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## Altered lipid homeostasis is associated with cerebellar neurodegeneration in SNX14 deficiency

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28 The authors have declared that no conflict of interest exists.

29

#### 30 Abstract

Dysregulated lipid homeostasis is emerging as a potential cause of neurodegenerative disorders. However, 31 32 evidence of errors in lipid homeostasis as a pathogenic mechanism of neurodegeneration remains limited. Here, we show that cerebellar neurodegeneration caused by Sorting Nexin 14 (SNX14) deficiency is 33 associated with lipid homeostasis defects. Recent studies indicate that SNX14 is an inter-organelle lipid 34 transfer protein that regulates lipid transport, lipid droplet (LD) biogenesis, and fatty acid desaturation, 35 suggesting that human SNX14 deficiency belongs to an expanding class of cerebellar neurodegenerative 36 disorders caused by altered cellular lipid homeostasis. To test this hypothesis, we generated a mouse model 37 that recapitulates human SNX14 deficiency at a genetic and phenotypic level. We demonstrate that 38 cerebellar Purkinje cells (PCs) are selectively vulnerable to SNX14 deficiency while forebrain regions 39 40 preserve their neuronal content. Ultrastructure and lipidomic studies reveal widespread lipid storage and metabolism defects in SNX14 deficient mice. However, pre-degenerating SNX14 deficient cerebella show 41 42 a unique accumulation of acylcarnitines and depletion of triglycerides. Furthermore, defects in LD content 43 and telolysosome enlargement in pre-degenerating PCs, suggest lipotoxicity as a pathogenic mechanism of SNX14 deficiency. Our work shows a selective cerebellar vulnerability to altered lipid homeostasis and 44 45 provides a mouse model for future therapeutic studies.

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#### 47 MAIN TEXT

#### 48 Introduction

Neurodegenerative disorders are characterized by a progressive loss of specific neuronal types often associated with the accumulation of toxic protein aggregates (1). To better understand disease mechanisms and find therapeutic alternatives, the field has principally focused on the study of protein quality control pathways, including autophagy (2, 3). In contrast, little attention has been paid to lipid homeostasis pathways despite their well-established association with neurodegeneration and relevance for the function and integrity of cellular organelles (4-7).

Genetic disorders affecting regulators of lipid homeostasis often show neurodegeneration, particularly 55 affecting the cerebellum and spinal cord (8, 9). The cerebellum integrates motor function with cognition, 56 emotion, and language, and its dysfunction is documented in a wide spectrum of neurological disorders 57 (10-12). Among cerebellar disorders, childhood onset spinocerebellar ataxias are the most severe. In 58 addition to impaired motor coordination and balance, spinocerebellar ataxia in children is often 59 60 accompanied by additional neurologic and systemic symptoms, including neurodevelopmental delay and intellectual disability (13, 14). Recent efforts that combine patient registry assemblies with advances in 61 sequencing technologies are revealing a new class of childhood cerebellar neurodegenerative disorders 62 caused by disfunction of lipid homeostasis pathways (8, 15). 63

Mutations in *Sorting Nexin 14* (*SNX14*) are the cause of a childhood-onset ataxia known as Spinocerebellar Ataxia Recessive 20 (SCAR20), characterized by progressive cerebellar degeneration and severe intellectual disability (16-18). We previously discovered that SCAR20 is associated with enlarged lysosomes and altered autophagy in neural cells derived from patients (16). These findings were also reproduced in patient skin fibroblasts and SNX14 deficient U2OS cell lines but deemed secondary to defects in cholesterol distribution and neutral lipid metabolism (19). Subsequent studies identified SNX14

as a regulator of cholesterol homeostasis in two independent genome wide perturbation screens (20, 21). 70 Although the mechanisms by which SNX14 regulates cholesterol trafficking is still unknown, recent 71 72 reports demonstrate that SNX14 is recruited to the endoplasmic reticulum (ER)-lipid droplet (LD) contact sites to facilitate the incorporation of fatty acids (FA) into triglycerides (TGs) of growing LDs (22). In 73 this process, SNX14 interacts with SCD1, an ER anchored FA desaturase, to cooperate in FA 74 75 incorporation into LDs (23). Consequently, SNX14 deficient cells show enhanced toxicity to saturated FAs and defective FA-stimulated LD biogenesis (22, 23). Furthermore, recent structural predictions 76 suggest that SNX14 and its SNX-RGS family members may be involved in intracellular lipid transfer 77 (24). However, it is currently unknown if the role of SNX14 in lipid homeostasis regulation is implicated 78 in the pathogenesis of SCAR20. 79

To shed light on the cellular and molecular mechanisms that lead to cerebellar degeneration and 80 intellectual disability in SNX14 deficiency, we generated the first Snx14 full body knock out mouse (Snx14 81 82 KO) that survives to adulthood. Our work shows that *Snx14* KO mice recapitulate cerebellar atrophy, and 83 motor and cognitive defects of SCAR20 patients. Whereas cerebellar atrophy is associated with Purkinje cell (PC) degeneration, forebrain regions responsible for cognitive behavior remain protected from 84 neurodegeneration. Guided by transcriptomic analyses that pointed to lipid dysregulation as a potential 85 cause of selective cerebellar degeneration, we identify tissue specific alterations of lipid profiles in Snx1486 cortices 87 KO mice. Particularly, non-degenerating Snx14 KO cerebral exhibit reduced phosphatidylethanolamine (PE) levels that may be associated with synaptic dysfunction, while 88 accumulation of Acylcarnitines (AcCa-s) is unique to pre-degenerating cerebella and likely associated 89 90 with selective cerebellar neurodegeneration. Finally, we show that SNX14 deficiency reduces LD content 91 and causes lipid storage defects in cerebellar PCs. Together, our work provides evidence for the

involvement of lipid homeostasis defects in selective neurodegeneration and uncovers lipid targets for
 therapeutic interventions.

94 **Results** 

#### 95 SNX14 deficiency causes partial embryonic lethality and developmental delay in mice

SCAR20 patients share clinical features of developmental delay and perinatal onset 96 neurodegeneration of the cerebellum. Previous work suggested that the severity of developmental 97 98 phenotypes is species-specific, with SNX14 deficient mice showing fully penetrant embryonic lethality, while dogs and zebrafish display neurological and metabolic defects reminiscent of SCAR20 patients (25). 99 However, by randomly introducing a frameshift 1 bp deletion in the exon 14 of Snx14 (c.1432delG; 100 101 p.Glu478Argfs\*18), we successfully generated SNX14 deficient mice (Snx14 KO) that are viable and thrive despite a complete loss of SNX14 protein and 90% reduction of the transcript when the mutation is 102 in homozygosity (Figure 1A and Supplemental Figure 1A-D). 103

Although Snx14 KO mice survive to adulthood, we noticed that they were born in lower than the 104 expected Mendelian ratio (observed 9.9% vs expected 25%) (Figure 1B). To test if the reduced birth ratio 105 was due to embryonic lethality, we genotyped embryos produced by heterozygous breeding pairs and 106 uncovered that about half of Snx14 KO embryos die between embryonic day (E)10 and E15. The other 107 108 half were distinguishable by their small size, a feature that persisted throughout neonate and adulthood 109 (Figure 1C-E). Notably, similar to SCAR20 patients (16-18), adult Snx14 KO mice showed dysmorphic 110 facial features characterized by an upturned nose, bulging forehead, and eye defects (Figure 1F-G). These 111 data indicate that SNX14 deficiency in mice causes developmental delay phenotypes reminiscent of 112 SCAR20.

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#### 114 Snx14 KO mice display motor and cognitive behavioral defects

Unlike SCAR20 patients who show severe gait abnormalities typical of cerebellar degeneration, 115 Snx14 KO mice were undistinguishable from their wild type (WT) littermates based on their home cage 116 walking activity. However, Catwalk gait analysis revealed a mild gait disruption characterized by longer 117 paw stand time and faster swing speed of the limbs (Figure 2A-B). Functional gait disruption was seen on 118 the horizontal Metz ladder where mice cross a series of rungs separated by varying distances. Here, Snx14 119 120 KO mice had significantly more foot slips than control mice (Figure 2C). Moreover, Snx14 KO mice underperformed when challenged with complex motor tasks that require coordination and balance. On the 121 accelerating Rotarod, Snx14 KO mice showed difficulty maintaining balance (Figure 2D) similar to other 122 cerebellar ataxia mouse models (26). In addition, the accelerating Rotarod procedure was performed in 123 three consecutive days to assess motor learning. While WT mice improved their performance over trial, 124 Snx14 KO learning rate was low, especially for females (Figure 2E). 125

Given that intellectual disability is also a hallmark of SCAR20, we wondered whether *Snx14* KO mice had broader behavioral deficits. To answer this question, we performed a test for social preference and recall (27). During a choice phase of the procedure, the *Snx14* KO mice showed typical preference for a social cue relative to an inanimate object. However, in the recall phase, *Snx14* KO mice failed to discriminate between a familiar and a stranger mouse (Figure 2G). Thus, *Snx14* KO mice showed similar preference to the social cue, but their lack of preference toward exploration of the novel mouse suggests a social memory deficit likely caused by dysfunction of brain regions, including the cerebellum (28, 29).

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#### 134 Behavioral defects are associated with cerebellar atrophy

Having established that SNX14 deficient mice recapitulate developmental, motor, and behavioral delays of SCAR20, we looked for the underlying neuropathologic causes. Similar to humans, in mice, SNX14 is widely expressed in the developing and adult brain with a slight enrichment in older brains (Supplemental Figure 2A). In line with the expression pattern, the gross brain morphology of *Snx14* KO mice appeared normal during the first month of life but showed defects as mice became older. Specifically, we found that *Snx14* KO mice had smaller cerebella than WT littermates starting at 2.5 months of age while forebrain areas were mostly intact (Figure 3A), suggesting that the cerebellum is particularly vulnerable to SNX14 deficiency.

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#### SNX14 deficiency causes selective PCs degeneration

To further determine vulnerabilities of SNX14 deficiency at a cellular level, we histologically 145 analyzed cerebellar and forebrain tissue. Recent single cell transcriptomic data show that within the 146 cerebellum, Snx14 expression is enriched in Golgi cells and PCs (30) (Supplemental Figure 2B). 147 Accordingly, RNAscope in situ hybridization showed an enrichment of Snx14 in PCs (Supplemental 148 Figure 2C-D). PCs are some of the largest neurons in the nervous system and their loss is a hallmark of 149 cerebellar ataxias (31). Thus, we first analyzed PCs in 1-, 2.5-, and 4-month-old cerebellar sections by 150 immunofluorescence (IF) staining with Calbindin 1 (CALB1) antibody. At 1 month of age, both WT and 151 Snx14 KO cerebellar stainings showed perfectly aligned somas in the PC layer and PC dendrites extended 152 into the molecular layer (ML). However, by 2.5 months of age, patches of missing PCs were evident in 153 154 Snx14 KO cerebella (Figure 3B). Quantification of PC number per mm of PC layer confirmed significantly lower PC density in lobule III of 2.5- and 4-month-old Snx14 KO cerebella compared to WT (Figure 3B 155 156 bottom left graph). The loss of PCs in Snx14 KO cerebella was followed by a reduced thickness of the 157 molecular layer that was first detectable at 4 months of age (Figure 3B bottom right graph). Upon closer examination of CALB1 staining, we identified vacuole-like structures within Snx14 KO PC dendrites and 158 159 soma (Figure 3C, Supplemental Figure 3A). Although, these vacuoles were more abundant and larger in 160 older cerebella, they were sparsely detected in 1-month-old PCs, suggesting that these vacuoles may be a

pathological sign that precedes PC neurodegeneration. Remarkably, IF staining with anti-LAMP1 antibody revealed that enlarged vacuoles overlap with lysosomal structures and *Snx14* KO PCs display larger lysosomes in comparison to WT (Figure 3D).

Given that PC degeneration is often followed by disorganization of Bergmann Glia (BG) processes 164 and gliosis, we also immunostained sagittal cerebellar sections with anti-GFAP and anti-IBA1 antibodies. 165 166 Concurrent with PC loss, anterior lobes of 2.5-month-old Snx14 KO cerebella showed abnormal branching of GFAP<sup>+</sup>BG processes (Figure 3E) and an accumulation of IBA1<sup>+</sup> microglia within the ML (Figure 3F). 167 Moreover, we found that reactive astrocytes progressively accumulate nearby the PC layer from 2.5 to 4.5 168 months of age (Figure 3E). Interestingly, these findings were specific of the anterior lobes of Snx14 KO 169 cerebella while posterior lobes (VIII and IX) did not show signs of neurodegeneration until 11 months of 170 age (Supplemental Figure 3B-D). Notably, we did not detect neuronal loss or signs of gliosis in cortical 171 and hippocampal regions of the forebrain (Figure 3G and Supplemental Figure 4A-C). 172

Taken together, our results indicate that despite the wide expression of SNX14 in the whole brain, the forebrain and posterior cerebellum are protected from neurodegeneration, while anterior PCs selectively neurodegenerate in SNX14 deficient mice after 2 months of age.

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#### 177 Lipid response genes are dysregulated in pre-degenerating *Snx14* KO mice cerebella

To gain insights into the molecular mechanisms of selective cerebellar PC degeneration, we next analyzed the transcriptome of Snx14 KO mice cerebella at pre- and post-degenerating stages (1-monthold and 1-year-old, respectively) and compared them with cerebral cortices, which do not show signs of neurodegeneration. After RNA sequencing, we defined differentially expressed genes (DEG) as those showing absolute log<sub>2</sub>(FC)>0.50 with p-adj<0.05 between Snx14 KO and WT tissue. As expected, Snx14was downregulated in all Snx14 KO samples (Figure 4A-B and Supplemental Figure 1C-D). Few

differences between Snx14 KO and WT cerebral cortex transcriptomes were detected at <2 month of age 184 (7 DEGs including Snx14) and only 37 downregulated and 3 upregulated DEGs at 1 year of age (Figure 185 4A). None of these DEGs suggested changes in specific cell type composition, which is consistent with 186 the lack of neurodegeneration or neuroinflammation in our histological analyses (Supplemental Data 1). 187 We then tested if cortical DEGs were enriched in specific cellular and molecular functional annotations. 188 189 Given the short list of DEGs at 1-month-old cortices, we only performed functional annotation analysis on the 37 downregulated genes at 1 year of age. Results revealed a significant enrichment for genes 190 involved in synaptic vesicle membrane (i.e. Doc2b, Sv2c) (Figure 4A, C). Accordingly, SNX14 has been 191 192 shown to promote synaptic transmission in mouse cortical neuronal cultures (32). To further validate these data, we analyzed cortical sections by IF staining of pre-and post-synaptic puncta markers and by WB 193 analysis of synaptic vesicle protein, SV2A, levels. Remarkably, IF and WB results were consistent with a 194 reduction of excitatory and inhibitory synaptic puncta (Supplemental Figure 4D) and SV2A protein levels 195 (Supplemental Figure 4E), suggesting that cognitive behavioral deficits in SNX14 deficiency are likely 196 197 caused by defects in synaptic signaling of forebrain cortical neurons.

Unlike cerebral cortices, cerebellar transcriptomes were markedly different between Snx14 KO 198 and WT mice, with 160 upregulated and 6 downregulated DEGs at 1 month of age and 142 up- and 222 199 200 downregulated DEGs at 1-year Snx14 KO (Figure 4B, D-G and Supplemental Figure 5A). We reasoned that the increase in the amount of downregulated DEGs from 1 month to 1 year of age could reflect the 201 202 progressive PC loss in Snx14 KO cerebella. Accordingly, most downregulated DEGs in 1-year Snx14 KO 203 cerebella correspond to PC markers, such as Calb1, Pcp2, Car8, and Rgs8 (Figure 4B). To unbiasedly test this observation, we analyzed 1-year-old DEGs for functional annotation enrichments. Furthermore, we 204 compared DEGs with a list of a recently reported mouse cerebellar single nuclear RNAseq dataset (30). 205 206 Results confirmed that downregulated DEGs are enriched in genes predominantly expressed in PCs

(Supplemental Figure 5B-C). In contrast, most of the upregulated DEGs genes are sparsely expressed 207 across various cerebellar cell types, with a group of them typically expressed in astrocytes and 208 macrophage/microglia (Lyz2, C4b, Cd68, Trem2, ApoE, Gfap) or associated with cell death (Casp3) 209 (Figure 4B and Supplemental Figure 5B- C). Notably, 1-month-old DEGs were not enriched for PC or 210 astroglia specific functional annotations indicating a later onset of neurodegeneration, consistent with our 211 212 histological analyses (Figure 4B and Supplemental Figure 5B). Additionally, functional annotation analysis revealed enrichments of genes localized in synaptic, dendritic, and ER compartments in pre- and 213 214 post-degenerating cerebella (Figure 4D and Supplemental Data 2).

Given the lack of neurodegenerative signs in histology or transcriptomic data (Figure 3 and 4), we 215 anticipated that DEGs at 1-month-old cerebella could point us to the molecular causes that precede PC 216 neurodegeneration. However, considering that PCs only constitute  $\sim 1\%$  of the total cerebellum, 217 transcriptomic changes in PCs may only contribute to small fold changes in bulk transcriptomic data. To 218 account for these small changes, we analyzed our RNAseq data with Gene Set Enrichment Analysis 219 220 (GSEA). Interestingly, GSEA revealed cerebellar specific enrichments in biological processes involved in oxidative stress (e.g. 'response to oxygen containing compounds' in 1-month-old cerebella and 221 'response to reactive oxygen species' in 1-year-old cerebella), fatty acid or lipid homeostasis regulation 222 223 (e.g. 'response to positive regulation of unsaturated fatty acid biosynthetic process' in 1-month-old cerebella and 'response to lipid' in 1-year-old cerebella) and iron accumulation (i.e. 'regulation of iron 224 225 ion transmembrane transport' and 'iron ion binding' in 1-year-old cerebella) (Figure 4E-G). Remarkably, 226 genes contributing to GSEA enrichments in pre-degenerating cerebella include upregulated Fabp5, which encodes a protein involved in interorganelle lipid transport (33), and *Dcn* encoding a protein released by 227 228 cells dying by ferroptosis (34) (Figure 4B, E-G). These data suggest that lipid homeostatic defects may 229 precede selective cerebellar degeneration in SNX14 deficiency.

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#### 1 Snx14 deletion alters lipid metabolite levels in a tissue specific manner

We next set out to analyze lipid metabolite composition of pre-degenerating cerebella in 2-monthold WT and *Snx14* KO mice by unbiased lipidomic analysis. As a control of a non-degenerating tissue, we included their cerebral cortices in the analysis. Since the liver is a lipid rich organ with high content of TGs stored in LDs, we included liver lipid extracts as a control for lipid metabolite detection. Finally, to distinguish tissue specific lipids from those circulated by their blood supply, we also extracted plasma lipids from circulating blood.

The lipid extracts were analyzed by ultraperformance liquid chromatography-high resolution mass 238 spectrometry (UPLC-HRMS) as previously described (35) and after normalization with lipid internal 239 standards, we quantitatively identified >200 lipid species per sample (Supplemental Data 3). Overall, 240 Snx14 KO and WT tissues had similar total lipid concentrations (Supplemental Figure 6A) and each tissue 241 analyzed was distinguishable by their relative lipid class abundance. For instance, liver displayed the 242 243 highest abundance of TGs while the cerebral cortex and cerebellum had phosphatidylcholines (PCh) as the most abundant lipid class (Supplemental Figure 6B). This data is consistent with the literature (36, 244 37), thus validating our methodology. 245

Next, we aimed to determine how SNX14 deficiency affects tissue specific lipid composition. To this end, we compared the concentration of each lipid specie in Snx14 KO and corresponding WT tissue. Given SNX14's role in facilitating the incorporation of FAs into TG during LD biogenesis (22), we hypothesized that SNX14 deficiency would result in a depletion of TG levels. Although TGs were undetectable in all cerebellar and cortical samples, Snx14 KO livers displayed a significant reduction of TGs (Figure 5A and Supplemental Figure 6B), further confirming our hypothesis and the reliability of our lipidomic analysis. Additionally, results showed that the cerebral cortex and cerebellum are the tissues with the largest amount of altered lipid species upon SNX14 depletion (Figure 5A). Using p-value < 0.05 as a cutoff, we identified 58 and 36 altered lipid species in cerebral cortices and cerebella, respectively. Furthermore, only cerebellar samples clustered by genotype in a principal component analysis (Supplemental Figure 6C), suggesting SNX14 has a larger impact on lipid homeostasis in cerebella than in the other tissues we analyzed.

259 Among the 58 altered lipids in cerebral cortices, 54 had lower concentrations in KOs, and 40 belong to the phosphatidylethanolamine (PE) class (Figure 5A-C). PEs provide fluidity and curvatures to 260 261 membranes which may facilitate vesicular budding and membrane fusion essential for synaptic vesicle formation (38). Thus, changes in PE species may alter cerebral cortex-dependent behaviors and executive 262 functions in SNX14 deficiency. The remaining 4 lipid species had higher concentrations in Snx14 KO 263 than in WT and all were sphingomyelins (SMs) (Figure 5A and Supplemental Figure 6E). Similarly, Snx14 264 KO cerebella exhibited increased levels of total SM concentrations (Figure 5A and Supplemental Figure 265 6E). While some PEs were lower in Snx14 KO cerebella, these did not influence total PE concentration 266 (Figure 5A-B). In addition, Snx14 KO cerebella were distinguishable from the cortex, liver, and plasma 267 268 given the increased levels of several acylcarnitine (AcCa) species (Figure 5A, E-F). Specifically, 6 out of 16 increased lipids in Snx14 KO cerebella were AcCa-s. This accounted for the majority of AcCa-s 269 detected in cerebella (6 out of 8) and resulted in an overall increase of total AcCa concentration in Snx14 270 271 KO cerebella. Snx14 KO cerebellar samples were also the ones with the largest amount of accumulated lipid species among all the analyzed tissues. 272

To further determine region specific differences in lipid metabolite abundance *in situ*, we next analyzed brain sections with matrix-assisted laser desorption ionization and mass spectrometry imaging (MALDI-MSI). Results uncovered differences between WT and *Snx14* KO brain lipid patterns consistent

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with UPLC-HRMS results, including reduced PE C38:2 levels (Figure 5F). Furthermore, two TGs were reduced in Snx14 KO cerebella, one of which (TG 53:2) is specific to the outermost layer of the cerebellar cortex comprised by PC soma and dendrites (Figure 5F). L-carnitine signal also overlapped this area and was more intense in Snx14 KO than WT (Figure 5F). Given L-carnitine's involvement in AcCa metabolism, this increase in signal may be associated with the accumulation of AcCa in this cerebellar region.

Taken together, the bulk and *in situ* lipidomic analyses show tissue specific lipid metabolite defects including a cerebellar specific AcCa accumulation that may be associated with the selective cerebellar neurodegeneration characteristic of SNX14 deficiency.

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#### 286 Snx14 deletion impairs lipid storage in vivo

Under conditions of high energy demand or nutrient deprivation, AcCa-s carry FAs into the 287 mitochondria for beta oxidation. Elevated concentrations of AcCa, however, can become cytotoxic and 288 289 disrupt mitochondrial function. Here, LDs are vital by storing excessive FAs and preventing AcCa induced toxicity (39). Accordingly, increased AcCa and decreased TG levels in Snx14 KO cerebella could be a 290 consequence of defects in LD biogenesis. In line with this idea, SNX14 interacts with LDs and its 291 292 deficiency leads to impaired LD content and morphology in cell cultures (22). Thus, we investigated whether Snx14 deletion alters LD biogenesis in the cerebellum by staining Snx14 KO and WT mice 293 294 cerebella with Bodipy 493/503 (BD493), a fluorescent dye that stains neutral lipids typically stored in 295 LDs. As a control, we stained the liver, a LD rich tissue, and detected abundant BD493 positive LDs in WT sections. Results also revealed a prominent reduction of LD amounts in Snx14 KO liver (Figure 6A), 296 297 which is consistent with the reduction of TG levels in Snx14 KO liver lipidomics (Figure 5A and Supplemental Figure 6B). These data suggest that SNX14 is necessary for LD biogenesis *in vivo*, at least
in the liver.

Next, we focused our attention on the cerebellum. Here, BD493 staining showed few, if any, structures resembling LDs, even in WT PCs (Supplemental Figure 7A). To further explore the possibility that *Snx14* deletion affects LD biogenesis in PCs, we stimulated LD biogenesis in cerebellar cultures by supplementation with oleic acid (OA). As expected, OA induced LD biogenesis in WT PCs (Figure 6B). In contrast, the number of LDs detected in *Snx14* KO PCs was half the number in WT PCs (Figure 6B), indicating that SNX14 is necessary for LD biogenesis also in PCs.

To assess for LDs or alternative lipid storage defects in Snx14 KO PCs in tissue, we analyzed 306 cerebellar sections by transmission electron microscopy (TEM) after imidazole-buffered osmium 307 tetroxide staining to highlight LDs (40) (Figure 6C-I). An overview of PC integrity in TEM images 308 confirmed that most Snx14 KO PCs are still intact at pre-degenerating ages (2 months) (Figure 6C) while 309 a gradient of degenerating PCs is observed at 6 months of age (Figure 6D and Supplemental Figure 7B). 310 311 Again, TEM studies failed to identify LDs in the cerebellum at pre-degenerating (2 months) or postdegenerating stages (6 months). Nonetheless, TEM results revealed that at pre-degenerating ages, Snx14 312 KO PCs have less and larger telolysosomes, which are lipid rich lysosomal storage organelles (Figure 6E-313 314 F and Supplemental Figure 7B). Interestingly, this is consistent with larger lysosome compartments we observed in Snx14 KO PCs (Figure 3D) and in SCAR20 patient neural cell lines (16). These results 315 316 suggests that SNX14 may have a specialized function regulating lipid clearance or storage through the 317 lysosomal compartment in PCs.

Less, yet enlarged telolysosomes in PCs, and elevated AcCa-s at pre-degenerating *Snx14* KO cerebella, suggest that lipid homeostasis defects underlay PC degeneration in SNX14 deficiency (39). Nevertheless, increased AcCa can also be a consequence of mitochondrial damage. Therefore, to

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determine if AcCa accumulation is a consequence of lipid storage defects or caused by mitochondrial damage, we assessed mitochondrial ultrastructure of PCs by TEM. Results showed mostly intact mitochondria in pre-degenerating *Snx14* KO PCs (Figure 6G and Supplemental Figure 7C), suggesting that AcCa accumulation is the result of lipid storage and clearance defects, not mitochondrial damage. Furthermore, at 6 months, ultrastructure analysis also revealed a progressive enlargement of ER as PCs degenerate. Damaged mitochondria with enlarged and disorganized cristae were only observed in most degenerated PCs (Figure 6I).

Together, our work indicates that lipid storage and clearance defects are associated with PC neurodegeneration in SNX14 deficiency contributing to the expanding list of neurodegenerative disorders associated with lipid homeostasis defects.

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#### 332 **Discussion**

SNX14 deficiency causes a childhood onset cerebellar degeneration syndrome clinically defined 333 334 as SCAR20 and characterized by cerebellar ataxia and intellectual disability. Previous work identified lysosome and autophagy specific defects in cultured patient neural progenitor like cells (16, 17) and recent 335 evidence implicates SNX14 in LD biogenesis, FA desaturation, and non-vesicular interorganelle lipid 336 transport (19, 22-24). However, most of these studies were performed in cultured cells with unclear 337 338 relevance for SCAR20 pathology. To overcome this limitation and study pathogenic mechanisms that 339 selectively affect the cerebellum, we generated a *Snx14* KO mouse that closely recapitulates SCAR20 at genetic and phenotypic level. Consistent with a widespread expression of SNX14, we find that SNX14 340 341 deficiency in vivo leads to tissue specific lipid metabolite and storage defects that likely result from cell type specific lipid homeostatic requirements. Remarkably, pre-degenerating Snx14 KO cerebella is 342 distinguishable from non-degenerating cerebral cortex by a unique accumulation of AcCa and L-Carnitine, 343

including depletion of TG levels. These data, combined with reduced LD numbers and enlarged telolysosomes in pre-degenerating PCs, suggest that lipid homeostasis defects cause cerebellar degeneration in SNX14 deficiency. However, due to lack of cellular resolution, our data does not rule out the possibility that other cell types in the cerebellum may also contribute to the lipidomic changes and neurodegeneration in *Snx14* KO cerebella.

Lipid homeostasis disruption is associated with many cerebellar neurodegenerative disorders (8). 349 350 Little is known, however, about the mechanisms that preserve lipid homeostasis in the cerebellum. PCs are fast and high frequency spiking neurons with a large membrane area, which makes them particularly 351 susceptible to oxidative stress induced by membrane lipid peroxidation (41). Portions of membranes that 352 353 contain peroxidated lipids are often cleared by autophagy, which leads to an overproduction of FAs and their storage in LDs as a protective mechanism from excess FA induced damage (39, 42, 43). Although 354 neurons produce few LDs, recent evidence implicates autolysosome derived structures in the clearance of 355 toxic lipids through exocytosis in neurons (44). Therefore, our data showing abnormal lipidomic profiles 356 357 and lipid storage and clearance organelles in pre-degenerating cerebella, fit with a model implicating SNX14 in the storage (lipid droplet) and clearance (lysosome) of toxic lipids generated in PCs. Notably, 358 SNX14 has been associated with lysosome function regulation (16) and recent structural predictions 359 suggest a role in inter-organelle lipid transport (24) that may be important for PC specific lipid 360 361 homeostasis. Furthermore, lipid clearance and storage defects have recently been associated with neuronal 362 ferroptosis (44) and our transcriptomic data show upregulation of ferroptosis associated genes (i.e. Dcn and Fabp5) in pre-degenerating Snx14 KO cerebella and genes associated with iron at older ages. These 363 364 data suggest an exciting hypothesis implicating lipotoxicity induced ferroptosis as a pathogenic mechanism of cerebellar degeneration in SCAR20 that warrants future investigation. 365

Given the widespread expression of Snx14, it is possible that other cell types contribute to PC 366 degeneration. Indeed, glia cells have a central role in the clearance and metabolism of neuronal lipids (44-367 368 46). Remarkably, loss of PCs in *Snx14* KO cerebella overlaps with a robust gliosis. Given the enrichment of Snx14 expression in cerebellar PCs reported in the literature and our RNAscope analyses, we predict 369 that PC degeneration is primary to SNX14 deficiency which then triggers gliosis in Snx14 KO cerebella. 370 371 In agreement, the pre-degenerating Snx14 KO cerebellar transcriptomic data shows an upregulation of Dcn, which encodes a protein that stimulates the immune response after being released by cells dying 372 from ferroptosis (34). Recent reports also suggest that gliosis is induced by PC degeneration in cerebellar 373 ataxias. For example, PC specific expression of mutant *ataxin1* in Sca1154Q/2Q mice is enough to induce 374 astrogliosis and microgliosis (47) and deletion of mutant ataxin-7 from PCs prevents gliosis in SCA7-92Q 375 BAC mice (48). Future studies will investigate if the loss of SNX14 affects lipid homeostasis in glia and 376 whether this contributes to the selective cerebellar degeneration in SNX14 deficiency. 377

Similar to recently reported SNX14 deficient mice (25, 49), the homozygous 1bp deletion in our 378 379 Snx14 KO mice causes loss of full length SNX14 protein and low RNA counts across all coding exons. Unlike previous models that showed fully penetrant embryonic lethality (25, 49), ~a third of our Snx14 380 KO mice develop and survive to adulthood with a phenotype that resembles SCAR20. This finding 381 suggests that SNX14 deficiency in humans may also interrupt embryonic development, and cause 382 383 SCAR20 only when embryonic lethality is circumvented. Although we still do not know what factors 384 determine the developmental success or failure in SNX14 deficiency, there is a striking difference in the genetic architecture of Snx14 mutations between organisms that show full and partial embryonic lethality. 385 386 For instance, SNX14 deficient mice that completely fail to develop carry deletions of at least one full exon while SCAR20 patients and animal models, including our Snx14 KO mice, dogs (50) and zebrafish (25) 387 carry truncating point mutations or small indels. This observation has interesting implications for the 388

generation of animal models of human disorders and for the pathogenic prediction of truncating genetic
 mutations that warrant further investigation.

Another factor that can influence the outcome of SNX14 deficient embryos is the environment and 391 diet lipid composition. In line with this idea, SNX14 deficient cells are more vulnerable than control cells 392 to saturated FAs (25) and treatment with valproic acid, a branched short-chain FA, partially rescued PC 393 degeneration in a conditional mouse model (49). Furthermore, previous studies have shown that maternal 394 395 diet lipid composition can modulate brain lipidome either embryonically by maternal feeding or in adult mice (37). Altogether, these data open a window to alter the course of SCAR20 through therapeutic diets. 396 In this regard, further elucidating mechanisms that preserve lipid homeostasis in neurons, and particularly 397 398 in the cerebellum is of crucial relevance.

Overall, our work highlights the relevance of lipid homeostasis for neurodegenerative disorders and suggest a mechanism for increased susceptibility of the cerebellum to the expanding class of disorders caused by disrupted lipid metabolism pathways. Furthermore, our study provides a mouse model and molecular targets for future therapeutic studies.

403

#### 404 Materials and Methods

405 Detailed materials and methods are available in Supplemental Methods.

406

#### 407 Sex as a biological variable

408 Our study examined male and female animals, and similar findings are reported for both sexes.409

410 *Animals* 

411 Generation of mouse model

412 *Snx14* KO mice were generated by pronuclear injection of 5ng/ul Cas9 mRNA and 2.5ng/ul sgRNA (5'-413 GTAAACACGTTCTCCAAC-3') in 1 cell stage fertilized embryos obtained from superovulated 414 C57BL/6J females mated with C57BL/6J males. Pups carrying *Snx14* indel alleles were selected for 415 backcross with WT C57BL/6J mice for 3-6 generations (to filter out potential off targets) and further 416 expanded as an experimental model. Only the *Snx14* c.1432delG carriers generated homozygous pups.

417

#### 418 **Behavior analysis**

#### 419 Experimental design

Behavior analysis was performed with three cohorts of WT and *Snx14* KO littermates starting at 8 months of age. Each cohort contained mixed genotype and sex of animals. Behavior tests were performed in the following order: accelerating Rotarod, Catwalk, Metz Ladder and Social choice/recall. Investigators were blinded during scoring of behavioral assessments. Whenever possible, offline analysis by computer software was utilized to enhance rigor.

425

#### 426 Accelerating Rotarod

On day 1, mice were habituated to the stationary Rotarod for 2 minutes. This was immediately followed 427 428 by a trial where rotation was programmed to rise from 4-40rpm in 300 seconds. After a 30-minute intertrial interval (ITI), a second trial was performed, followed by another ITI and third trials. Three additional 429 430 trials were performed on the next 2 consecutive days, for a total of 9 trials. A trial was terminated when a 431 mouse fell, made one complete revolution while hanging onto the rod, or after 300s. Latency to fall (time stayed until falling or riding the rod for a single revolution) was determined. Learning rate was calculated 432 433 as followed: learning rate = (Trial 9 latency to fall – Trial 1 latency to fall)/8, 8 is the number of inter-trial 434 intervals in this study.

435

#### 436 Catwalk gait analysis

In the Catwalk gait analysis assay, mice were placed on a meter-long illuminated glass plate walkway in
a dark room. A high-speed video camera below the plate recorded the paw prints, as the mice traversed a
20cm section of the alley. The paw print footage was analyzed by CatWalk XT program (Noldus,
Leesburg, VA).

441

#### 442 *Metz ladder rung waking test*

The Metz procedure used a 1-meter-long horizontal ladder, which was about 1cm wider than the mice. 443 The Plexiglas walls were drilled with 3mm holes to accept the metal rungs. The gaps between the rungs 444 were randomly spaced 1-5 cm apart so that the mice had to adjust the projection of the landing of each 445 paw. Mice were trained to run the ladder with all rungs in place, 1cm apart before the test trials began. In 446 the test, each mouse was placed at the beginning of the ladder. Five trials were performed on consecutive 447 448 days and videotaped. The pattern of the rungs was changed after each trial to prevent animals from adapting. Trials were recorded by a high-definition digital camera. Foot slip(s) of each trial was quantified 449 later by an investigator blinded to group designation with video. 450

451

#### 452 Social choice and recall test

Mice were tested for social preference and recall as described previously (51). The testing apparatus was a rectangular Plexiglas three chamber arena (60 cm (L)  $\times$  40 cm (W)  $\times$  20 cm (H)). The chamber was continuous with areas at the ends designated for the placement of vented cylinders to hold the cues. The social cues were juvenile, sex-matched C57BL/6J mice. The inanimate cues were smooth rocks that approximate the size of the social cues. The procedure consisted of a habituation phase whereby the

experimental mouse was placed into the center chamber with empty cylinders in the side chambers for 10 458 minutes. After habituation, the choice phase immediately began. The cylinders were loaded with either a 459 social cue (young mouse, M1) or inanimate cue. The experimental mouse was allowed to explore the cues 460 for 10 minutes. Immediately after the choice phase, the recall phase was performed. The now familiar 461 social cue, M1 remained in a cylinder while a novel mouse, M2 was loaded into the cylinder that 462 463 previously held the inanimate cue. The experimental mouse was allowed to freely explore the 2 social cues for 10 min. The bouts and duration of explorations (nose <1 cm proximity) with the cylinders was 464 determined with ANYmaze software (Stoelting Co. Wood Dale II.). 465

- 466
- 467 *Histology*

#### 468 Immunofluorescence staining

Mice were anesthetized with isoflurane (Terrell) and perfused trans-cardially with 20ml 1X PBS and 20ml 4% paraformaldehyde (PFA) (Electron Microscopy Sciences). Brains dissected out from scalp were postfixed in 4% paraformaldehyde for 18h in RT and washed 3 x 10 mins in 1X PBS. Brains were sliced into 50um sections using a vibratome (Leica).

473

On the day of staining, slides were washed with 1X PBS, permeabilized and blocked with PBS+0.3% Triton X-100 (PBST) and 4% goat serum (G9023, Sigma-Aldrich) for 45 min at room temperature. Slides were then incubated with primary antibodies (see Supplemental Methods) in 2% goat serum in PBST at 4°C on the shaker overnight. Next day, slides were washed with PBST 3 x 10 mins and incubated with Alexa Fluor-conjugated secondary antibodies at 1:500 in 2% normal goat serum in PBST for 2h at room temperature (RT). Slides were washed in PBST 3 x 10 mins, incubated with 300 nM DAPI (D3571, Invitrogen) for 10 min at RT and mounted on microscope slides with ProLong Gold antifade (P36930,

481	Invitrogen) or Mowiol (#81381, Sigma) covered with a coverslip. Immunostainings were imaged with a
482	Leica TCS SP8 X confocal microscope and images processed and quantified with ImageJ (NIH).
483	
484	RNAscope In situ hybridization
485	The RNAscope in situ hybridization was performed as recommended by the manufacturer with reagents
486	from Advanced Cell Diagnostic (USA) (see Supplemental Methods). Once RNAscope was completed,
487	immunofluorescent staining was immediately performed as described above. Sections were imaged with

488 a Leica TCS SP8 X confocal microscope and images processed and quantified with ImageJ.

489

#### 490 **BODIPY staining**

Fixed brain and liver tissue was sliced into 50  $\mu$ m sections using vibratome, rinsed in PBS and incubated with 2  $\mu$ M BODIPY 493/503 (D3922, Invitrogen) for 30 min at RT with gentle rocking. Then, the sections were rinsed in PBS 3 x 10 mins and mounted on microscope slides with Mowiol and covered with coverslips.

495

#### 496 Transmission Electron Microscopy (TEM)

Mice were perfused with 20 mL of PBS, followed by 20 mL 2% PFA and 2% glutaraldehyde in sodium cacodylate buffer. Cerebella were dissected, trimmed to 1 mm thickness, and processed for TEM at the University of Delaware's Bio-Imaging Center. Briefly, tissues were washed 3 x 15 min in 0.1M sodium cacodylate buffer pH 7.4 and post-fixed for 2 h with freshly prepared 1% osmium tetroxide and 1.5% potassium ferrocyanide in 0.1M sodium cacodylate buffer pH 7.4 or alternatively, to improve lipid droplet detection, with 1% osmium tetroxide in 0.1M imidazole pH 7.5. The tissue was washed with water, dehydrated through an ascending acetone series, and then infiltrated with Embed-812 resin. The next day, samples were embedded in flat-bottom capsules and polymerized at 60°C overnight. Ultrathin sections were cut using a Leica UC7 ultramicrotome and placed onto single hole 1500-micron copper aperture grids with a formvar/carbon film. Sections were post-stained with 2% uranyl acetate in 50% methanol and Reynolds' lead citrate and examined on a ThermoFisher Scientific Talos L120C transmission electron microscope operating at 120kV. Images were acquired with a ThermoFisher Scientific Ceta 16M camera. Quantification of area and numbers was done by ImageJ.

510

### 511 *Matrix assisted laser desorption ionization coupled to time-of-flight mass spectrometry (MALDI-TOF*

512 MS) Imaging

MALDI-TOF MS imaging was carried out in MALDI MS Imaging Joint Facility at Advanced Science
Research Center of City University of New York Graduate Center.

8-weeks-old mouse brains were cryosectioned (10 µm thickness) sagittally and gently transferred onto 515 the pre-cooled conductive side of indium tin oxide (ITO)-coated glass slides (Bruker Daltonics) for 516 MALDI imaging. Mounted cryosections were desiccated in vacuum for 45 min at RT, followed by matrix 517 deposition using HTX M5 sprayer (HTX Technologies, LLC) on DHB matrix (40 mg/mL in 518 519 methanol/water (70/30, v/v), flow rate of 0.05 mL/min and a nozzle temperature of 85 °C for 8 cycles) to detect metabolites and lipids. MALDI mass spectra were acquired in positive ion mode (DHB) acquired 520 by MALDI-TOF MS Autoflex (Bruker Daltonics). MALDI imaging data were recorded and processed 521 522 using FlexImaging v3.0, and further analyzed using SCiLS (2015b). Ion images were generated with rootmean square (RMS) normalization and a bin width of  $\pm 0.15$  Da. The spectra were interpreted manually, 523 and analyte assignment was achieved by comparing with LC-MS/MS experiment results (52). The signal 524 525 intensity of the cortex and cerebellum regions of three animals of each genotype were quantified using

SCiLS and further analyzed using GraphPad. P-value between control and mutant animals were analyzed
by Student's t-test using three animals of each group.

528

#### 529 Cell Culture

#### 530 Purkinje Cell Culture

531 Primary mixed cerebellar cultures were generated and maintained as described (53). Briefly, cerebellums were isolated from E16.5 of WT or Snx14 KO mice, dissociated and plated at 50,000 cells on coverslips 532 coated with 0.1 mg/mL poly-D-lysine in recovery media (DMEM/F-12, (#11330032, Gibco) 533 supplemented with 1% Penicillin-Streptomycin (#15140122, Gibco), 1X B-27 (# 17504044, Gibco), 10% 534 FBS (#101, Tissue Culture Biologicals), 20 ug/mL Insulin (#I9278, MilliPore Sigma), and 100 ug/mL 535 IGF-1 (#100-11, PeproTech). Two hours later, recovery media was removed and replaced with complete 536 media (DMEM/F-12, supplemented with 1% Penicillin-Streptomycin, 1X B-27, 1% FBS, 20 ug/mL 537 Insulin, and 100 ug/mL IGF-1). Purkinje cells were cultured for 7 days in vitro before processing for 538 539 experiments.

540

#### 541 Purkinje Cell Lipid droplet Analysis

To promote LD biogenesis, cerebellar cultures were incubated with 600 uM Oleic Acid (#O1008, Sigma) conjugated to 100 uM fatty acid-free BSA (A1595, Sigma) overnight. Cells were fixed 10 min with 4% PFA at RT, washed with 1X PBS and blocked in blocking buffer (1.5% Glycine, 3% BSA, 0.01% Saponin in 1X PBS) for 1h at RT. Cells were incubated overnight at 4°C with primary antibodies in antibody solution (1% BSA, 0.01% Saponin in 1X PBS). The following day cells were washed, incubated in secondary antibodies with 300nM DAPI, and 2 $\mu$ M BODIPY 493/503 for 2h at RT and mounted with Fluoromount-G (#00-4958-02, Invitrogen). Images for quantification were captured with Leica TCS SP8 X confocal microscope and BODIPY 493/503 positive puncta quantified with 'analyze particles' plug-in
 in Fiji-ImageJ after processing with "Intermodes" algorithm.

551

#### 552 Biochemical studies

#### 553 Western blot

554 Mouse tissue was dissected, fast-froze, and stored in -80°C until use. On the experiment day, tissue was homogenized in RIPA buffer (#9806, Cell Signaling) supplemented with a protease inhibitor cocktail 555 (P8340, Sigma-Aldrich) and incubated for 15 minutes at 4C. After centrifugation at 13,200 rpm, 556 supernatant containing protein extract was collected, mixed with 1X LDS loading buffer (B0007, 557 Invitrogen) supplemented with 200 mM DTT (BP172-5, Fisher Scientific) and loaded on a 4-15% Mini-558 Protean TGX Precast Protein Gel. Proteins were transferred onto PVDF membranes in Mini Gel Tank at 559 80V for 180 min. Membranes were blocked with 5% milk-TBST or EveryBlot Blocking Buffer 560 (#12010020, Bio-Rad) for 1h at RT then probed with primary antibodies diluted in 5% milk-TBST or 561 EveryBlot Blocking Buffer solution overnight at 4°C. The next day, membranes were washed and probed 562 with horseradish-peroxidase-conjugated secondary antibodies for 1h at RT, incubated in either Pierce<sup>TM</sup> 563 ECL Western Blotting Substrate kit (#32106, Thermo Scientific) or SuperSignal<sup>TM</sup> west dura extended 564 565 duration substrate (34076, Invitrogen) and exposed on autoradiography film following development in AFP Mini-Med 90 X-Ray Fil Processor. Exposed films were scanned, and protein bands were quantified 566 567 using ImageJ.

568

#### 569 **RNA-seq**

1-month-old or 1-year-old mice were euthanized, and tissue was dissected on ice, fast frozen, and stored
in -80°C until RNA extraction. Total RNA from the cerebellum or cerebral cortex were isolated using

TRIzol (15596026, Invitrogen) reagent according to the manufacturer's instructions. Strand-specific 572 mRNA-seq libraries for the Illumina platform were generated and sequenced at GENEWIZ or Novogene 573 following the manufacturer's protocol with sample specific barcodes for pooled sequencing. After 574 sequencing in Illumina HiSeq or Novoseq platform with 2x150 PE configuration at an average of 15 575 million reads per sample, sequenced reads were trimmed to remove possible adapter sequences and poor-576 577 quality nucleotides and trimmed reads mapped to the Mus musculus GRCm38 reference genome using Spliced Transcripts Alignment to a Reference (STAR v2.7.3a) software. Reads were counted using 578 FeatureCounts from the subread package (v2.0.1) (54). Transcripts Per Million (TPM) values were 579 580 calculated from featureCounts-derived counts. Heatmap of gene expression were generated using the tidyverse R package with z-score of the log2(tpm+1). Differential gene expression analysis was performed 581 with DEseq2 (v1.38.3). Raw p-values were adjusted using the Benjamini-Hochberg method. Differentially 582 expressed genes (DEGs) were defined as having an adjusted p value (Padj)<0.05. Volcano plots were 583 generated with the EnhancedVolcano R package. Functional enrichment analysis was conducted utilizing 584 585 the enrichR R package. Gene Set Enrichment Analysis (GSEA) was performed on the Mus musculus msigdbr database in the C5 ontology category. Relevant lipid, oxygen, or iron -related terms were 586 manually selected and displayed in the waterfall plots, generated through the tidyverse R package. 587

588

#### 589 UPLC-HRMS whole lipidome analysis

#### 590 Sample preparation

2-months-old mice were euthanized and, after heart blood collection, cortex, cerebellum, and liver were dissected, snap-frozen in liquid nitrogen and stored at -80 until lipid extraction. For lipid extraction, plasma samples were prepared as previously reported (55).  $\sim$ 10 mg of frozen tissue fragments were weighted, chopped and mixed with 0.6 mL 80% methanol (MeOH) and 10 µL on internal standard mix

(SPLASH® LIPIDOMIX #330707 from Avanti Polar Lipids, Alabaster, AL). Samples were pulse 595 sonicated in ice for 30x 0.5 second, incubated for additional 20 min in ice, vortexed 3x 30 seconds and 596 tissue homogenates transferred to a 10 mL glass Pyrex tube with screw cap. Then, 5 mL methyl tert-butyl 597 ether (MTBE) was added to each tube and vigorously shacken for 30 minutes, followed by the addition 598 of 1.2 mL water and 30 second vortex. Samples were centrifuged for 10 min at 1000 g and the top clear 599 600 phase was collected to a clean glass Pyrex tube and dried down under nitrogen. For the analysis, dried samples were resuspended in 100  $\mu$ L MTBE/MeOH=1/3 (v/v), spun down at 10,000g for 10 min at 4°C. 601 The top 50 µL were transferred to a HPLC vial and 2ul were injected for LC-MS analysis. 602

#### 603 Liquid chromatography high resolution -mass spectrometry (LC-HRMS) for lipids

604 Separations were conducted on an Ultimate 3000 (Thermo Fisher Scientific) using an Ascentis Express C18,  $2.1 \times 150$  mm 2.7µm column (Sigma-Aldrich, St. Louis, MO). For the HRMS analysis, a recently 605 calibrated QE Exactive-HF mass spectrometer (Thermo Fisher Scientific) was used in positive ion mode 606 with an HESI source. Control extraction blanks were made in the same way using just the solvents instead 607 608 of the tissue homogenate. Untargeted analysis and targeted peak integration was conducted using LipidsSearch 4.2 (Thermo Fisher Scientific) as described by Wang et al (56). Lipids quantification was 609 done from the full scan data. The areas were normalized based on the amount of the internal standard 610 611 added for each class. All amounts were then normalized to the original tissue weight.

612

#### 613 *Statistics*

614 Statistical analyses were performed using GraphPad Prism 8 (GraphPad Software, Inc.). When possible, 615 data was analyzed blind to the genotype. Sample size for each experiment was determined based on similar 616 studies. To compare the means of groups where normal distribution and similar variance between groups 617 was confirmed, two-tailed Student's *t*-test (for two samples), one-way ANOVA (for more than two

618	samples) or two-way ANOVA followed by Sidak's or Tukey's post hoc test (for multiple variables) was
619	used. A $P$ value less than 0.05 was considered statistically significant. Outliers were removed in two
620	behavioral studies using the ROUT method with Q=1%, $p < 0.0002$ .
621	
622	<u>Study approval</u>
623	All animal procedures were performed according to NIH guidelines and approved by the Institutional
624	Animal Care and Use Committee (IACUC) at Children's Hospital of Philadelphia.
625	
626	Data availability
627	RNAseq data was deposited in GEO under the GSE215834 reference. Whole data from lipidomic
628	analysis is available in Lipidomic data file. All other data are available in the Supporting Data Values
629	file.
630	
631	Author contributions
632	Study conceptualization and design: Y.Z., V.S., and N.A. Validation and maintenance of mouse colony:
633	Y.Z., T.J., and N.A. Behavioral study design, execution, and data collection: B.C. and T.O. Behavioral
634	data analysis: B.C., T.O., Y.Z., and H.T. Histology, cell culture, and TEM studies: Y.Z., V.S., M.F., and
635	D.Y. RNA extraction and RNAseq analysis: Y.Z. Lipidomic analysis: Y.Z., P.X., and C.M. MALDI MS
636	Imaging analysis: S.L. and Y.H. Data interpretation: Y.Z., V.S., M.H., and N.A. Supervision and project
637	administration: N.A. Manuscript preparation: Y.Z., V.S., and N.A. Manuscript edit and review: All
638	authors.
639	

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651

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#### 778 Figure 1





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Figure 2. SNX14 deficiency in mice recapitulates motor and behavioral deficits of SCAR20. (A-B) Catwalk analysis shows altered gait of KO mice with a longer stand (A) and shorter swing (B) than WT mice. Bar graphs represent mean ±S.E.M of n=24 WT and n=18 KO mice. Two-tailed Welch's t-test. (C) Metz ladder rung test shows altered limb placing and coordination of KO males and females. Bar graphs represent mean foot slip of 5 trials performed in consecutive days ±S.E.M of n=10 WT males, n=7 KO males, n=12 WT females, n=12 KO females. Two-way ANOVA followed by Sidak's test. (D) Accelerating rotarod reveals defects in motor performance of KO mice in the 9 trials performed over 3 consecutive days. Graphs show mean latency to fall ±S.E.M of n=11 WT males, n=7 KO males, n=13 WT females, n=11 KO. Two-way ANOVA test shows significant effect of genotype. (E) KO females show impaired learning rate on accelerating rotarod performance over time (between trial 1 and 9). Bar graph shows mean learning rate ±S.E.M of n=9 WT males, n=7 KO males, n=13 WT females, n=10 KO females. Two-way ANOVA followed by Sidak's test. (F-G) Three-chamber social interaction test showing similar preference for a mouse over an object between WT and KO mice (F) but impaired social novelty preference in KO mice (G). Bar graphs represent mean and S.E.M of WT n=24, KO n=17. Two-way ANOVA followed by Tukey's test. n.s. = non-significant, \**P* <0.05, \*\**P* < 0.01, \*\*\**P* < 0.001, \*\*\*\**P* < 0.0001. 

Figure 3 822





824 Figure 3. SNX14 deficiency causes selective cerebellar degeneration. (A) Representative brain images from WT and KO 825 mice at indicated age shows shrinkage of KO cerebellum (CB) over time. Ruler marks separated by 1mm. Bar graphs show 826 percentage area of CB or cerebral cortex (CX) relative to the whole brain (WB) in n=3-5 mice. Two-way ANOVA followed 827 by Sidak's test. (B) Representative cerebellar sagittal sections immunostained with PC specific anti-CALB1 antibody reveal 828 progressive loss of PCs in KO mice. Bar graphs show PC linear density (right) and thickness of the molecular layer (left) in the Cerebellar Lobule III of n=3-4 mice. Two-way ANOVA followed by Sidak's test. (C) Representative immunostaining of PCs 829 830 with anti-CALB1 antibody reveals progressive accumulation of vacuoles in KO mice. (D) Immunostaining of PCs with anti-831 CALB1 and lysosomes with anti-LAMP1 show enlarged lysosomes in KO mice. Bar graph shows average lysosome size per 832 mouse. n=3 mice (in WT = 29 PCs and 4,033 lysosomes were counted and in KO = 30 PCs and 3,247 lysosomes). Two-tailed t-test. (E-F) Representative immunostaining showing progressive accumulation of astrocytes labeled with anti-GFAP (E) and 833 834 microglia with anti-IBA1 (F) in degenerating KO cerebella (base of Lobule III & IV). (G) Coronal sections of cerebral cortices 835 immunostained with anti-NeuN do not show differences between WT and KO mice. Bar graphs show percentage thickness 836 occupied by each cortical layers (I-VI) in 4-5 cortical regions of 2 mice per genotype and age. Two-way ANOVA followed by Sidak's test. In all graphs, data represent mean  $\pm$ S.E.M. n.s. = non-significant, \*P < 0.05, \*\*P < 0.01, \*\*\*\*P < 0.001, \*\*\*\*P < 0.001837 838 0.0001.



Figure 4. Genes involved in lipid response are differentially expressed in SNX14 deficient cerebella. (A-B) Volcano plots of differentially expressed genes (DEGs) in the Cortex and Cerebellum of WT vs. KO mice at 1 month and 1 year. Dashed lines indicate statistical significance cut off  $(-\log 10(\text{Padj}) > 1.301 \text{ and } \log 2(\text{FC}) = \pm 0.5)$ . Number of significantly down and up -regulated genes are displayed on the top of each plot in blue and red, respectively. (C-D) Dot plots of gene ontology (GO) analysis of the DEGs, with down- and upregulated genes marked in blue and red, respectively. Dot size indicates proportion of DEGs relative to the total number of genes in each category. (E) Waterfall plots of Gene Set Enrichment Analysis (GSEA) of cerebellar specific significant gene ontology terms. Terms in orange, magenta, and black, are related to lipid, oxygen, and iron, respectively. (F) Heatmap of the top 20 leading edge genes of each term displayed in the 1-month Cerebellum GSEA (E). (G) Heatmap of the top 10 leading edge genes of each term displayed in the 1-year Cerebellum GSEA (E).





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Figure 5. Unique deregulation of lipid metabolites in pre-degenerating KO cerebella. (A) Volcano plots show deregulated lipids in 2-month-old Snx14 KO cerebellum (CB), cerebral cortex (CX), liver, and plasma. Horizontal gray lines indicate P <858 0.05 cut-off. Data shows increased concentrations of Acylcarnitine (AcCa) species specifically in KO CB. (B) Bar graphs show 859 total PE concentrations per tissue in n=8 WT and n=10 KO mice. Two-tailed t-test. PEs are significantly reduced in Snx14 KO 860 CX. (C) Dotplot depicting fold change (FC) (proportional to dot size) and p-value (in grey intensity scale) of PE species 861 detected in cerebral cortices for all analyzed tissues. Red dots represent significantly increased lipids while blue dots represent 862 significantly decreased lipids. (D) Bar graphs show total AcCa concentrations in n=8 WT and n=10 KO mice. Two-tailed t-863 test. AcCa-s are significantly increased only in KO CB. (E) Dotplot depicting FC and p-value of AcCa species detected in 864 cerebellar samples for all analyzed tissues. Red dots represent significantly increased lipids while blue dots represent 865 significantly decreased lipids. (F) MALDI-MS imaging of brain cryosections show reduction of PE C38:2, TG 46:1 and TG 866 53:2, and cerebellar accumulation of L-carnitine in KO. The molecules were revealed in positive ion mode using DHB matrix 867 and the m/z (mass-to-charge ratio) of  $[M+H]^+$  are indicated. Heatmap colors depict the relative abundance of each metabolic species. Bar graphs show cerebellar or cortical intensity of each lipid species in n=3 per genotype. In all panels, graphs show 868 mean  $\pm$ S.E.M. n.s. = non-significant, \*P <0.05, \*\*P < 0.01. Key of lipid class is found in supplemental data. 869



871 872 Figure 6. Lipid storage organelles are affected in SNX14 deficient tissue. (A) Representative BODIPY 493/503 (BD493) 873 labeling shows less lipid droplets (LDs) in 2-month-old KO mice liver sections. (B) Representative BD493 and anti-CALB1 874 labeling shows less LDs in Snx14 KO primary cerebellar culture PCs. Bar graphs show average number of LDs per CALB1<sup>+</sup> 875 PC in n=6 mice per genotype used for PC cultures. Total number of CALB1<sup>+</sup> PC quantified: n=69 WT and n=50 KO. Twotailed t-test. (C-D) Representative TEM image of PC layer in WT and KO mice at 2 (C) and 6 (D) months of age. (E) 876 877 Representative TEM images of PCs show less but larger telolysosomes in 2-month-old KO mice. Bottom graphs show the 878 average area of telolysosomes (left) and the percentage of PCs with indicated number of telolysosomes (right) in n=3 mice per 879 genotype (6-10 PCs per mouse). Two-tailed t-test (left) and Two-way ANOVA followed by Sidak's test (right). (F) 880 Representative TEM image of PCs showing less but larger telolysosomes in in 6-month-old KO mice. Bottom graphs show the average area of telolysosomes (left), and percentage of PCs with indicated number of telolysosomes (right) in n=3 mice per 881 882 genotype (6-10 PCs per mouse). Two-tailed t-test (left) and Two-way ANOVA followed by Sidak's test (right). (G) 883 Representative TEM image of PC mitochondria at 6 months of age. Bottom bar graphs show the average area (left) and 884 roundness (right) of mitochondria in n=3 mice per genotype (10 PCs per mouse). Two-tailed *t*-tests. (I) Representative TEM 885 images show a spectrum of less to more degenerating PCs from 6-month-old KO mice. Yellow arrowheads point to insets of 886 mitochondria and enlarged telolysosomes. ER swelling highlighted in magenta and indicated with an asterisk. In all panels, data represent mean  $\pm$ S.E.M. n.s. = non-significant, \**P* < 0.05, \*\**P* < 0.01, \*\*\**P* < 0.001, \*\*\*\**P* < 0.0001. 887