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# **Proteomic mapping of diferentially OPENvulnerable pre-synaptic populations identifes regulators of neuronal stability** *in vivo*

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**Synapses are an early pathological target in many neurodegenerative diseases ranging from wellknown adult onset conditions such as Alzheimer and Parkinson disease to neurodegenerative conditions of childhood such as spinal muscular atrophy (SMA) and neuronal ceroid lipofuscinosis (NCLs). However, the reasons why synapses are particularly vulnerable to such a broad range of neurodegeneration inducing stimuli remains unknown. To identify molecular modulators of synaptic stability and degeneration, we have used the** *Cln3***<sup>−</sup>/<sup>−</sup> mouse model of a juvenile form of NCL. We profled and compared the molecular composition of anatomically-distinct, diferentially-afected pre-synaptic populations from the** *Cln3***<sup>−</sup>/<sup>−</sup> mouse brain using proteomics followed by bioinformatic analyses. Identifed protein candidates were then tested using a** *Drosophila* **CLN3 model to study their ability to modify the CLN3-neurodegenerative phenotype** *in vivo***. We identifed diferential perturbations in a range of molecular cascades correlating with synaptic vulnerability, including valine catabolism and rho signalling pathways. Genetic and pharmacological targeting of key 'hub' proteins**  in such pathways was sufficient to modulate phenotypic presentation in a *Drosophila* CLN3 model. **We propose that such a workfow provides a target rich method for the identifcation of novel disease regulators which could be applicable to the study of other conditions where appropriate models exist.**

Synapses are an early pathological target in a range of diseases $^{1,2}$  $^{1,2}$  $^{1,2}$  including conditions associated with advanc-ing age (e.g. Alzheimer (AD)<sup>[3](#page-15-2),[4](#page-15-3)</sup> and Parkinson disease<sup>[5,](#page-15-4)[6](#page-15-5)</sup>), neurodevelopmental conditions (e.g. spinal muscular atrophy (SMA)<sup>7[,8](#page-15-7)</sup>), protein misfolding/accumulation diseases (e.g. Huntington disease (HD)<sup>[9](#page-15-8)</sup>), prion diseases<sup>10</sup>, spinocerebellar ataxias (SCA)<sup>[11](#page-15-10)</sup> and lysosomal storage disorders (Neuronal ceroid lipofuscinosis (NCLs or Batten disease)[12](#page-15-11)[–16.](#page-15-12) However, our understanding of the reasons why specifc synaptic populations are so vulnerable to such a broad range of neurodegenerative stimuli, and the mechanisms that govern their stability, remains in its infanc[y2](#page-15-1),[17](#page-15-13),[18](#page-15-14).

In this study, we sought to defne the molecular regulators of synaptic stability, using animal models of CLN3 disease (a.k.a. juvenile NCL or JNCL, OMIM # 204200). The NCLs, are the most frequent autosomal-recessive neurodegenerative disease and form of dementia in childhood[19.](#page-15-15) Incidence in the USA is estimated at 1.6–  $2.4/100,000$  whereas in Scandinavian countries it is  $2-7/100,000^{20,21}$  $2-7/100,000^{20,21}$  $2-7/100,000^{20,21}$ . The term "NCL" currently encompasses up to 14 disease subtypes, which are grouped together due the lysosomal accumulation of autofuorescent storage material, distinct ultrastructural properties, broadly similar pathology and clinical features and a severe

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neurodegenerative phenotype<sup>[16,](#page-15-12)22</sup>. Crucially for the purposes of this study, pre-synaptic disruption is a key early event in NCL, accurately predicting the distribution of subsequent neuronal loss $12-16$  $12-16$ .

The knowledge of the underlying genetic cause and/or storage material composition have provided a base for the basic understandying of the pathogenesis and their correlation to the clinical progression of the disease, the design of gene replacement therapies and the development of animal models<sup>[23](#page-15-19)–[25](#page-15-20)</sup>. However, the pathways and molecular cascades leading to neurodegeneration in NCL, as in many other neruodegenerative diseases, are still unknown.

Mutations in *CLN3* underlie a juvenile form of NCL, the most prevalent form worldwide<sup>26</sup>. The *CLN3* gene encodes a putative transmembrane protein whose function is not completely understood. However, experiments in yeast and *in vitro* studies have revealed possible functions relating to vacuolar pH regulation<sup>[27](#page-15-22)</sup> and endocytic membrane trafcking[28](#page-15-23). Although *CLN3* is ubiquitously expressed throughout the body, the most obviously afected tissues are neurologic based. Tis feature is shared by other monogenetic neurodegenerative conditions such as SMA<sup>[29](#page-15-24)</sup>. Yet, the reasons why neurons appear to be particularly vulnerable to defects in such broadly expressed proteins is not understood.

In humans, the onset of CLN3 disease occurs typically between 4–7 years of age, when loss of visual acuity is identified. Learning deficits are followed by speech and motor difficulties and seizures. As no treatment is currently available, premature death results at a mean age of 24 years old<sup>30</sup>. Unlike other more complex neurodegenerative diseases, the growing knowledge of the NCL-causative genes has facilitated the development of powerful animal models in recent years. Tese models have signifcatively improved our understanding of the progressive nature of the different forms of NCL. Cln3 null mice (*Cln3<sup>−/−</sup>*) reproduce various aspects of the human disor- $der<sup>31,32</sup>$  $der<sup>31,32</sup>$  $der<sup>31,32</sup>$  $der<sup>31,32</sup>$  $der<sup>31,32</sup>$ . Although neuronal loss is widespread in terminal disease<sup>31,[33](#page-16-3)</sup>, differential vulnerability can be detected across brain regions and their respective resident cell types/subcellular compartments at specifc time-points. Tis diferential degenerative progression follows the same pattern in the vast majority of NCL murine models: during pre-symptomatic stages there is an early selectivity for relay neurons within the thalamic nuclei followed by the corresponding cortical areas<sup>14[,31](#page-16-1)-35</sup>, as well as GABAergic hippocampal interneurons and Purkinje neurons<sup>[31,](#page-16-1)32</sup>. These reports in murine models regarding the vulnerability of the thalamus correlates with MRI studies in human patients showing alterations in thalamic areas at "pre-clinical" stages<sup>30[,36](#page-16-5),[37](#page-16-6)</sup>. Furthermore, differential regional/ neuronal vulnerability is also a shared event in a great number of neurodegenerative diseases in which some brain regions seem affected earlier than others, such as in Alzheimer<sup>[38](#page-16-7)–40</sup>, Parkinson<sup>41</sup>, Huntington diseases<sup>[42,](#page-16-10)43</sup>.

In this study, we aimed to defne the molecular regulators of both synaptic stability and vulnerability using animal models of NCL. We initially characterized diferential patterns of pre-synaptic pathology in *Cln3*<sup>−</sup>/<sup>−</sup> mice<sup>31</sup> across two different time-points. This enabled the subsequent application of high-throughput proteomics in order to map the molecular fngerprint of diferentially vulnerable biochemically-isolated pre-synaptic populations. We identify multiple cascades correlating with synaptic vulnerability and describe valine degradation and rho signalling pathways as two major regulators of synaptic vulnerability. The direct contribution of these pathways to neurodegeneration was confrmed *in vivo* using a *Drosophila* CLN3 model. Finally, we are confdent that regulators of synaptic vulnerability and degeneration identifed in the context of NCL are also conserved in their expression across other neurodegenerative conditions, in which pre-synaptic alterations are an early event<sup>[2](#page-15-1)</sup>. This research therefore opens a window for further investigation into common molecular therapeutic targets and strategies for novel interventions across a range of neurodegenerative conditions during early disease onset.

#### **Results**

**Quantitative immunohistochemistry identifes diferential rates of pre-synaptic pathology between brain regions in** *Cln3<sup>−/−</sup>* **mice.** Although the spatio-temporal pattern of neuron loss in *Cln3*<sup>−</sup> mice has already been studied<sup>32,33</sup>, little is known about the progression of synaptic pathology across differentially vulnerable brain regions. Studies in other NCLs using *Ppt1*<sup>−</sup>/<sup>−</sup>, *Cln6*<sup>−</sup>/<sup>−</sup> or *CathD*<sup>−</sup>/<sup>−</sup> mice have revealed progres-sive synaptic loss starting in the thalamus and followed later in corresponding cortical areas<sup>[12,](#page-15-11)14</sup>. Hippocampal structures seem to be also affected early, although to a lesser extent than seen in the thalamus<sup>[33](#page-16-3)</sup>. To investigate whether a similar pattern of synaptic pathology might be present in *Cln3*<sup>−</sup>/<sup>−</sup> mice we studied the expression of the presynaptic marker synaptophysin (Syp; as previously described in[12](#page-15-11)) in three brain regions that exhibit diferent degrees of neuronal vulnerability using quantitative immunohistochemistry (at 6.5 and 13 months; see methods – Fig. [1](#page-3-0) and Supplementary Fig. [S1\)](http://S1). The three brain regions studied were: 1. Thalamus – ventral posteromedial/ ventral posterolateral nuclei (VPM/VPL) and lateral geniculate nucleus (LGNd); 2. The corresponding cortical target regions in the primary somatosensory barrel feld (S1BF) and primary visual areas (V1) respectively, and 3; Hippocampus- stratum radiatum and stratum oriens (Fig. [1A](#page-3-0)). At 6.5 months no signifcant changes in Syp expression were detected between *Cln3*<sup>−</sup>/<sup>−</sup> and control mice in any of the brain regions studied (Supplementary Fig. [S1\)](http://S1). However, by 13 months, some diferences between genotypes were apparent (Fig. [1B and C](#page-3-0)). Similar to mouse models of other NCLs, thalamic nuclei demonstrated greater synaptic pathology at 13 months as indicated by the lower Syp immunoreactivity to synaptophysin in *Cln3*<sup>−</sup>/<sup>−</sup> vs. controls. Hippocampal structures were also afected, although as predicted, the diference in Syp immunoreactivity between genotypes in both hippocampal subfelds was less than in thalamic nuclei. In contrast, cortical regions such as S1BF and V1 (corresponding to the thalamic nuclei VPM/VPL and LGNd) showed no statistical diference between genotypes in Syp immunoreactivity at 13 months. Immunohistochemistry using antibodies against VAMP2 and SNAP-25 (two more pre-synaptic markers) also hint at pre-synaptic alterations in the thalamus (Fig. [S2](http://S2)–[S4\)](http://S4).

Tus, the pre-synaptic alterations seen here (as determined with the markers detailed above) appears to follow a similar pattern to that previously reported for neuronal loss in mouse models of other forms of NCL $^{12,14}$  $^{12,14}$  $^{12,14}$  $^{12,14}$  $^{12,14}$ , with the thalamus being the most afected region, followed by the hippocampus, whereas cortical measurements were still unafected at this stage of disease. We would categorize these alterations in pre-synaptic markers as being moderate at 13 months, as the pathology as quantifed here is less than 10% (9.57% relative to controls).



<span id="page-3-0"></span>**Figure 1.** Spatio-temporal synaptic loss study in *Cln3<sup>−</sup>/<sup>−</sup>* detected diferentially vulnerable synaptic populations across brain regions. (**A**) Brain region schematic showing the brain areas measured in grey. Talamic regions includes the ventral posterior medial/ventral posterior lateral thalamic nucleus (VPM/VPL) (top lef) and the dorsal lateral geniculate nucleus (LGNd) (bottom left); their respective cortical projections in the primary somatosensory cortex (S1BF) (top right) and primary visual cortex (V1) respectively (bottom right); hippocampal regions measured within the CA1-3 were the stratum radiatum and stratum oriens (bottom). (**B** and **C**) Representative photomicrographs of coronal sections of the same brain regions immunostained with synaptophysin (Syp) and bar chart showing its corresponding quantifcation based on the area of immunoreactivity in 13 month old control and *Cln3*<sup>−</sup>/<sup>−</sup> mice. Syp immunoreactivity was lower in thalamic regions (VPM/VPL and LGNd) in the *Cln3*<sup>−</sup>/<sup>−</sup> mice when compared to controls indicating more pathology, detectable earlier in the thalamus. Hippocampal stratum oriens and stratum radiatum also showed reduced Syp immunostaining, although the diference between genotypes was smaller. Cortical regions did not show diference in immunoreactivity for synaptophysin indicating that no synaptic loss is happening in these cortical areas at 13 months. (Mean  $\pm$  SEM; \*P < 0.05; \*\*P < 0.01; \*\*\*P < 0.001; ns P > 0.05, Student T test, 5 mice per each genotype and time-point were used, Scale bar = 200 um (left) and 20 um (right).

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<span id="page-4-0"></span>**Figure 2.** Experimental design workfow. Experiments are divided into *in vivo* (animal models), MS processing and *in-silico* where the different bioinformatics tools are detailed. \*This schematic was reproduced for the three brain regions of interest: cortex, hippocampus, and thalamus at 13 months of age.

**Comparative molecular profiling reveals proteomic perturbations that correlate with the extent of pre-synaptic alterations.** Despite advances in the characterization of the mammalian synaptic proteome, little is known about how these proteins interact and the molecular mechanisms that govern synaptic vulnerability. Afer confrming the synaptic vulnerability pattern across three distinct brain regions, we wanted to examine if the onset of pre-synaptic pathology correlated with distinct molecular alterations in the proteome. To address this, we examined biochemically isolated pre-synaptically enriched fractions (through the generation of "crude" synaptosomes – see methods) produced from microdissected brain regions from *Cln3*<sup>−</sup>/<sup>−</sup> mice and controls at 13 months of age where we characterise the pre-synaptic pathology as thalamus  $>$  hippocampus  $>$  cortex.

Afer synaptosome production and protein extraction, iTRAQ proteomics was carried out as detailed in Fig. [2.](#page-4-0) 1536 total proteins were identifed in hippocampus and thalamic samples whereas 2068 were detected in cortex. We then pre-fltered the proteins by those which were identifed by at least 2 peptides and therefore are more likely to be reliable identifcations. 914 proteins passed that flter in hippocampus and thalamus and 1295 in cortex. Almost 40% of our pre-fltered thalamic proteome was altered when compared to control littermates. In contrast, only 20% and 17.6% of the synaptic proteome was altered in the cortex and hippocampus relative to controls respectively. The number of protein alterations also correlates with pre-synaptic pathology being more apparent in thalamic areas at this "moderate" stage of disease progression.

Next, these pre-fltered proteins from the three pre-synaptic populations were aligned to look for overlapping proteins. 709 proteins were identifed in all three brain regions and these were taken forward for comparative analysis (Fig. [3A\)](#page-6-0).

In order to gain a broad and unbiased overview for the types of processes that may be represented by the protein alterations detected in all three pre-synaptic populations, we used *Ingenuity pathway analysis* (*IPA*) (see Methods – Fig. [3B](#page-6-0)). Interestingly, we identifed multiple canonical pathways which IPA designated as perturbed in a strikingly progressive manner, consistent with the degree of pre-synaptic pathology outlined above i.e. −log P value=higher in thalamus>hippocampus>cortex (Fig. [3B\)](#page-6-0). An example cascade can be found in more detail in Supplemetary Fig. [S5](http://S5). Tis analysis therefore indicates that in general, alterations in specifc molecular cascades were increasing with pre-synaptic pathology, further confrming the diferential vulnerability pattern previously identified by IHC. Thus, the synaptic vulnerability pattern described here appears consistent at both the morphological and that molecular level.

Whilst canonical cascade analysis is an interesting tool to use at the outset, multiple members of the same cascade do not necessarily need to change in the same manner, and whilst the trend of the majority of canonical cascades seen in Fig. [3B](#page-6-0) are consistent with the regional pathology reported, the individual protein alterations are far more complex (ass seen in the heatmap in Fig. [3C](#page-6-0)). Tus, to better understand and visualize the diferent molecular expression trends occurring across the three datasets, individual proteins were tracked using *BioLayout Express 3D* (see Methods). Simplistically, *BioLayout* is a complex pattern recognition sofware which generates a visual representation of the data based on protein abundance<sup>[44](#page-16-12)</sup>. In this case the software interrogates a regional expression profle generated by arranging the individual proteomic sets in order of increasing pre-synaptic pathology/vulnerability (cortex < hippocampus < thalamus). The outcome of this is that proteins were found to cluster into 14 diferent groups according to similarities in their relative expression across these three regions (Fig.  $3D$ )<sup>[45](#page-16-13)</sup>. Of interest to us were Clusters 3 (70 proteins), 4 (68 proteins), 6 (51 proteins) and 7 (50 proteins) comprise proteins whose expression can be grouped (or clustered) together as having either a gradual upregulation or downregulation, therefore correlating (directly or inversely with) with pre-synaptic vulnerability (Fig. [3D and E,](#page-6-0) Supplementary Table [S1](http://S1)). Next, we applied an enrichment analysis to these distinct clusters in order to determine if each expression trend could be associated with a specifc biological or functional category (using the *DAVID* software tool – see methods and Fig.  $3E$ ). Unexpectedly, clusters with proteins that were increasingly upregulated in vulnerable regions were enriched for structural associated candidates like cytoskeletal-related proteins, and specifc biological functions such as endocytosis (Cluster 4 and 7). Perhaps more surprisingly, clusters whose protein expression decreased with increasing vulnerability were related to specifc mitochondrial functions (Clusters 3 and 6) (Fig. [3D and E](#page-6-0) and Supplementary Table [S2\)](http://S2). Tus, comparative proteomic profling of diferentially vulnerable pre-synaptic populations revealed that signifcant alterations to the synaptic proteome correlated with the degree of pathology seen at these relatively moderate disease stages.

*In silico* **analysis highlights alterations in valine catabolic and ROCK2 signalling cascades in vulnerable pre-synaptic compartments.** Afer confrming that the pre-synaptic proteome changes in a manner consistent with the diferential vulnerability/regional disease progression in *Cln3*<sup>−</sup>/<sup>−</sup> mice, we next sought to tease out the potential molecular regulators of neuronal stability from our complex proteomic datasets. To do this, we focused our analyses on the thalamus, because pre-synaptic compartments here were perturbed to a greater extent than those in the hippocampus or cortex (according to the pre-synaptic markers we employed). In the thalamic dataset, 1536 total proteins were identifed following iTRAQ processing. Further *DAVID* enrichment analysis was applied to the unfltered thalamic data to confrm that the starting material was suitably enriched for synaptic proteins (Table [1](#page-7-0)). As described above, we applied fltering criteria based on the number of peptides (more than 2 unique peptides), those changed  $>$  20% (1.2 fold-change) and those which were mapped by *Ingenuity Pathway Analysis* (IPA) sofware. A total of 374 proteins met these criteria (Fig. [4A](#page-8-0) and Supplementary Tables [S3–](http://S3)[S4\)](http://S4). Next, we confrmed the veracity of the fltered data by validating a range of proteins using quantitative fuorescent Western blotting (QWB) (Fig. [4B–G\)](#page-8-0).

*Higher order functional clustering highlights similarities with other neurodegenerative conditions.* Here, we used *ingenuity pathway analysis* (*IPA*) to elucidate which molecular pathways and biological networks were disrupted in "afected" thalamic pre-synaptic compartments. Here we input the total pre-fltered thalamic data into IPA and applied the 1.2 fold-change cut-of in the sofware. Of the three-hundred and seventy-four proteins recognised by *IPA*, 68.3% have previously been reported in the literature as being associated with neurological disease (Fig. [5A](#page-9-0)). The majority of the diseases and disorders which fall under the category of "neurological disease" are known to demonstrate synaptic pathology as an early event<sup>[1](#page-15-0)</sup>. Interestingly, the main molecular and cellular functions identified in *IPA* were "molecular transport" and "cellular assembly and organization". These functional categories are consistent with the comparative analysis of diferentially vulnerable synaptic populations carried out above (see Fig. [3E](#page-6-0)), where clusters showing increases in protein expression from the levels detected in the comparatively spared cortical synapses through to greater levels again in the more vulnerable thalamic populations highlighted cytoskeletal protein alterations.

*Molecular cascade tracking identified potentially conserved regulators of vulnerability.* The "Canonical pathways" function in *IPA* revealed that, mitochondrial dysfunction was the most signifcant canonical pathway identifed (see Methods). The contribution of synaptic mitochondria in neurodegeneration has been highlighted in recent years<sup>46</sup>, and mitochondrial abnormalities have also been related to CLN3 disease and other NCLs<sup>[47](#page-16-15)[–50](#page-16-16)</sup>, and our data is therefore consistent with these previous fndings. However, in this study we wanted to consider pathways that have not previously been related to the NCLs before. Within the top canonical pathways identifed with *IPA*, we also detect perturbations in ROCK signalling and valine degradation pathways (Fig. [5B and C](#page-9-0)).



<span id="page-6-0"></span>**Figure 3.** Diferentially vulnerable synaptic population molecular profling. (**a**) Venn diagram and heat map show the 709 common proteins identifed and overlapped in cortical, hippocampus and thalamic synaptic proteomic datasets. (**B**) Heat map representing the signifcance (−log P value) of the canonical pathways identifed in IPA across cortex, hippocampus and thalamus datasets. It is observed a preogressive increase in signifcance correlating to the synaptic vulnerability pattern described previously. (**C**) Heat map showing the 709 common proteins identifed and overlapping in cortical, hippocampus and thalamic synaptic proteomic datasets. (**D**) *BioLayout* clustering 3D representation of proteomic expression data across diferentially vulnerable synaptic populations orientated at 3 diferent angles. Each sphere represents a single protein and the edge represents how similar their expression trend is towards the other proteins in the dataset. The closer the spheres are the more similar expression trend they have. The colours represent the different clusters of co-expressed proteins. (**E**) Expression profle means in log scale (*Cln3*<sup>−</sup>/<sup>−</sup>*/*WT) of co-expressed proteins in clusters 3, 4, 6 and 7 (Supplementary Tables [S1\)](http://S1) and its main biological function/subcellular compartment identifed by *DAVID* enrichment analysis (See Supplementary Table [S2\)](http://S2). Clusters highlighted show steady up or downregulation across cortex (C), hippocampus (H) and thalamic (T) regions correlating with the vulnerability status of synapses.

<span id="page-7-0"></span>

**Table 1.** *DAVID* analysis of proteomic data confrms synaptic protein enrichment.

Examining these pathways, proteins such as ROCK2 and HIBCH were diferentially expressed key "hub" components. Teir expression was identifed as tracking in *BioLayout* clusters 4 and 6 respectively correlating to the degree of pre-synaptic pathology in *Cln3*<sup>−</sup>/<sup>−</sup> mice (see above, Figs [3D](#page-6-0) and [5](#page-9-0)). Moreover, we have previously identifed ROCK2 and HIBCH expression alterations in pre-synaptic striatal isolates following cortico-striatal lesion injury (Fig. [5E](#page-9-0)), and demonstrated their potential to alter neurodegeneration in an injury specifc context *in vivo*[2](#page-15-1) . Collectively these fndings hinted that there may be a common molecular cascade underlying pre-synaptic pathology following a range of neurodegeneration inducing insults (from injury through to genetic). ROCK2 signaling and valine degradation pathways and specifcally proteins such as ROCK2 and HIBCH, could therefore represent potentially novel regulators of synaptic stability in an NCL disease context, which have not been investigated yet. So we next sought to determine if such candidates could have the ability to moderate NCL disease progression *in vivo*.

**Targeting the valine catabolic cascade is sufficient to modulate CLN3-induced neurodegener-<br><b>ation** *in vivo*. HIBCH is a 3-hydroxyisobutyryl-CoA hydrolase protein that fulfils a core function within the valine degradation cascade<sup>[51](#page-16-17)</sup>. Mutations in *HIBCH* cause a progressive infantile neurodegeneration in humans, characterized by hypotonia, motor delay and neurological regression $51-53$  $51-53$ . Moreover, manipulation of HIBCH has been suggested to modulate injury-induced axonal degeneration<sup>2</sup>. However, despite this apparently restricted function, very little is known about its role (if any) in synaptic compartments or its binding/interaction partners (see Fig.  $6A$ ).

Using QWB techniques we confrmed that HIBCH protein was less abundant in "crude" synaptosome extracts from thalamus in *Cln3*<sup>−</sup>/<sup>−</sup> mice relative to wild type controls. Interestingly, HIBCH protein levels trended towards a slight up-regulation of HIBCH in "non-synaptic" compartments. Whilst not statistically signifcant, this non-synaptic increase may indicate a redistribution or altered trafficking of this protein in response to *Cln3* defciency (Fig. [6B\)](#page-10-0). Importantly, examination of human *post mortem* brain samples showed that HIBCH protein levels were signifcantly reduced in JNCL patients (Fig. [6C\)](#page-10-0).

From comparing synaptic vulnerability (Fig. [3](#page-6-0)), HIBCH belonged to Cluster 6 (gradual downregulation profle) and was co-expressed with other mitochondria-related proteins (Fig. [3E\)](#page-6-0) including Hydroxyacyl-CoA Dehydrogenase/3-Ketoacyl-CoA Thiolase/Enoyl-CoA Hydratase (Trifunctional Protein), Alpha Subunit (HADHA), which also fulfls an enzymatic role within the valine degradation pathway (Fig. [6D\)](#page-10-0). HADHA sits upstream of HIBCH in the valine catabolism cascade (Fig. [6E](#page-10-0)) and catalyzes the fnal three steps of mitochondrial long chain fatty acid β-oxidation<sup>54</sup>. HADHA codes for the α subunit of the mitochondrial trifunctional protein (αMTP), and its defciency causes a metabolic disease that presents with Reye-like syndrome, with cardiomyopathy, neuromyopathy and sudden death in infancy<sup>55</sup>. More recent reports have indicated that HADHA deficiency results in a pigmentary retinopathy leading to vision loss<sup>56</sup>.

Given the important role of these two proteins in the valine catabolism cascade, we wanted to explore the consequences of experimental suppression of HIBCH and HADHA levels *in vivo* in order to assess their infuence on synaptic and neuronal stability in an NCL-disease specifc context. To do so, we used a *CLN3* gain of function fy model previously characterized by Tuxworth *et al*. in which the *Drosophila* eye undergoes degeneration (becoming smaller and rougher)<sup>57</sup>. Although this is an accepted model for investigations into the cascades underpinning CLN3, the human disease is normally caused by loss of function. As with the human disease where dose is crucial to phenotype (i.e. heterozygous humans are not obviously afected) we can also demonstrate that degeneration ocuring in the *Drosophila* eye is CLN3 dose dependent (Fig. [S6](http://S6)) as is the case for other neurodegeneration associated proteins (i.e.  $TDP43$ )<sup>58</sup>.

Tus, using this *Drosphila* model as a rapid screen for candidate ability to efect CLN3 driven phenotype (whilst bearing in mind its potential limitations) we frst established a recombinant fy containing the *GMR-GAL4* eye driver and the *UAS*-*CLN3* gene in the second chromosome (*GMR-GAL4;UAS-CLN3/CyOGFP*, see Table [2](#page-10-1)), and crossed this with fy lines containing RNA interference (RNAi) of HIBCH and HADHA mouse orthologs. CG5044 and CG4389/MTPα orthologs were selected using the DRSC Integrative Ortholog Prediction Tool (DIOPT) (see Methods and Table [2](#page-10-1)). Progeny of the relevant genotype was counted (*GMR-GAL4;UAS-CLN3/ HibchRNAi* and *GMR-GAL4;UAS-CLN3/MTPαRNAi*), eyes photographed and total surface area measured with



<span id="page-8-0"></span>**Figure 4.** Synaptic thalamic proteome fltering and validation. (**A**) Dot plot demonstrating the process of proteomic data fltering. Each data point represents an individual protein identifed using iTRAQ proteomic technique. LHS 1536 proteins were identifed across all thalamic samples. Following fltering (see Methods) a molecular fngerprint for thalamic synaptic alterations comprising 374 candidate proteins was produced (RHS; see also Supplementary Tables [S3](http://S3) and [S4\)](http://S4). (**B**) QWB representative bands for two upregulated (β-catenin and ROCK2), two downregulated (HIBCH and α-synuclein) and one unchanged protein candidate (calretinin) verifying the proteomic data. (**C** and **G**). Quantifcation and statistical QWB analysis showing the magnitude of alteration in *Cln3*<sup>−</sup>/<sup>−</sup> thalamic synaptic fractions. All the selected candidate proteins were altered as indicated by this iTRAQ analysis. Mean  $\pm$  SEM; \*P < 0.05; \*\*\*P < 0.001 (Student T test, n = 6 mice per each genotype).

*Image J* (see Methods). As expected, experimental suppression of either HIBCH or HADHA increased the neurodegenerative phenotypes observed (i.e. acting as enhancers of the CLN3-dependent phenotype) (Fig. [6F and](#page-10-0)  [G](#page-10-0)). Interestingly, the further upstream the valine degradation cascade is disrupted, the greater the efect on CLN3 phenotype (Fig. [6H](#page-10-0)). Interestingly, in this context neither HIBCH or HADHA were able to cause degeneration in the eye in the absence of the CLN3 disease background (Supplementary Fig. [S7](http://S7)). Tis suggests that their manipulation alone is not sufficient to affect general neuronal stability, and that the effect seen when crossing HIBCH and HADHA with the CLN3 fy is likely mediated by the *CLN3* allele. As such HIBCH and HADHA appear to be *in vivo* modifers of neuronal stability in a CLN3-disease context, and this provides further evidence that valine catabolism may play a role in CLN3-dependent neurodegeneration.

**Increased pre-synaptic ROCK2 expression appears conserved across a range of diseases and its downregulation can reduce CLN3-induced neurodegeneration** *in vivo***.** As discussed above, ROCK cascades were also identifed as being perturbed in degenerating thalamic pre-synaptic compartments and were predicted (by *in silico* analysis) to be activated when compared to wildtype controls (Fig. [5B and C\)](#page-9-0). ROCK2 is a *Rho*-kinase belonging to a family of serine/threonine kinases, of which isoform 2 is the most predominant in the brain<sup>59</sup>. ROCK2 directly interacts with, and is activated by, Rho GTPases. These are the central mediators of actin reorganization<sup>[59](#page-16-24)</sup>, which are reported to have a key role in synaptic plasticity and long term potentiation<sup>60</sup>. Activation of ROCK2 has been implicated in several adult-onset neurodegenerative conditions where synaptic pathology is present, such as HD, AD, ataxia, and in Purkinje cell degeneration<sup>2[,61–](#page-16-26)[64](#page-16-27)</sup>. Moreover, ROCK2 has been shown to infuence childhood neurodegenerative conditions such as SMA, where pharmacological inhibition partially recued symptoms and increases lifespan in an SMA mouse model<sup>[65,](#page-16-28)66</sup>. Moreover, we have previously reported that ROCK2 is capable of infuencing axonal degeneration *in vivo* following injur[y2](#page-15-1) . Taken together with our current data, this is consistent with a model whereby upregulation/activation of ROCK2 signalling in pre-synaptic compartments may correspond with increased neuronal vulnerability or degeneration in multiple infantile- and late-onset neurodegenerative conditions (Fig. [7D\)](#page-11-0).

The ROCK2 interactome (displayed in Fig. [7A\)](#page-11-0) illustrates its reported interactions with other gene/proteins, many of which were also altered in *Cln3*<sup>−</sup>/<sup>−</sup> thalamic synaptic extracts. In the *Cln3*<sup>−</sup>/<sup>−</sup> thalamus, QWB confrmed that ROCK2 was upregulated by 53.7% in the synaptic fractions, and by 27.7% in the corresponding "non







<span id="page-9-0"></span>**Figure 5.** IPA analysis of the thalamic synaptic proteome. (**A**) Table of top 5 "diseases and disorders" and "molecular and cellular functions". (**B** and **C**) Canonical pathways bar chart of representative pathways showing, (B) The percentage of upregulated and downregulated proteins within each pathway. Numbers on the top indicate the total number of proteins within the canonical pathway. (**C**) Signifcance of the association between the dataset and the canonical pathway (−log(p-value) and ratio) and z-score prediction of activation/inhibition (see Methods). Pathways highlighted in green and blue are associated to ROCK2 and HIBCH, respectively. (**D**) HIBCH and ROCK2 protein expression ratio across cortex ("less afected') and thalamic ("degenerating") synapses in these moderately afected *Cln3*<sup>−</sup>/<sup>−</sup> mice (*Cln3*<sup>−</sup>/<sup>−</sup>/WT). (**E**) HIBCH and ROCK2 protein expression is conserved in models of injury, following similar pattern of expression 24hours post-injury when synapses start to degenerate<sup>[2](#page-15-1)</sup>.



<span id="page-10-0"></span>**Figure 6.** CLN3 induced degeneration is enhanced by the genetic downregulation of HIBCH and HADHA *in vivo* impacting in valine catabolism pathway. (**A**) HIBCH proteome interactome represented by IPA indicating only 6 gene/proteins are known to interact with HIBCH. (**B**) QWB bands and quantifcation of HIBCH in synaptic and "non synaptic" fractions of *Cln3<sup>−</sup>/<sup>−</sup>* and control mice in thalamic protein extracts showing a downregulation in synapses and upregulation in non-synaptic parts \*\*P < 0.01, ns P > 0.05 (Student T test). (**C**) QWB bands and quantifcation of cortical "crude" synaptosomes isolated from *post mortem* human JNCL patients (P) and controls (C) showing its downregulation at end stages in human disease \*P<0.05 (Student T test). (**D**) HIBCH and HADHA are co-expressed and their expression trend correlates to the vulnerability status of synapses: unchanged in cortex ("spared") and downregulated in thalamus ("degenerating"). (**E**) Valine degradation pathway represented with *IPA*. The first protein which expression is perturbed in the degradation process of valine is HADHA; upstream of HIBCH. (**F–G**) Representative light microscope images of *Drosophila* eyes of disease model (DM), DM+HIBCH RNAi and DM+HADHA RNAi and corresponding quantifcation of the eye surface area. Scale bar=100um. \*\*\*P<0.001 (One-way ANOVA and Tukey's multiple comparison test as a post-hoc). (**H**) Schematic representation of the fourth and ffh steps of the valine catabolism cascade showing the enzymes HADHA and HIBCH catalyzing each step respectively. (**A,E & H**) Colored nodes illustrate proteins present in the thalamic dataset in *Cln3<sup>−</sup>/<sup>−</sup>* with respect WT. Nodes in grey represent proteins changed <20%, down-/up-regulated proteins by >20% are represented in green and red respectively. Orange box highlights the steps that are disrupted in the valine degradation cascade. Dotted lines on representative westerns indicate cropping of bands from the same membrane.

<span id="page-10-1"></span>

**Table 2.** *Drosophila* stock description. VDRC: Vienna *Drosophila* RNAi Centre. BDSC: Bloomington *Drosophila* Stock center.



<span id="page-11-0"></span>**Figure 7.** CLN3 induced degeneration is suppressed by the downregulation of ROCK2 genetically and pharmacologically (**A**) ROCK2 protein/gene interactome represented with IPA showing that some of the interactors of ROCK2 are also altered in "crude" thalamic synaptosomes. (**B**) QWB bands and quantifcation of control and *Cln3*<sup>−</sup>/<sup>−</sup> synaptic and "non synaptic" fraction. Upregulation of ROCK2 is higher in synapses than in the non-synaptic fraction indicating a potential synaptic specific response. \*P  $< 0.05$  (T-student test). (**C**) "Signaling by Rho family GTPases" canonical pathway represented by *IPA*. Upregulation/activation of ROCK2 impacts in actin nucleation and polymerization that may disrupt actin dynamics in the synapse. (**A&C**) Coloured nodes illustrate proteins present in the thalamic dataset in *Cln3*<sup>−</sup>/<sup>−</sup> with respect WT. Nodes in grey represent proteins changed <20%, down-/up-regulated proteins by >20% are represented in green and red respectively, orange indicates predicted activation and blue; predicted inhibition. (**D**) ROCK2 upregulation at early stage of disease is a converved event across animal models of injury, two adult-onset neurodegenerative diseases (Huntington and Spinocerebelar ataxia)<sup>2</sup> and two childhood neurodegenerative conditions (SMA and the lysosomal storage disorders-NCLs). (**E** and **H**) Representative light microscope images of *Drosophila* eyes and corresponding 200x zoom of eye structure of (**A**) Disease Model (DM) (**B**) DM+1.5mM Fasudil. (**C**) DM+Rok RNAi TRiP and (**D**) Control fy (Canton S). (**I**) Bar chart representing the % of normalized suppression calculated from the average eye surface areas of three independent experiments as "(x- average eye surface area of the disease model)/(average of eye surface area of the control - average eye surface area of disease model)  $\times$  100" for each of the three independent experiments. Scale bar = 100um. \*\*\*P < 0.001, \*P < 0.05 (One-way ANOVA and Tukey's multiple comparison test as a post-hoc).

synaptic" *Cln3<sup>−/−</sup>* isolates, relative to controls (Fig. [7B\)](#page-11-0), suggesting a more synaptically-focussed response to altered CLN3 expression. Our *in silico* analysis added support for this model by highlighting specifc sub-cascades under multiple pathways related to ROCK2 such as "Signalling by Rho Family GTPases" and "Rho A signalling" (Fig. [5B and C](#page-9-0)). Tese cascades regulate actin dynamics and cytoskeletal organization (as shown in Fig. [7C](#page-11-0)) and can impact synaptic structure and afect synaptic transmission. Figure [7D](#page-11-0) summarises previously published data generated by our laboratory where ROCK2 was identifed to be upregulated in a range of conditions at early stages of disease progression<sup>[2](#page-15-1)</sup> and new synaptic data regarding its role in CLN3 disease. Thus, the conserved increased expression of ROCK2 in *Cln3*<sup>−</sup>/<sup>−</sup> mice, and in other related conditions and its ability to change degeneration, is highly suggestive of a potentially conserved contribution to synaptic vulnerability.

Thus, in order to investigate whether ROCK2 is capable of directly modulating the process of neurodegeneration in a CLN3-specifc disease context *in vivo*, we again turned to the CLN3 *Drosophila* model. RNAi GD and RNAi TRiP fy lines for a ROCK2 ortholog were crossed to our established CLN3 recombinant *Drosophila* (see Table [2](#page-10-1)). Genetic downregulation of ROCK2 with two independently generated RNAi lines were tested and both were found to drastically improve their CLN3*-*induced small eye phenotype, although was more obvious with the TRiP library line (Fig. [7G–I\)](#page-11-0). The total surface area of the eyes of the offspring (*GMR-GAL4*;*UAS-CLN3/*  *RokRNAi*) were non-signifcant with respect the control eyes and therefore the phenotype was fully-recued. Qualitatively, there was also an obvious reduction/absence of black patches (indicative of cell death), and clear recovery of the gross structure of the *Drosophila* compound eye. With the RNAi GD line the eye size was also improved (25.22% of recue) (Fig. [7I](#page-11-0)). Thus, experimental manipulation of ROCK2 was sufficient to modulate neurodegeneration in a CLN3 model *in vivo*. To exclude the possibility of the suppression efect being due to the dilution of *GAL4* among two UAS transgenes, a control experiment was carried out, where the CLN3 recombinant fy was crossed to a UAS-*GFP* line. No signifcant changes were detected in the eye of the ofspring, indicating that the results showed in our RNAi experiments are likely a consequence of the downregulation of ROCK2.

#### **The ROCK inhibitor Fasudil ameliorates CLN3-dependent neurodegeneration** *in vivo***.** Following the rescue of the CLN3-dependent eye degeneration by means of genetic tools we therefore wanted to test the possibility of targeting ROCK2 pharmacologically in *Drosophila*.

Several ROCK inhibitors have been characterized, but their specifcity for individual ROCK kinases is still unclear. Fasudil is a small molecule that unspecifcally inhibits ROCK2, originally discovered as a blocker of cerebral vasospasm in animal models<sup>67</sup>. Importantly, fasudil has been demonstrated to be safe and effective in clinical trials of cerebral vasospasm, pulmonary hypertension, Raynaud phenomenon and cancer<sup>[68](#page-16-31)-[73](#page-17-0)</sup>. Moreover, fasudil has previously been applied to murine models of SMA showing an improvement of the phenotype and lifespan of these mice<sup>65,66</sup>. However, fasudil also targets other kinases such as MAP4K4, PKC, PRKAA1 or PRKAA2<sup>74</sup>, which are predicted to be also activated by IPA (Supplementary Fig. [S8A\)](http://S8A). The activation of these kinases seem to have downstream consequences on biological functions relevant to NCL such as autophagy (Supplementary Fig. [S8A\)](http://S8A) or and "Protein kinase A signalling" (Supplementary Fig. [S8B and C\)](http://S8B and C). Given the success of the compound in mammalian systems<sup>65,[66,](#page-16-29)73</sup>, including humans<sup>[69](#page-16-32),[70](#page-16-33)</sup>, to date and the apparently conserved ROCK2 alterations across multiple-neurodegenerative paradigms, we decided to test Fasudil in CLN3 *Drosophila* (*GMR-GAL4/UAS-CLN3*). The drug was added into the fly food in order to obtain 1.5 mM of fasudil in the food and the *Drosophila* were raised as normal (see Methods). Pharmacological inhibition with fasudil caused a signifcant, albeit modest suppression of the CLN3*-*derived eye phenotype. Quantifcation of eye surface areas confrmed a 17% rescue of the CLN3 eye phenotype in the treated animals (Fig. [7E,F and I](#page-11-0)). Tus, targeting of ROCK cascades with fasudil was sufficient to ameliorate neurodegeneration in a CLN3 model *in vivo*.

#### **Discussion**

Here we show what may be the frst proteomic profling carried out across multiple diferentially vulnerable synaptic populations at an early stage of disease progression. We have demonstrated that such unbiased proteomic mapping of distinct pre-synaptic populations coupled with *in silico* analysis, and *in vivo* rapid phenotypic screening in *Drosophila*, is an efective target-rich workfow for the identifcation of novel molecular alterations that regulate synaptic/neuronal stability.

Here we have confirmed that *Cln3<sup>−/−</sup>* murine pre-synaptic pathology appears to follow the same neuronal diferential vulnerability pattern as other NCL subtypes[12](#page-15-11),[14](#page-15-25). We show that, at 13 months of age in *Cln3*<sup>−</sup>/<sup>−</sup> mice the synaptic pathology is more pronounced in the thalamus (although still <10%), followed by hippocampal pre-synaptic compartments, while cortical synaptic populations remain relatively unafected (Fig. [1](#page-3-0)). iTRAQ proteomic and bioinformatics showed that diferential synaptic pathology was also refected at the molecular level, showing a progressive increase in the disruption of multiple canonical pathways correlating with our obser-vations at the immunohistological level (Fig. [3B](#page-6-0)). Further analyses by means of complex pattern recognition software identifed alterations in the pre-synaptic proteome (identifed by iTRAQ analysis of "crude" synaptosomes) that correlated with the assigned vulnerability status of pre-synaptic populations in the *Cln3*<sup>−</sup>/<sup>−</sup> mice. Pathway analysis highlighted perturbations in valine catabolism and *rho* signaling pathways, with proteins including HIBCH, HADHA and ROCK2 as key players, correlating with synaptic vulnerability. Furthermore, HIBCH and ROCK2 displayed a conserved direction of change in multiple neurodegenerative conditions (Figs [5E](#page-9-0) and [7D](#page-11-0)), and we have previously demonstrated that they can alter axonal degeneration in an injury dependent context<sup>[2](#page-15-1)</sup>. Importantly, this suggests that not only do conserved pathways regulating degenerative processes exist, but they may be targetable across multiple conditions (Fig. [7D](#page-11-0)).

In contrast, no lysosomal specifc pathways were identifed in our *in silico* analysis in synapses, indicating that loss of CLN3 may be impacting in other pathways beyond lysosomal function. Tis is supported by studies in which CLN3 was found to be expressed in other locations such as in synaptosomes, lipid rafs, Golgi, mito-chondria, glia and endothelial cells<sup>[75–](#page-17-2)[79](#page-17-3)</sup>, where it may play other roles such as regulation of oxidative stress<sup>[78,](#page-17-4)[80](#page-17-5)</sup> and glial function<sup>32[,34](#page-16-34)</sup> or blood-brain barrier homeostasis<sup>[79](#page-17-3)</sup>. However, individual NCL and lysosomal-storage disorder-related proteins were identified, such as cathepsin  $D^{81}$ , prosaposin $82,83$  $82,83$  $82,83$  and acid ceramidase $84$  confirming the veractity of the data generated.

By infuencing the valine catabolic cascade through genetic disruption of HIBCH and HADHA it was possible to alter the CLN3 degenerative phenotype in *Drosophila in vivo*. Moreover, the further upstream in the catabolic cascade we intervened, the greater the resulting efect on disease phenotype. In humans, both HIBCH and HADHA deficiency cause infantile onset diseases with neurologic clinical scheme<sup>53,[85,](#page-17-10)[86](#page-17-11)</sup>. Crucially, this could indicate that perturbations in valine catabolic pathways might be a shared event across multiple neurodegenerative conditions and therefore targets identifed here may be transferable to other diseases.

Interestingly, the upregulation of ROCK2 has been reported in a range of conditions by our laboratory and other[s2,](#page-15-1)[61,](#page-16-26)[62,](#page-16-35)[64](#page-16-27) (Fig. [7D](#page-11-0)). ROCK2 upregulation seemed to be a more synaptic-specifc perturbation (at least at early stages) in the *Cln3*<sup>−</sup>/<sup>−</sup> mice and its magnitude of change tracked across diferentially vulnerable brain regions (i.e. Talamus>Hippocampus>Cortex). Its genetic and/or pharmacological downregulation has also been shown to ameliorate the phenotype of several conditions including SMA, another relatively early onset neurodegenerative disease[65](#page-16-28),[66](#page-16-29). In this study we demonstrate for the frst time that downregulation of ROCK2 also recues the

phenotype in a CLN3 *in vivo* model when manipulated genetically, and to a lesser degree with available pharmacological agents (Fig. [7E–I\)](#page-11-0).

Cortical synaptic extracts from human patients have shown that ROCK2 was downregulated in JNCL patients with respect to controls (Supplementary Fig. [S9](http://S9)). Nevertheless, the interpretation of the results from *post mortem* tissue is challenging as at end stage of the disease brain atrophy is much more pronounced, making it difficult to distinguish which molecular alterations are more likely to be causative, and which are a consequence of the ongoing degeneration taking place. ROCK2 upregulation could be a disease stage-specifc change that contributes to the initiation of degenerative stimuli, and it is not necessarily globally upregulated throughout disease progression. It would be advantageous to study the molecular alterations and pathways taking place throughout the time-course of disease progression in NCL as this would further our understanding of the mechanisms underlying the initiation and progression of neurodegeneration. In general, ROCK2 appears to be an attractive target for the development of new therapeutic strategies, not only for NCL, but also in a wide range of more common neurodegenerative diseases to potentially halt or delay disease progression (Fig. [7D\)](#page-11-0). There are currently multiple ROCK inhibitors with different degrees of specificity available<sup>68,74</sup>. However, in order to avoid off-target effects in future studies, it will be necessary to develop a compound that targets ROCK2 more specifcally.

It would be of interest for future investigations to study the molecular overlaps between valine degradation and ROCK2 signaling pathways contributing to synaptic vulnerability. *In silico* analysis based on published literature, suggests potential overlaps, although the level of complexity in the published data allowing us to draw links between these candidates would require carefully planned experiments in order to unravel such interactions (See Fig. [S10\)](http://S10).

We suggest that common mechanisms regulating the process of neurodegeneration and/or vulnerability of synapses may occur across a range of neurodegenerative conditions triggered by diferent insults. Unlike more common adult-onset complex polygenetic diseases such as Alzheimer or Parkinson disease for which the genetic insult triggering the neurodegeneration is not well understood, the NCLs are monogenetic diseases for which there are available well-characterized mice models that more accurately replicate the human disease<sup>14,[31,](#page-16-1)[32,](#page-16-2)[35,](#page-16-4)[87,](#page-17-12)88</sup>. Here we demonstrate that we can use the NCLs as a model to study a specifc neurodegenerative process (i.e. synaptic loss) which is likely goverened by mechanisms which may indeed be conserved across multiple conditions ranging from injury through to chronic neurodegenerative conditions (i.e. Huntington disease and spinocerebellar ataxia) and diseases of childhood (i.e. SMA and the NCLs).

The workflows highlighted here should not be viewed as a mass screening "stamp collecting" exercise. Instead it should be clear that combining mammalian model "–omic" screening of diferentially vulnerable tissues with *in silico* candidate identification and *in vivo* phenotypic assessment in *Drosophila* is an efficient pipeline for elucidating the mechanistic cascades governing neurodegenerative processes, and a "target-rich" way of identifying factors which are capable of modulating neurodegeneration, and therefore may be therapeutically targetable.

#### **Materials and Methods**

**Tissue harvesting and processing.** *Ethics statement*. All animal experiments were approved by a University of Edinburgh internal ethics committee and were performed under license by the UK Home Office (project license numbers 70/6567 and 70/7364).

*Mice. Cln3<sup>−/−</sup>* mice<sup>[31](#page-16-1)</sup> inbred on a C57BL/6 background for at least 10 generations and control (+/+) littermates resulting from heterozygous crosses were used in this study. Mice were genotyped as described i[n31.](#page-16-1) Five *Cln3*<sup>−</sup>/<sup>−</sup> and fve wild-type control mice (WT) at 6.5 months and 12–13 months of age were used for immunohistochemistry experiment. Four *Cln3*<sup>−</sup>/<sup>−</sup> and four control mice (WT) at 12–13 months were used for the proteomics and biochemical experiments.

*Human post mortem samples.* Human brain samples were obtained from the Human Brain and Spinal Fluid Resource Center, Los Angeles and from The MRC London Neurodegenerative Disease Brain Bank, Institute of Psychiatry, Psychology & Neuroscience, King's College London. Samples were obtained at routine autopsy with fully informed written consent by the families. Study protocols for the use of this human material were approved by the Ethical Research Committees of the Institute of Psychiatry under the approval numbers 223/00 and 181/02. All experiments using human tissue samples and/or derivatives of, were performed in accordance with the relevant guidelines and regulations governing their storage, handling and disposal. Details about the samples are described in Supplementary Table [S5](http://S5).

**Immunohistochemistry (IHC).** Five *Cln3*<sup>−</sup>/<sup>−</sup> and fve wild-type control mice (WT) at 6.5 months and 12-13 months of age were terminated and brains rapidly removed and immersed for fxation in 4% paraformaldhyde (pH 7.4). The IHC protocol used allows quantitative and qualitative comparisons between animal tissues as previously described in<sup>[89](#page-17-14),[90](#page-17-15)</sup>. Please refer to Supplemental Experimental Procedures for more details.

**Biochemical separation of neuronal compartments.** Brains were harvested from six *Cln3*<sup>−</sup>/<sup>−</sup> and wild-type (WT) controls mice at 13 months old and briefy chilled in ice cold ACSF (125 mM NaCl, 26 mM NaHCO3, 25mM glucose, 2.5mM KCl, 1.25mM NaH2PO4, 1mM CaCl2, 4mM MgCl2) before regional microdissection. Talamus, cortex (predominantly pre-/frontal) and hippocampus and were microdissected bilat-erally<sup>12[,17](#page-15-13),91</sup> pooled by mouse and immediately processed for "crude" synaptosome production as previously described<sup>2,[17](#page-15-13),[92](#page-17-17)</sup>. Please refer to Supplemental Experimental Procedures for more details.

For the *post mortem* human samples, a portion of the total sample was obtained from frozen tissue (stored at −80 °C) and transferred in an eppendorf tube containing cold isotonic sucrose solution. Please refer to Supplemental Experimental Procedures for more details.

**Protein extraction.** *Mice*. For iTRAQ proteomics, protein extraction from "crude" synaptosomes was carried out as previously described in Fuller *et al.*[93](#page-17-18). Please refer to Supplemental Experimental Procedures for more details.

*Human.* "crude" synaptosomes were resuspended and homogenized in a solution containing 100mM Tris-HCl, 4% (w/v) SDS and 1% protease inhibitor cocktail (Roche). Samples were spun at 20,000g and 4 °C for 20minutes. Supernatant was aspirated and collected as extracted sample. Pellets and extracted samples were stored at −80 °C for QWB. Extracted samples concentration was determined using a BCA assay (Thermo).

**Proteomic processing.** Sample preparation and protein identifcation and quantifcation analysis by mass spectrometry was carried out as previously described in Fuller *et al.*[93.](#page-17-18) Please refer to Supplemental Experimental Procedures for more details.

**Quantitative fuorescent western blotting (QWB).** QWB was performed as previously described in refs[2,](#page-15-1)[91,](#page-17-16)[92,](#page-17-17)[94](#page-17-19)[,95.](#page-17-20) Briefy, samples were denatured in NuPage® LDS Sample bufer 4X (Invitrogen, UK) at 98 °C and 15ug of protein loaded and run on 4–12% Bis-Tris gel (Invitrogen). Accuracy of loading and protein estimation was confirmed by total protein analysis of Instant Blue (Expedeon) stained gels as previously described<sup>94</sup>. Protein transfer to a polyvinylidene fluoride (PDVF) membrane was carried out using the I-Blot® transfer system (Invitrogen, UK). Membranes were incubated with Odyssey blocking bufer (Li-Cor) for 30minutes. Next, membranes were incubated in primary antibodies overnight at 4 °C and secondary antibodies for 1 h at room temperature. Please refer to Supplemental Experimental Procedures for more details.

*In silico* **proteomic analysis.** *BioLayout Express 3D*. BioLayout sofware incorporates a complex pattern recognition algorithm which groups protein data based only on expression profle. It allows the visualization and graphing of expression trends of co-expressed proteins to allow better characterization and understanding of complex large datasets<sup>[44](#page-16-12)</sup>. Please refer to Supplemental Experimental Procedures for more details.

*Enrichment analysis.* To obtain an indication of the level of sample enrichment aforded through the process of synaptosome production, un-filtered mass spectrometry data was processed using The *Database for Annotation*, *Visualization and Integrated Discovery* (*DAVID*) sofware (available at [http://david.abcc.ncifcrf.gov\)](http://david.abcc.ncifcrf.gov). *DAVID* provides a relatively comprehensive set of functional annotation tools for large data set interpretation<sup>96,97</sup>. *DAVID* was also used to characterize the functions associated to the protein clusters correlating to vulnerability mapped using *BioLayout*. For analysis to confirm synaptic protein enrichment we have applied a cut off of four fold as previously described in $98$  (see Table [1](#page-7-0)).

*Ingenuity Pathway analysis.* To obtain further insight into potential cellular pathways that may be perturbed in the *Cln3*<sup>−</sup>/<sup>−</sup> thalamic pre-synaptic compartments compared to control mice, the *Ingenuity Pathways Analysis* (*IPA*) application (Ingenuity Systems) was used, as previously described[2,](#page-15-1)[91](#page-17-16)[,92](#page-17-17),[99](#page-17-24). Please refer to Supplemental Experimental Procedures for more details.

**Phenotypic assessment methods and** *Drosophila* **husbandry.** *Drosophila* mouse orthologs for candidate proteins were identifed using *DRSC Integrative Ortholog Prediction Tool* (*DIOPT*)[100](#page-17-25) and RNAi lines were chosen from Flybase and obtained from Vienna *Drosophila* stock centre (VDSC) and from Bloomington *Drosophila* Stock Centre (BDSC) (see Table [2](#page-10-1)). UAS-*CLN3* mutant fy was donated by Richard Tuxwort[h57](#page-16-22). A stock carrying *GMR-GAL4* and UAS-*CLN3* on the second chromosome was established by conventional recombination methods and used as a tester line for the screen (*GMR-GAL4;UAS-CLN3/CyO GFP*). RNAi candidate lines were crossed to this line and the F1 progeny was assessed for suppression or enhancement of the CLN3-derived small and rough eye phenotype. To have a quantitative read out of the severity of the phenotypes obtained in our experiments we measured the eye surface area, as previously described<sup>[101](#page-17-26)</sup>. Stocks were maintained on standard cornmeal food at room temperature. For all the crosses, fies were raised at 25 **°**C in a circadiarian light incubator. Details of the lines used can be found in Table [2.](#page-10-1)

**Fasudil drug assay.** The rock inhibitor used was Fasudil, Monohydrochloride Salt (LC laboratories, *F-4660*), which was previously used in mice models of SMA improving survival and skeletal muscle developmen[t65.](#page-16-28) *GMR-GAL4* x UAS-*CLN3* crosses were raised in Nutri-Fly instant food (Genesee Scientifc, *66-118*) containing 1.5 mM of Fasudil Monohydrochloride Salt and blue food colouring diluted in water, as described in<sup>102</sup>. Eyes of the ofspring were photographed and total surface area was measured with *Image J*. Suppression of the CLN3-degenerative eye were calculated from 3 independent experiments as "(x- average eye surface area of the disease model)/(average of eye surface area of the control−average eye surface area of disease model)×100" for each experiment.

**Imaging.** *Immunohistochemistry*. Micrographs were taken with Leica DMRB × 5/0.12 objective and AxioCam HRC Zeiss from the following brain regions: 1. The thalamic nuclei ventral posterior medial/ventral posterior lateral thalamic nucleus (VPM/VPL) and the dorsal lateral geniculate nucleus (LGNd); 2. Teir respective cortical projections in the primary somatosensory cortex (S1BF) and primary visual cortex (V1) respectively, and 3; the hippocampal stratum radiatum and stratum oriens (See Fig. [1A\)](#page-3-0). Immunoreactivity measurements were carried out using a thresholding image analysis in *Image J* (National Institute of Health, Bethesda, MD), USA), as previously described  $32,34,103$  $32,34,103$  $32,34,103$  $32,34,103$ . Thirty non-overlapping images (x40) were taken, on three consecutive sections, through the brain areas described above. All parameters including lamp intensity, video camera setup and calibration were kept constant throughout image capturing.

*Drosophila eye images. For the HIBCH and HADHA experiments* fruit fy eye images were photographed with a Nikon D5100 camera attached to a SZX9 Nikon stereomicroscope. For the ROCK2 experiments eye images were taken with an AxioCam ERc 5s Rev.2 attached to a Zeiss Stemi 305 trino stereomicroscope. Images were analyzed with *Image J* software (National Institute of Health, Bethesda, MD, USA) as previously described in<sup>101</sup>.

**Data analysis and fgure production.** QWB data was analysed using *Odyssey* sofware as per manufac-tures guidelines and as previously described<sup>[2,](#page-15-1)[17,](#page-15-13)[104](#page-17-29)</sup>. Data was graphed and statistical comparisons for IHC thresholding, QWB protein intensity and *Drosophila* eye surface area quantifcation were carried out using *GraphPad Prizm* as previously described<sup>[91](#page-17-16),[101](#page-17-26)</sup>. For QWB and IHC Student's T test was applied. For *Drosophila* eye quantifications one-way ANOVA was carried out followed by Tukey's multiple comparison test as a post-hoc when a significant difference was found in the ANOVA. P values < 0.05 were considered to be statistically significant for all analyses (\*P<0.05; \*\*P<0.01; \*\*\*P<0.001). Statistical analysis of proteomic data was automatically carried out by *MASCOT*. P & Benjamini values to determine enrichment were automatically calculated by *DAVID* sofware (see above and [http://david.abcc.ncifcrf.gov\)](http://david.abcc.ncifcrf.gov). P values/Fishers exact tests for pathway and networking analysis were automatically determined by *IPA* (see above and<http://www.ingenuity.com/>). Heatmap representing the differentially expressed proteins and pathways across brain regions was produced using "R" [\(http://www.R-project.](http://www.R-project.org)  $\text{org})^{105}$  $\text{org})^{105}$  $\text{org})^{105}$ .

**Availability of data and material.** The raw datasets used and/or analysed during the current study are available from the corresponding author on reasonable request.

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#### **Author Contributions**

T.M.W. conceived and designed the study. M.L.H., H.R.F., A.M.S.W., S.L.E. and T.M.W. carried out experiments. T.M.W., and J.D.C. contributed reagents. M.L.H., J.D.C., T.H.G. and T.M.W. wrote the manuscript. G.P. contributed to the interpretation of the fy data. All authors contributed to the data analysis, interpretation and approved the fnal manuscript.

#### **Additional Information**

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**Competing Interests:** The authors declare that they have no competing interests.

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