \textit{smpd1^{−/−}} compared to WT levels (p=0.0198, Figure 5A) but restored to 92% of WT levels in \textit{gba1^{−/−};smpd1^{−/−}} (p=0.0445, Figure 6A). Complex II was not significantly altered in any of the
genotypes (Figure 6B). Complex III activity in gba1⁻/⁻ was reduced by 45% compared to WT levels (p=0.0091, Figure 6C) as previously observed. Complex III activity in the gba1⁻/⁻;smpd1⁻/⁻ double mutant zebrafish was reduced by only 9% compared to WT levels however this did not reach significance when compared to gba1⁻/⁻ (p=0.1688). Complex IV activity was unchanged in smpd1⁻/⁻ compared to WT, but reduced by 40% in gba1⁻/⁻ compared to WT as previously reported (p=0.0491, Figure 6D). Remarkably, there was a marked improvement of complex IV activity in gba1⁻/⁻ ;smpd1⁻/⁻ with an increase in activity of 69% compared to gba1⁻/⁻ (p= 0.0005, Figure 6D). Thus, there is rescue of mitochondrial respiratory chain function in mitochondrial Complex I and IV, where ASM deficiency normalises mitochondrial Complex IV function in gba1⁻/⁻ and GCase deficiency normalises mitochondrial Complex I function in smpd1⁻/⁻. Malfunction of the mitochondrial respiratory chain can result in oxidative stress and subsequent lipid peroxidation. We therefore investigated next whether the observed rescue in mitochondrial Complex I and IV results in reduced oxidative stress-related damage. Mitochondrial lipid peroxidation was increased in whole gba1⁻/⁻ adult fish by 63% above WT levels (p= 0.0214, Figure 6E). As predicted, lipid peroxidation levels were reduced by 70% in gba1⁻/⁻;smpd1⁻/⁻ double mutants compared to gba1⁻/⁻ and thus effectively normalised (p=0.0094, Figure 6E).

Discussion

Biochemically, GCase and acid sphingomyelinase both play a key role in sphingolipid metabolism. Unexpectedly, we observed a rescue of motor behaviour and a marked prolongation of life expectancy following combined GCase and ASM deficiency, despite clear evidence of an additive effect on the intracellular level of key sphingolipids and their metabolites. The remarkable rescue effect of mitochondrial Complex I and IV in gba1⁻/⁻;smpd1⁻/⁻ on behaviour and survival suggests a central role of mitochondrial dysfunction in GCase deficiency. The profound normalisation of neuronal function in gba1⁻/⁻;smpd1⁻/⁻, as indicated in our RNA-seq based pathway analysis, is in keeping with the observation of the rescued motor phenotype. The normalisation of intracellular homeostasis is also reflected by the normalisation of both lysosomal and mitochondrial transcriptional pathways.

Mitochondrial dysfunction is a key feature of both familial and sporadic PD as well as in LSD’s, the mitochondrial network and lysosomal system are also known to be tightly interlinked. ASM activity must also be tightly controlled, as either too much or too little has been shown to negatively affect mitochondrial function, depending on the cell type, tissue or experimental paradigm understudy. However, when
focussed on neuronal health, ASM inhibition consistently ameliorates phenotypes in the context of neuronal loss.\textsuperscript{39–41} A plausible rescue mechanism of mitochondrial function in DKO, could be glutamate/calcium signalling. Neuronal GCase deficiency \textit{in vitro} has been shown to sensitise mitochondria to physiological levels of glutamate.\textsuperscript{33} This leads to pathological responses in calcium signalling and downstream mitochondrial dysfunction.\textsuperscript{33} Conversely, ASM deficiency \textit{in vitro} has been shown to cause a decreased vulnerability to glutamate linked excitotoxicity in neurons.\textsuperscript{42} This was not only linked to a decrease in intracellular calcium levels, but also of oxidative stress.\textsuperscript{42} Inhibiting ASM function in primary oligodendrocyte culture, can also rescue glutamate induced mitochondrial dysfunction.\textsuperscript{39}

Of note, 9 genes involved in calcium ion regulated exocytosis were downregulated in \textit{gba1}\textsuperscript{-/} single mutants but subsequently normalised in the DKO, namely syt2a, syt7a, cplx3a, cadpsa, snap47 cacna1hb, rims1b, cbarpb, napbb.

An alternative mechanism could be intracellular redistribution of the sphingolipid profile in double mutants. We only detected a synergistic increase in sphingolipid levels in DKO compared to single \textit{gba1}\textsuperscript{-/}, but not a normalisation. However, this does not exclude an effect on subcellular localisation of a specific sphingolipid metabolite which may underpin the observed rescue mechanism. Sphingolipid signalling is vitally important for many varied inter- and intracellular processes.\textsuperscript{29} Sphingolipid signalling must remain highly compartmentalised due to its pleiotropic effects.\textsuperscript{43–45} However, due to technical reasons we used bulk brain tissue for our metabolite analysis, which would therefore not allow for the detection inter- and intracellular sphingolipid differences. Future work should involve metabolic analyses of sphingolipids separated by cell type and by specific cellular fractions, to produce a spatial understanding of the distinct sphingolipid metabolism and distribution across the different \textit{gba1}\textsuperscript{-/}/\textit{smpd1}\textsuperscript{-/} genotypes. Furthermore, whole body analyses using recently developed techniques to monitor in-situ glucosylceramide generation would give novel insights into the glycolipid dysregulation in our double mutants.\textsuperscript{46}

Intriguingly, deficiency of another LSD gene, \textit{asah1}, which functions on a separate arm of the ceramide pathway to \textit{smpd1} (Figure 1), also ameliorates \textit{gba1} deficient phenotypes \textit{in vivo} and \textit{in vitro}. Biallelic \textit{ASAH1} mutations cause the LSD Farber disease in humans. By developing a DKO zebrafish for \textit{gba1}\textsuperscript{-/} and \textit{asah1}\textsuperscript{-/}, Lelieveld and colleagues demonstrated that \textit{asah1} deficiency also led to a rescue of behavioural and neuronal phenotypes in a similar manner to our DKO \textit{gba1}\textsuperscript{-/};\textit{smpd1}\textsuperscript{-/}. 
zebrafish. In keeping with our own data, the rescue effect observed was not due to an amelioration of neuroinflammation, as DKO retained the upregulation of tnfβ, il1β and apoeb exhibited by gba1−/−. Similarly, Kim et al demonstrated that pharmacological inhibition of ASAH1 led to a significant reduction in GBA1 linked cellular phenotypes including accumulation of ubiquitinated proteins and alpha synuclein in dopaminergic neuronal cultures derived from PD-GBA1+− patients.

For practical reasons, we modelled combined enzymatic deficiency using homozygous, and not heterozygous mutants for gba1 and smpd1, intrinsically modelling LSDs and not PD. However, the unexpected nature of our results demonstrates the need for future characterisation of combined partial LSD gene deficiencies in the wider context of PD.

**Abbreviations**
Parkinson’s Disease (PD), Glucocerebrosidase 1 (GBA1), acid sphingomyelinase (SMPD1), Glucocerebrosidase enzyme (GCase), Acid Sphingomyelinase enzyme (ASM), Genome Wide Association Study (GWAS), Lysosomal storage disorder (LSD), RNA Integrity Number (RIN), Gene Set Enrichment (GSEA), Double Knockout (DKO).

**Supplemental data**
Supplemental data can be found online at

**Declaration**
Ethics approval and consent to participate

All larval and adult zebrafish were housed at the University of Sheffield; experimental procedures being in accordance UK Home Office Animals (Scientific Procedures) Act 1986 (Project license PPL 70/8437, held by Dr Oliver Bandmann).

Consent for publication

Not applicable.
Availability of data and materials

All data generated or analysed during this study are included in this published article [and its supplementary information files]. The exception is the raw data of the gene expression which is available from the corresponding author on reasonable request.

Competing interests

The authors declare that they have no competing interests.

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Authors’ contributions

Conducted and analysed experiments: MK, MG, LW, HM, NL, MD, DA, HB, AR
Experimental design: MK, MG, LW, HM, NL, MD, DA, HB, AR, DL, RM, AS, OB
MK and OB wrote the manuscript.
All authors read and approved the final manuscript.

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References:


Fig. 1. A representative image of the ceramide pathway. Enzymes of the Ceramide pathway under study (GBA1 and SMPD1) are highlighted in red, whilst *enzymes are linked to PD.
Fig. 2. Genetic and biochemical characterisation of the *smpd1*^-/- mutant zebrafish line.

(A) Representative genotyping gel of the *smpd1*^-/- alleles (5bp del and 136bp insertion) demonstrating WT, *smpd1*^+/- and *smpd1*^-/-.

(B) Acid sphingomyelinase (ASM) enzymatic activity compared to WT controls, n=6 larvae at 5 dpf per genotype, *p*=0.006 by Welch’s t-test.

(C) Quantification of sphingolipid metabolites, namely Sphingomyelin (Sph), dihydro-sphingomyelin (DHSph), ceramide (Cer), lactosylceramide (LacCer), Sphinganine (Spgn) and Sphinganine 1 phosphate (Sa1P). All metabolites shown are the C18 neuronal species. Data represented are the mean ±SD. ***p<0.001, **p<0.01 and *p<0.05 by two-tailed t-test.
Fig. 3. Accumulation of key glycolipids across *gba1*<sup>−/−</sup> and *smpd1*<sup>−/−</sup> single mutant and *gba1*<sup>−/−</sup>; *smpd1*<sup>−/−</sup> double mutant genotypes

(A) Glucosylceramide, (B) Lactosylceramide, (C) Ceramide (D), Sphinganine, (E) Sphingosine and (F) Sphingomyelin levels in WT, single mutant *gba1*<sup>−/−</sup> vs *smpd1*<sup>−/−</sup> zebrafish and *gba1*<sup>−/−</sup>; *smpd1*<sup>−/−</sup> double mutant zebrafish. n=10 of 12 week old zebrafish brains used per group. Data represented are the mean ±SD. ****p<0.0001, ***p<0.001, **p<0.01 and *p<0.05 by 2 way Anova test with Tukey’s multiple comparison.
Fig. 4. Gaucher cell accumulation is present in the retina of gba1<sup>−/−</sup>; smpd1<sup>−/−</sup> double mutants.

(A) Analysis of 4C4 positive cells within the retina reveals comparatively significant increases of ~50% in both gba1<sup>−/−</sup> (p=0.0026) and gba1<sup>−/−</sup>; smpd1<sup>−/−</sup> double mutants (p=0.0193) compared to WT, despite enhancement of key glycolipids in the double mutants. 4C4 counts in the smpd1<sup>−/−</sup> were not statistically different to WT. Data represented are the mean ±SD. **p<0.01 and *p<0.05 by 2-way Anova test, n= 4 for
all groups. (B) 4C4 cells in the retina of the \( gba1^{+/+} \) and \( gba1^{+/+}; smpd1^{+/+} \) double mutants were localised throughout the retina and appeared larger and rounder (arrows), when compared to ramified WT 4C4 cells (arrowheads).
Fig. 5. Acid sphingomyelinase deficiency improves survival and rescues neuronal dysfunction in $gba^+$ zebrafish.

(A) Disease free survival analysis of $gba1^+$; $smpd1^+$ double-mutant (n=13) compared to $gba1^+$ single mutant zebrafish (n=12). p= 0.0055 by Gehan-Breslow-Wilcoxon test. Animals were culled for human reasons when they could no longer consistently maintain buoyancy (B) The Comparative gene ontology analysis indicates marked global impairment of neuronal function in $gba1^+$ but restoration of neuronal health in $gba1^+$; $smpd1^+$. A differential expression analysis was first used to identify genes with statistically significant difference between $gba1^+$ and $gba1^+$; $smpd1^+$. Genes with adjusted p-value < 0.05 and log2 fold-change > 1 or <-1 were used to identify enriched biological processes amongst genes over- and under-expressed in $gba1^+$. ClusterProfiler identified significant enriched GO (gene ontology) terms within the gene expression changes which were plotted by GeneRatio (ratio of the differentially expressed genes in one particular GO term to the total number of differentially expressed genes). The lists of over- and under-expressed genes were analysed separately. Shown are the ten GO terms with highest gene ratios amongst under-expressed genes; all relating to key aspects of normal neuronal function and homeostasis. These pathways downregulated in $gba1^+$ compared to WT, were normalised in the double mutants. Each GO term is coloured according to the adjusted p-value and ranked according to gene ratio. The size of the point is then
scaled according to the number of differentially-expressed genes in the GO term. (C+D) Lysosomal pathway genes and oxidative phosphorylation pathway genes are upregulated in gba<sup>−/−</sup> but normalised in gba1<sup>−/−</sup>;smpd1<sup>−/−</sup>. The comparison of RNAseq-based transcription levels in the respective pathways between gba<sup>−/−</sup> and gba1<sup>−/−</sup>;smpd1<sup>−/−</sup> revealed that both (C) lysosomal pathway genes and (D) oxidative phosphorylation pathway genes were enriched with marked upregulation of both pathways in gba1<sup>−/−</sup> compared to wild-type and gba1<sup>−/−</sup>;smpd1<sup>−/−</sup>. The x-axis ranks all differentially expressed genes based on the rank metric score from the most upregulated (left) to the most downregulated (right) for either pathway. The vertical black lines show the location of pathway genes in the entire ranked list from the gba1<sup>−/−</sup> expression changes, compared to WT and double mutants. The y-axis is the degree to which a set of pathway genes is overrepresented at the extremes (up or down-regulated) of the entire ranked list of all differentially expressed genes within the genome. A peak in enrichment score (green line) demonstrates an enrichment of pathway genes amongst all over- or under-represented genes. A sharp peak demonstrates how highly upregulated each pathway is within the gba1<sup>−/−</sup> group compared to WT and double mutants.
Fig. 6. Mitochondrial respiratory chain function and lipid peroxidation.

(A) Complex I activity was reduced in smpd1<sup>-/-</sup> by 64±34.77% compared to WT (p=0.0198). Complex I activity was normalised in gba1<sup>-/-</sup>;smpd1<sup>-/-</sup> with an increase by 56±21.9% compared to smpd1<sup>-/-</sup> (p=0.0445). (B) Complex II activity was similar across the different genotypes (p>0.05). (C) Complex III activity was reduced in gba1<sup>-/-</sup> compared to WT by 45%±14.99 (p=0.0091) and increased by 35±35.2% in gba1<sup>-/-</sup>;smpd1<sup>-/-</sup> compared to gba1<sup>-/-</sup> but this did not reach significance (p>0.05). (D) Complex IV activity was reduced in gba1<sup>-/-</sup> by 40%±26.79 compared to WT (p=0.0491), but completely rescued in gba1<sup>-/-</sup>;smpd1<sup>-/-</sup> double mutants with an increase by 69±26.79% compared to gba1<sup>-/-</sup> (p=0.0005). For all mitochondrial complex activity measurements, 6 brains were used for each genotype. (E) Mitochondrial lipid peroxidation levels were increased 63±51% in gba1<sup>-/-</sup> compared to WT (p=0.0214), but reduced by 71±43.48% compared to gba1<sup>-/-</sup> and thus effectively normalised to WT levels in gba1<sup>-/-</sup>;smpd1<sup>-/-</sup> double mutants (p=0.0094). For lipid peroxidation experiments, n= 6-8 zebrafish bodies were used for each genotype. Significance in both mitochondrial respiratory chain assays and lipid peroxidation levels was determined by two way ANOVA with Tukey’s multiple comparison test using 12 week brain material. Data represented are the mean ±SD. ***p<0.001, **p<0.01 and *p<0.05.
Fig. S1. CRISPR/Cas9 generated smpd1 allele.
Using CRISPR/Cas9, we isolated a mutant allele in exon 3 containing a 5bp deletion (red script) with a 136bp insertion (blue script) leading to a frame shift and generation of a premature stop codon at codon 426 (full length WT smpd1: 676 amino acids).
Fig. S2. GD biomarker activation is not substantially altered in gba1<sup>−/−</sup>;smpd1<sup>−/−</sup>. (A) Hexosaminidase activity was not significantly changed between smpd1<sup>−/−</sup> and WT, but markedly elevated by 85% in gba1<sup>−/−</sup> (2284±200 nmol/hr/mg protein) compared to WT (1234±47.3nmol/hr/mg protein; p<0.0001). gba1<sup>−/−</sup>;smpd1<sup>−/−</sup> displayed a similar increase in activity of 70% above WT levels (2131±272.2nmol/hr/mg protein, p<0.0001) compared to gba1<sup>−/−</sup>. (B) Chitotriosidase activity was not significantly changed in smpd1<sup>−/−</sup> compared to WT, but increased in gba1<sup>−/−</sup> by 389% (31.8±11.95 nm/hr/mg protein, p=0.0002) compared to WT (6.5±4.0 nm/hr/mg protein) and by 230% in gba1<sup>−/−</sup>;smpd1<sup>−/−</sup> showed (21.5±5.1nm/hr/mg protein, p=0.0177) compared to WT. (C) Beta galactosidase activity in all genotypes showed comparable activities of approximately 50nm/hr/mg protein. Significance in all enzyme activities assays was determined by two way ANOVA with Tukey’s multiple comparison test with an n of 5 per group using 12 week brain material. Data represented are the mean ±SD. ****p<0.0001, ***p<0.001, **p<0.01 and *p<0.05.
**Fig. S3. Over-represented GO terms in the up-regulated genes.** The top 10 GO terms as identified by ClusterProfiler from the list of statistically-significant genes (adjusted $p$-value < 0.05 and log2 fold-change $> 1$ or $<-1$ respectively. Each point is coloured according to the adjusted $p$-value for the GO term being over-represented, and the size of the point is scaled according to the number of differentially-expressed genes in the GO term. GO terms are ordered on the y-axis according to the size of the GO-term and adjusted $p$-value in the case of GO terms with the same number of genes.
### Table S1. Observed and expected genotypes from \( gba1^{+/-};smpd1^{+/-} \) incrosses from 4 pooled clutches.

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<thead>
<tr>
<th>Genotype</th>
<th>WT</th>
<th>( gba1^{+/-} )</th>
<th>( gba1^{+/-} )</th>
<th>WT; ( smpd1^{+/-} )</th>
<th>WT; ( smpd1^{+/-} )</th>
<th>WT; ( smpd1^{+/-} )</th>
<th>WT; ( smpd1^{+/-} )</th>
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<td>( n )</td>
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<td>48</td>
<td>15</td>
<td>26</td>
<td>48</td>
<td>27</td>
<td>35</td>
<td>15</td>
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<td>Observed %</td>
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<td>19.12%</td>
<td>5.98%</td>
<td>10.36%</td>
<td>19.12%</td>
<td>10.76%</td>
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<tr>
<td>Expected %</td>
<td>6.26%</td>
<td>12.50%</td>
<td>6.26%</td>
<td>12.50%</td>
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### Table S2. The leading-edge gene subset in lysosome pathway.

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<th>Rank Metric Score</th>
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<td>cathepsin L.1</td>
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<td>cd63</td>
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<td>atp6ap1b</td>
<td>ATPase H+ transporting accessory protein 1b</td>
<td>5.114915371</td>
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<td>lipf</td>
<td>lipase, gastric</td>
<td>3.06033802</td>
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<td>fuca1.2</td>
<td>alpha-L-fucosidase 1, tandem duplicate 2</td>
<td>2.934919357</td>
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<td>ctsba</td>
<td>cathepsin Ba</td>
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<tr>
<td>sftpbb</td>
<td>surfactant protein Bb</td>
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<td>hexosaminidase A (alpha polypeptide)</td>
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<td>zgc:110239</td>
<td>zgc:110239</td>
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### Table S3. The leading-edge gene subset in oxidative phosphorylation pathway.

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<th>Symbol</th>
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<th>Rank Metric Score</th>
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<td>atp5pb</td>
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<td>ATPase H+ transporting V0 subunit a1a</td>
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<td>cox4i1</td>
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All videos were taken when the respective zebrafish were 12 weeks of age.

**Movie 1.** Swimming behaviour in WT. All adults placed into a novel tank swim to the bottom momentarily before resuming standard swimming, balance and buoyancy maintained.
Movie 3. Markedly abnormal swimming behaviour in gba"/- with typical “corkscrew” swimming pattern. All adults placed into a novel tank swim in circular motions with balance and buoyancy defects. These increase with frequency and duration at end stage until they need to be culled for humane reasons.

Movie 2. Swimming behaviour in smpd"/. All adults placed into a novel tank swim to the bottom momentarily before resuming standard swimming, balance and buoyancy maintained.
**Movie 4.** Swimming behaviour in *gba*<sup>−/−</sup>;<*smpd*<sup>−/−</sup>. All adults placed into a novel tank swim to the bottom momentarily before resuming standard swimming, balance and buoyancy maintained.