Quantitative Peptidomics of Purkinje Cell Degeneration Mice

Iryna Berezniuk1, Juan J. Sironi2, Jonathan Wardman2, Raymond C. Pasek3, Nicolas F. Berbari3, Bradley K. Yoder3, Lloyd D. Fricker1,2

1 Department of Neuroscience, Albert Einstein College of Medicine, Bronx, New York, United States of America, 2 Department of Molecular Pharmacology, Albert Einstein College of Medicine, Bronx, New York, United States of America, 3 Department of Cell, Development, and Integrative Biology, University of Alabama at Birmingham Medical School, Birmingham, Alabama, United States of America

Abstract

Cytosolic carboxypeptidase 1 (CCP1) is a metallopeptidase that removes C-terminal and side-chain glutamates from tubulin. The Purkinje cell degeneration (pcd) mouse lacks CCP1 due to a mutation. Previously, elevated levels of peptides derived from cytosolic and mitochondrial proteins were found in adult pcd mouse brain, raising the possibility that CCP1 functions in the degradation of intracellular peptides. To test this hypothesis, we used a quantitative peptidomics technique to compare peptide levels in wild-type and pcd mice, examining adult heart, spleen, and brain, and presymptomatic 3-week-old amygdala and cerebellum. Contrary to adult mouse brain, young pcd brain and adult heart and spleen did not show a large increase in levels of intracellular peptides. Unexpectedly, levels of peptides derived from secretory pathway proteins were altered in adult pcd mouse brain. The pattern of changes for the intracellular and secretory pathway peptides in pcd mice was generally similar to the pattern observed in mice lacking primary cilia. Collectively, these results suggest that intracellular peptide accumulation in adult pcd mouse brain is a secondary effect and is not due to a role of CCP1 in peptide turnover.

Introduction

In the 1970s, a spontaneous mutant mouse was discovered and named Purkinje cell degeneration (pcd) due to the loss of cerebellar Purkinje cells starting around 3 weeks after birth [1]. A small number of other cell types undergo degeneration in pcd mice, including olfactory bulb mitral cells, retinal photoreceptor cells, and spermatocytes [1]. The mutation responsible for the pcd phenotype was mapped to the gene encoding cytosolic carboxypeptidase 1 (CCP1, also known as Nna1), and the gene was named Agtpbp1 because the protein was initially considered to be an ATP/GTP binding protein [2]. CCP1 was discovered in a search for mRNAs upregulated in spinal motor neurons during regeneration after axotomy [3]. Thus, CCP1 is linked to both degeneration and regeneration. CCP1 has sequence homology to metallocarboxypeptidases, including conservation of critical active site residues, but lacks a signal peptide and is expressed in the cytosol [4]. Five additional members of the CCP1 subfamily were discovered and named CCP2 through CCP6 [4,5]. CCP1 is the most abundant of the CCPs in mouse brain [4].

Altogether, nine independent phenotypic alleles of pcd have been characterized that are due to mutations in the Agtpbp1 gene [6–8]. The pcd3 allele results from a deletion of exons 6–8, and the splicing of exon 5 to exon 9 introduces an inframe stop codon that results in a truncated protein lacking the active CP domain. The CP domain is critical for CCP1 function and normal mouse phenotype; expression of CCP1, but not CCP1 with mutated catalytic residues, in Purkinje cells of pcd mice prevents loss of these cells [9,10]. Many studies have been done to characterize Purkinje cell death in mutant mice, and a number of potential mechanisms of neurodegeneration have been proposed [8,11–15].

Based on the broad distribution of CCP1 mRNA in many tissues and cell types, and the expression of the protein in the cytosol, two potential functions for CCP1 were proposed. One is a role in protein/peptide turnover within a cell. Proteins are degraded by the proteasome into peptides, which need to be converted into amino acids and subsequently be recycled into proteins. Although evidence suggests that peptide degradation is largely carried out by aminopeptidases [16,17], in theory it is possible that cytosolic carboxypeptidases also contribute to this process. Previously, some of the authors of the present study reported that adult pcd mouse brains have greatly elevated levels of peptides derived from intracellular proteins (i.e. those present in cytosol, mitochondria, and other non-secretory pathway compartments) [18]. This finding was interpreted as evidence that CCP1 participates in intracellular peptide degradation [18]. The other function proposed for CCP1 involved tubulin processing [4]. The alpha chain of tubulin undergoes trimming of the C-terminal Tyr and Glu residues. In addition, both alpha and beta tubulin are modified by addition of Glu or Gly to the gamma carboxyl group of a Glu located near the C-terminus, and these side chains are subsequently removed in a dynamic
process [19–21]. Recently it was shown that CCP1 is capable of removing Glu residues from the C-terminus and polyglutamyl side chains of tubulin [22,23]. Furthermore, the lack of CCP1 leads to tubulin hyperglutamylation. Knock-down or knock-out of tubulin tyrosine ligase-like-1, an enzyme that adds Glu to the side chain of tubulin, prevents neurodegeneration of Purkinje cells in pcd mice [22,23]. Impaired tubulin polyglutamylation in other animal models is known to lead to mislocalization of molecular motors and affect tubulin-dependent trafficking and synaptic transmission [24–26]. To test if CCP1 functions in the degradation of intracellular peptides, we compared peptide levels in brains of presymptomatic pcd mice, before Purkinje cell death, to age-matched wild-type (WT) mice. We also measured relative peptide levels in non-neuronal tissues of adult pcd mice, selecting two organs (heart, spleen) which have high levels of CCP1 relative to other CCPs. Because protein turnover is a fundamental cellular process, if CCP1 participates in the degradation of proteasome-generated peptides, the absence of CCP1 activity in the pcd mice would be expected to produce a change in the intracellular peptideome. We also expanded our analysis of the adult pcd mouse brain by including peptides derived from secretory pathway proteins. The pattern of changes of peptides in adult mouse brain pcd mice was similar to the pattern found in mice lacking a gene necessary for the formation of primary cilia, a sensory cell organelle [27]. Because the disruption of primarily cilia has recently been shown to affect tubulin modifications [28,29] and proteasome function [30], our finding further suggests that the altered peptideome in adult pcd mouse brain is a secondary change due to affected tubulin processing.

**Materials and Methods**

**Animals**

Animal use experiments were approved by the Institutional Animal Care and Use Committee of Albert Einstein College of Medicine (protocol #20090305). Multiple breeding pairs of pcd heterozygous mice (BALB/cByJ- Agtpbp1 pcdf/f) were purchased from The Jackson Laboratory. Homozygous pcd mice and WT littermates were produced from matings of heterozygous from The Jackson Laboratory. Homozygous (Ift88tm1.1Bky) College of Medicine. The analysis of peptides in heart, two adult WT and three adult pcd mice was used. For the cilia mutant and control mice,Brains from the cilia mutant and control mice were collected at three months of age and processed for peptidomics analysis as described previously [31]. To delete cilia, six-eight week old female gld88-/- mice on the inductive Cagg-CreER (B6.Cg-Tg(CAG-cre/ Esr1*)5Amc/J) background were injected with 6mg/40gm body weight tamoxifen suspended in corn oil for five consecutive days. Brains from the cilia mutant and control mice were collected at three months of age and processed for peptidomics analysis as described below. Cilia and IFT88 protein loss was confirmed by immunofluorescence microscopy in brain sections and by Western blot analysis on parallel sets of mice as described previously [32,33].

**Quantitative Peptidomics**

For the analysis of peptides in young amygdala and cerebellum, two WT and three pcd mice were used, each analyzed separately. All animals were 3-weeks old. For the analysis of peptides in spleen, three adult WT and two adult pcd mice were used. For the analysis of peptides in heart, two adult WT and three adult pcd mice were used. The analysis of peptides in adult brain was performed with two pcd and two WT mice. For the cilia mutant mice, brain peptides were analyzed in six cilia mutant and three control mice. For all peptidomics studies, tissues were heat-treated using microwave irradiation, as described [34]. Peptide extraction, labeling with isotopic tags, sample preparations, and peptide identification from tandem mass spectrometry (MS/MS) analysis were performed as described [18,35]. Briefly, peptides were extracted with hot water and cold HCl, and labeled with the isotopic tags D0-, D3-, D6-, D9-, or D12-trimethylammonium butyrate (TMAB)-N-hydroxysuccinimide ester. After labeling, excess TMAB was quenched with Gly, the samples were pooled, filtered through YM-10 filters, treated with hydroxylamine to remove unspecified TMAB binding to Tyr residues, and desalted using C18 spin columns. Samples were analyzed by liquid chromatography/mass spectrometry (LC/MS) on a Waters QTOF-Ultima Mass Spectrometer or an API Q-Star Pulsar-i quadrupole time-of-flight mass spectrometer. MS/MS data analysis was performed using Mascot search engine (Matrix Science Ltd., U.K.). The IPI mouse database, which consists of 56,934 sequences; 25,565,245 residues, was searched. Methionine oxidation, N-terminal acetylation, and the isotopic D0-, D3-, D6-, and D9-TMAB tags were considered during peptide identification (D12-TMAB is not a search option on Mascot). The Mascot search results were manually interpreted using the following criteria: (1) The Mascot ion score was the top score of all analyzed peptides. However, this alone was insufficient to distinguish true positives from false positives, and all of the following additional criteria were also considered. (2) The D0-, D3-, D6-, or D9-isotopic form of TMAB identified by Mascot matched the isotopic form identified by analysis of the peak set. This criterion was important because Mascot does not discriminate between peak sets and cannot determine which of the peaks correspond to the D0, D3, D6, D9, or D12-labeled peptides. False positives have a 4 in 5 chance of having the wrong isotopic tag if one tag was incorporated, a 24 in 25 chance of having the wrong isotopic tags if two tags were incorporated, and a 124 in 125 chance of having the wrong isotopic tags if three tags were incorporated. Thus, this is an important criterion that excludes many false positives. (3) The number of labels incorporated into the peptide corresponded to the number of free amines (unmodified N terminus and side chains of Lys). If a peptide was labeled with several tags, then all tags represented the same isotopic form. (4) The major MS/MS fragment ions corresponded to predicted a, b, or y ions, internal ions, or precursor ions with loss of trimethylamine. (5) At least five fragment ions matched b or y ions. (6) The mass accuracy of the fragment ions was within 40 ppm, the accepted specification for the q-TOF instruments. (7) The observed charge state matched the predicted charge state based on the peptide sequence. (8) The fragment ions matched the expected ions based on the particular peptide sequence. For example, cleavage of Xaa-Pro bonds was favored and produced strong fragment ions while cleavage at Pro-Xaa bonds was rarely detected.

**Quantitative Real-time PCR**

Total RNA was isolated from heart and spleen of adult WT and pcd mice using RNeasy Mini Kit (Qiagen). Superscript III first strand kit (Invitrogen) was used to synthesize cDNA from 2 μg of total RNA and random hexamers. Primers used for mouse CCP1 through CCP6 and GADPH were previously described [23]. SYBR green fluorescent tag was introduced using Power SYBR Green PCR Master Mix (Applied Biosystems). PCRs were carried out on a 7900HT Real Time Thermal cycler (Applied Biosystems). All samples were run in triplicate. The threshold cycle number (Ct) was used to calculate quantitative values. The fold-change in expression was calculated using DDCt method.
confirmed in our colony of

Results

2 brains were fixed in 4% PFA overnight, incubated in 30% sucrose
PBS and then with 4% paraformaldehyde (PFA). Tissue prepa-

tion and staining were performed as indicated [36]. Briefly,
brains were fixed in 4% PFA overnight, incubated in 30% sucrose
in PBS for 6 h, and frozen in optimal cutting temperature gel at
−50°C. Cerebellum coronal sections (14 μm thick) were probed
with anti-Calbindin-D28k (1:3000, Sigma-Aldrich) followed by
secondary Cy2-conjugated antibodies [Jackson Immunoresearch].
The cerebellum sections were mounted in antifade reagent
(Invitrogen) and then analyzed by fluorescence microscopy.

If CCP1 functions in peptide degradation in adult mouse brain,
we reasoned that because protein/peptide degradation is such
a fundamental process, CCP1 would also function in this capacity
in young mouse brain, and therefore a peptidomics analysis of
young pcd mouse brain would also show marked changes in levels
of intracellular peptides. In contrast, if the previously observed
increase in levels of many brain peptides was a consequence of the
neurodegeneration in the adult pcd mice, then young mice
analyzed prior to the onset of neurodegeneration would not show
the increase in intracellular peptide levels. Purkinje cells start to
degenerate in pcd mice when animals are 3 weeks old [1]; this was
confirmed in our colony of pcd mice by immunohistochemistry
(Figure S1). To test if young presymptomatic mice show greatly
altered levels of the brain peptidome, a quantitative peptidomics
approach involving TMAB isotopic labels was used to compare
amygdala and cerebellum of 3 week old WT mice versus pcd mice.
The five distinct masses of the TMAB labels allow 2–3 mutant
mice to be compared to 2–3 WT mice in the same experiment;
this allows for variation among replicates of WT animals to be
determined along with the ratio of peptides in mutant versus WT
mice. Representative MS data are shown in Figure 1, comparing
relative levels of the same peptide in three different LC/MS runs.
Panel A shows the LC/MS run for adult amygdala in which two
pcd mice were compared to three WT mice. Panel B shows the
LC/MS run for 3 week old amygdala in which three pcd mice
were compared to two WT mice. Panel C shows the LC/MS run
for 3 week old cerebellum in which three pcd mice were compared
to two WT mice. For this peptide, levels are generally comparable
between the pcd and WT mice when tested at 3 weeks of age, but
clearly much higher in the adult pcd mice than the WT mice.
Altogether, 146 distinct peptides arising from intracellular proteins
were identified in the young amygdala and 177 distinct in-
tracellular peptides identified in the young cerebellum; there was
considerable overlap between the two sets of peptides (Figure 2).
These two sets of peptides also showed considerable overlap with
the intracellular peptides previously identified in the study
comparing the amygdala of adult WT and pcd mice [36]. Data
are provided in Table S1.

A graphic representation of the data was prepared by plotting
individual data points, sorted by the relative ratio of each
intracellular peptide in a pcd mouse tissue, relative to the average
level of that peptide in the same tissue of WT mice (Figure 3). For
this analysis, each data point represents a peptide detected in one
of the samples, the y-axis represents the ratio relative to the
average level in the controls, and the x-axis represents the rank
order of each data point (sorted from smallest ratio to largest ratio).
This representation of the data shows a dramatic increase in the
levels of most intracellular peptides previously identified in adult
pcd mouse amygdala (Figure 3A, grey circles). For this analysis,
the ratio of peptides seen only in the pcd mouse tissue and not in WT
tissue was capped at 5, reflecting the typical signal-to-noise ratio
of representative data (see Figure 1A). Conversely, peptides seen only
in the WT tissue but not in the mutant tissue would be capped at
a ratio of 0.2, also reflecting a 5-fold difference in the relative
levels. The actual changes, increases or decreases, may be much
larger than 5-fold, and this value is simply a conservative estimate
based on the typical baseline signal. In figure 3A, it is clear that the
majority of the identified peptides in the adult amygdala are like
those in Figure 1A; detectable only in the pcd mice and not in the
WT controls. For those peptides observed in WT animals, it was
possible to determine the peptide level in each animal relative to
the average WT value; this provides an estimate of mouse-to-
mouse variation. Most of the peptides detected in WT mouse
amygdala had ratios between 0.5 and 2.0 (Figure 3A, black circles).
Contrary to the results with adult mouse amygdala, only a few of
the intracellular peptides were greatly elevated in amygdala
(Figure 3B, grey line) and cerebellum of young presymptomatic pcd
mice (Figure 3G, grey line). Therefore, the absence of large
changes in intracellular peptides in young pcd mouse brain regions
suggests that accumulation of intracellular peptides later in life
reflects secondary changes due to the CCP1 mutation in pcd mice.

In addition to examining levels of intracellular peptides in young
brain tissue, we also examined peptides in adult mouse brain and
other adult tissues. Before performing peptidomic analyses, we first
needed to identify appropriate tissues for this analysis. It is
important to consider levels of CCP2-6 because these other
enzymes can potentially compensate for the absence of CCP1 in
pcd mice. Levels of CCP1 mRNA in WT and pcd mouse are higher
than levels of mRNA for CCP2-6 based on in situ hybridization
and reverse transcription PCR [4]. This is also supported by
bioinformatic analysis of expressed sequence tag (EST) databases
(http://www.ncbi.nlm.nih.gov/UniGene/). In addition to brain,
many other tissue show relatively high levels of CCP1 based on
PCR analysis and bioinformatics analysis of EST databases. For
example, heart and spleen have levels of CCP1 mRNA generally
comparable to brain [4]. Heart and spleen have much lower levels
of CCP2-6 mRNA, based on previous PCR results [4] and EST
database analysis. To confirm, and to determine if the other CCP
mRNAs are up-regulated in pcd mice, we performed QRT-PCR
on the heart and spleen of adult WT and pcd mice. As predicted
from EST database analysis, CCP1 mRNA is reasonably
abundant in WT mouse heart and spleen (Figure 4A and B,
respectively). The relative levels of all other CCP mRNAs in these
organs are much lower than CCP1 mRNA (Figure 4). The level of
CCP1 transcript in heart is lower in pcd compared to WT mice
(Figure 4). The levels of other CCP mRNAs are not significantly
changed in pcd mice, suggesting that there is no compensation for
the absence of enzymatically active CCP1 in mutant animals.

Peptide levels in adult pcd mouse heart, spleen, and brain were
compared to levels in WT mice. Data are included in Table S1,
and graphic representations shown in Figure 5. Levels of peptides
in pcd mouse heart and spleen are generally comparable to the levels
of peptides in the corresponding WT mouse tissues (Figure 5A and
B). In contrast, the majority of peptides derived from cytosolic and
mitochondrial proteins observed in the pcd mouse brain are not
detectable in control mouse brain, defined as a pcd/WT ratio >5
(Figure 5C). For the analysis in Figure 5C, adult mouse brains
lacking the cerebellum and olfactory bulb were used in order to
avoid potential problems due to cell death that occurs in these
brain regions. The changes in peptide levels observed in adult pcd
mouse brains in the present study were similar to the changes
previously reported when levels of intracellular peptides were
analyzed in adult pcd mouse brain hypothalamus, amygdala,
cortex, striatum, and hippocampus [18]. Thus, including the two
additional *pcd* mice analyzed in the present study, the dramatic increase in levels of intracellular peptides in the brain of adult mice was detected in 8 distinct *pcd* mice, compared to 11 distinct WT mice.

Altogether, 152 peptides derived from intracellular proteins were identified in the new analysis of the adult *pcd* mouse brain, whereas only 64 peptides derived from intracellular proteins were identified in the WT mice included in the LC/MS analysis (Figure 6). In addition to analyzing peptides derived from intracellular proteins, the new analysis of adult *pcd* and WT mouse brain considered all peptides detected in the sample, which included peptides derived from secretory pathway proteins (i.e. neuropeptides and related molecules). A total of 52 secretory pathway peptides were found in the WT brains and 48 were found in the *pcd* brains (Figure 6). However, when the relative levels of secretory pathway peptides were plotted on rank plots, there was a clear decrease in levels of these peptides in the *pcd* mice (Figure 7, top right panel). For comparison, the rank plot of the peptides from intracellular proteins previously shown in Figure 5C is included in Figure 7 (top left panel). To test if the decrease in secretory pathway peptides was also observed in the data on the adult amygdala and hypothalamus (which were derived from different groups of animals than the study on whole brain), we reanalyzed the previous data for peptides derived from secretory pathway proteins. As found for whole brain, some of the secretory
pathway peptides detected in amygdala and hypothalamus showed a decrease in *pcd* mice, relative to WT mice (Figure 7, right panels). However, some secretory pathway peptides also showed a large increase in *pcd* mouse amygdala and hypothalamus (Figure 7). The identity of these peptides and relative levels are included in Table S1, and representative peptides highlighted in the next two figures described below.

The quantitative peptidomics approach used in the present study has previously been used for the analysis of brain peptides in other mutant mice, and with one exception (described below), the pattern of changes in intracellular and secretory pathway peptides of adult *pcd* mouse brain was unique. Some of the mutants analyzed showed large changes in levels of secretory pathway peptides, but no large changes in levels of most intracellular peptides (Figure S2). These mutants include mice lacking carboxypeptidase E (*Cpefat/fat* mice) and mice with targeted disruptions of the genes encoding prohormone convertase 2 [37–39]. Because these enzymes function in the biosynthesis of neuropeptides, the change in levels of secretory pathway peptides was expected in these mutant mice. Other mutant mice lines that have been examined include mice with disruptions in the genes encoding prohormone convertase 1 (also known as prohormone convertase 3), proprotein convertase 7, endothelin converting enzyme 2, and *ift88*. Prohormone convertase 1/3, proprotein convertase 7, and endothelin converting enzyme 2 are peptidases in the secretory pathway while Ift88 is a protein required for intraflagellar transport and the formation of cilia [40–45]. In most of these previous studies, the relative levels of peptides derived from intracellular proteins were generally similar between mutant and WT mice. However, there was one exception; Ift88 cilia mutant mice showed a dramatic increase in the levels of the majority of hypothalamic peptides derived from intracellular proteins (Figure 7, lower left panel). Analysis of striatal and thalamic peptides showed a similar effect in the Ift88 cilia mutant mice (Table S1). Interestingly, hypothalamic peptides derived from
secretory pathway proteins were also found to be altered in this mouse mutant, with some peptides elevated and other decreased, relative to WT control mice (Figure 7, lower right).

Although the rank plots shown in Figure 7 (and other figures) are useful to compare the overall peptidome between different groups of mice, these plots do not convey information about the individual peptides. In order to compare individual peptides, we used heat maps in which peptides commonly found in the various analyses were selected and the average ratio of peptide in mutant mice, relative to WT mice, was calculated. Then, the value was color-coded, with bright green representing large decreases (ratio of mutant to WT $\leq 0.50$), dark green representing small decreases (ratio 0.51 to 0.80), grey representing no substantial changes (ratio 0.81 to 1.19), dark red representing small increases (ratio 1.20 to 1.99) and bright red representing large increases (ratio $\geq 2.0$). This analysis was performed for peptides derived from intracellular proteins (Figure 8) and also for peptides derived from secretory pathway proteins (Figure 9). For the intracellular peptides, many of the peptides found to increase in the adult whole brain were also found to increase in the adult amygdala and cerebellum (Figure 8). Only small changes were found in peptides in the 3 week old $pcd$ mouse cerebellum and amygdala, consistent with the rank plots shown in other figures (Figure 8). Interestingly, the $Ift88$ cilia mutant mice showed changes in many of the same intracellular peptides as observed in the adult $pcd$ mice (Figure 8).

Heat map analysis of the secretory pathway peptides was performed, and because many of these peptides were detected in WT mice, it was possible to compare levels using statistical tests. Of the 8 peptides that decreased substantially in the $pcd$ mice that were also found in the other experiments, 4 of these were significantly decreased in the $Ift88$ cilia mutant mice (Figure 9). Of the 16 peptides that were not significantly altered in the $pcd$ mice, all 16 were not significantly affected in the $Ift88$ cilia mutant mice either. Furthermore, the 8 peptides elevated in the $pcd$ mice were also found to be elevated in the $Ift88$ cilia mutant mice, although for these peptides statistical testing of the changes between mutant and WT mice could not be performed (statistical testing requires a number, and a ratio of $>5$ is not a number). In contrast to the general similarity between the changes in secretory pathway peptides between $pcd$ mice and $Ift88$ cilia mutant mice, there was no obvious correlation between either of these mutants and the $Cpefat/fat^{+/--}$ mouse or prohormone convertase 2 KO mouse (Figure 9).

The secretory pathway peptides were analyzed further in an attempt to understand why some peptides increased while others...
Figure 7. Relative levels of intracellular and secretory pathway peptides in adult WT and mutant mice. Peptides identified in adult whole brain (minus the cerebellum and olfactory bulb) or in amygdala or hypothalamus were included in this analysis. For the top three panels, each dot in the graph shows the ratio between a peptide in one WT or pcd mouse versus the average level in the WT replicates. Large black circles represent pcd mice, small black circles represent WT mice. The x-axis reflects the number of peptides found in pcd mice. For the bottom panels, the
decreased in the adult pcd mouse brain. For this analysis, the ratio of results from all secretory pathway peptides identified in amygdala, hypothalamus, and/or whole brain were combined to provide a larger number of peptides, and the changes in peptide levels were averaged so that for each peptide a single value was used. Then, the ratio between peptide levels in the pcd mouse and the levels of the corresponding peptide in WT mice within each LC/MS run was used to divide the peptides into five groups, as used for the heat map analysis. Similar to the heat map shown in Figure 9, which focused on only those peptides found in multiple experiments, the analysis of all identified peptides showed relative few peptides in the “slight increase” category (Figure 10, dark red bars), and more peptides in the other 4 categories. Analysis of the C-terminal amino acid of the peptides in each group did not reveal any obvious differences (data not shown). However, consideration of the cleavage site(s) required to generate the peptide revealed that most of the peptides that were substantially decreased in adult pcd mice arose from cleavage at prohormone convertase 1/3 and/or prohormone convertase 2 consensus sites (i.e. RR, KR, RxxR, and related sequences) or at a site containing a basic residue other than a prohormone convertase consensus site (Figure 10, Lower panel). Many of these peptides are known neuropeptides that represent the mature form present within large dense-core secretory granules. The majority of unchanged peptides were also in this group (Figure 10, Lower panel). Nearly all of the peptides that increased in adult pcd mice were produced by processing at a non-basic site on either the N- or C-terminal side of the peptide (Figure 10, Lower panel). Many of these cleavages may represent extracellular cleavage by peptidases present in the synapse or lysosomal enzymes after fusion of lysosomes with vesicles.

**Discussion**

The overall goal of the present study was to determine whether CCP1 plays a role in peptide turnover within the cell. Previously, some of the authors of the present study proposed that CCP1 contributed to peptide degradation based on the dramatic increase in levels of intracellular peptides observed in pcd mice [18]. While these authors had proposed that CCP1 could also play a role in tubulin processing, the simplest explanation to account for the increase in peptides derived from intracellular proteins was that these peptides represented substrates of CCP1. This was found to
be the case for peptides that accumulate in the brains of Cpefat/fat mice, which lack carboxypeptidase E activity due to a point mutation [46]. However, the Cpefat/fat mice showed both an increase in levels of substrates and a decrease in levels of products, as expected for a normal substrate-product relationship [47]. In contrast, the adult pcd mouse brain peptides were mainly elevated, which was interpreted to indicate that CCP1 degraded all peptides that it came into contact with [18]. However, the peptides elevated in adult pcd mouse brains contain a wide range of C-terminal amino acids; if these peptides represent substrates, then CCP1 has a very broad substrate specificity. This was a surprising observation; even the metallocarboxypeptidases present in the digestive system show marked preference towards particular amino acids [48–52]. Additional problems with this hypothesized function for CCP1 were found from studies overexpressing and knocking down the protein in HEK293 cells; only a small number of peptides were significantly elevated by the knock-down of CCP1, and no peptides were significantly altered by the overexpression of the protein [23]. Because protein turnover is a fundamental process that occurs in every cell of the body, it was expected that if CCP1 performs this role in adult mouse brain, it would also perform this role in cultured cells as well as in young mouse brain and other tissues. Therefore, the goal of the present study was to address the function of CCP1 by examining peptide levels in pcd mouse tissues that do not undergo degeneration, and young brain, prior to the degeneration of Purkinje cells.

Our present studies show that accumulation of peptides does not occur in the brain of young pcd mice and in heart and spleen of adult mutants. Since the adult mouse heart and spleen express high levels of CCP1 mRNA and low levels of other CCPs, it was expected that the lack of CCP1 in pcd mice would cause the same changes as in adult pcd mouse brain. It is conceivable that the normal peptide profile in pcd heart and spleen is achieved through a compensatory mechanism, although our results demonstrate that levels of mRNA encoding other CCPs are not upregulated by the absence of CCP1 in adult pcd mice. The finding that intracellular peptides are unchanged in the brain of young pcd mice also complicates the hypothesis that CCP1 functions in peptide turnover. Unless peptide turnover proceeds by a different mechanism in adult brain versus young mouse brain, adult spleen, and adult heart, these results suggest that the previous hypothesized role for CCP1 in peptide turnover is not correct.

The other hypothesized role for CCP1 involved the processing of tubulin, which undergoes a series of C-terminal reactions. For example, the C-terminal Tyr and Glu of alpha-tubulin are removed to form delta-2 tubulin. In addition, both alpha- and beta-tubulin can be modified by the attachment of Gly or Glu residues to the gamma-carboxyl group of Glu residues near the C-terminus.

**Figure 9.** Heat map representation of relative levels of secretory pathway peptides in adult pcd and other mutant mouse brains. Peptides derived from secretory pathway proteins that were commonly detected in the hypothalamus of cilia mutants, pcd, WT, Cpefat/fat, and prohormone convertase 2 knockout mice were included in this analysis. Each row represents a different peptide. The color shows the ratio between a peptide in one WT or mutant mouse replicate versus the average level in WT mice within each LC/MS run. Additional abbreviations: s.d., standard deviation; n, number of replicates; sig, statistical significance of difference with the WT/WT value (calculated using Student’s t-test); *, p < 0.05; **, p < 0.01; ns, not significant; fat/WT, the ratio of peptides in Cpefat/fat mice relative to WT mice; PC2KO/WT, the ratio of peptides in prohormone convertase 2 knock-out mice relative to WT mice. See Table S1 for data.

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Impaired tubulin polyglutamylation causes cilia in kidney cells due to the absence of Kif3a [58]. Recently, it was shown that tubulin is hyperacetylated in mutant animals, which fits with a role for CCP1 in the modification of tubulin. This leads to the formation of delta-2 tubulin, and also to the deglutamylating activity of CCP1 may not be required until later in development. In agreement with this, it was shown that ciliogenesis is not affected in C. elegans CCP1 mutant, but ciliary defects develop gradually in an age-dependent manner [59].

Although initially unexpected, our observation that the peptidome of adult pcd mouse brain is altered has many similarities to that of the cilia mutant animals also fits with a role for CCP1 in the modification of tubulin. Recently, it was found that tubulin is hyperacetylated in a kidney cell line that lacks primary cilia due to the absence of Kif3a expression [54]. Impaired tubulin polyglutamylation causes cilia dysfunction which leads to primary ciliary dyskinesia [53] and is associated with the spermatid flagellar structure in mice [36]. Recently, it was reported that hyper-glutamylation of β-tubulin destabilizes ciliary microtubules in T. thermophila [57]. Moreover, it was shown that CCP1 down-regulation causes cilia shortening in cell culture [58]. In addition, it was demonstrated that CCP1 regulates cilia localization and velocity of molecular motors, and loss of CCP1 affects cilia stability but not ciliogenesis in C. elegans [59]. Very recently, CCPs were linked to cilia using a bioinformatics approach [60]. Tubulin within the axonemal of cilia is highly glutamylated, and the levels of this modification were measured as the cilia mature [61]. Thus, the deglutamylating activity of CCP1 may not be required until later in life when cilia are formed. In agreement with this, it was shown that ciliogenesis is not affected in C. elegans CCP1 mutant, but ciliary defects develop gradually in an age-dependent manner [59].

While it is clear that CCP1 is involved in tubulin processing, and this can potentially account for the similar peptide changes in cilia mutant and pcd mouse brain peptides, the mechanism by which altered tubulin processing affects the brain peptideome is less clear. The changes in secretory pathway peptides could potentially result from altered trafficking and/or storage of the peptide-containing secretory granules. It has been shown that impaired polyglutamylation of tubulin affects synaptic transmission [25]. The changes in secretory pathway peptides observed in our data are consistent with elevated secretion of peptides accompanied by extracellular processing. This would account for the reduced levels of many mature neuropeptides and the increased levels of peptides that are produced by processing at non-basic sites. For example, the melanin concentrating hormone peptide, known as peptide EI, is greatly reduced in the pcd mice while the shorter form of this peptide lacking the N-terminal residue is greatly elevated in pcd mice (Table S1). Interestingly, peptides from broadly expressed secretory pathway proteins such as chromogranin B and proSAAS represent many of those peptides unchanged in the pcd mice while peptides from prohormones expressed in more limited cell types, such as protachykinin A, proenkephalin, prodynorphin, proopiomelanocortin, and promelanin concentrating hormone all have several peptides that are greatly decreased in the pcd mouse. This suggests that the changes in secretory pathway peptide levels in the pcd mouse are more pronounced in selected neurons.

The mechanism by which levels of peptides derived from cytosolic and mitochondrial proteins are altered by changes in tubulin processing is less clear. These peptides are presumably generated by the proteasome, based on recent findings that the vast majority of peptides present in human cell lines (HEK293 and SH-SY5Y) are affected by the proteasome inhibitors epoxomicin [62] and bortezomib [63]. Although this has not been examined yet for brain peptides, many of the peptides found in mouse brain are identical to those found in the cell lines [64,65], and therefore it is likely that the brain peptides are also generated by the proteasome. If so, then the observed results could be explained by an effect of tubulin modification on proteasome activity, either directly or indirectly. Knockdown of proteins essential for the formation of the basal body influences proteasome function, leading to altered protein turnover [30]. The altered levels of intracellular peptides in cilia mutant mice observed in the present study is consistent with altered proteasome function. Taken together, one possible explanation of the altered levels of intracellular peptides in adult pcd mouse brain is that the defect in CCP1 leads to altered ciliary function which in turn affects proteasome activity and peptide levels. Alternatively, the defect in CCP1 activity in the adult pcd mouse brain may lead directly to changes in proteasome function and peptide levels, independent of the effect of CCP1 on ciliary function. While many details of the precise mechanism by which CCP1 defects lead to altered peptide levels in adult mouse brain, the original model that CCP1 plays...
a direct role in the degradation of intracellular peptides is not supported by the data reported in the present paper or by other recent results showing CCP1 selectively cleaves Glu from tubulin.

**Supporting Information**

**Figure S1** Distribution of calbindin staining in cerebellum of 3-week-old WT and pcd mice. WT and pcd cerebellar sections were probed with antibodies against calbindin, a marker for Purkinje cells. Staining shows Purkinje cell dendrites, Purkinje cell bodies, and Purkinje cell axons (all green) in WT (left panel) and pcd (right panel) cerebellum.

**Figure S2** Relative levels of intracellular and secretory pathway peptides in the hypothalamus of adult mice. Data were plotted as described for Figures 3, 5, and 7. All values represent the ratio of the level of a particular peptide in the mutant mouse hypothalamic extracts, relative to WT mouse hypothalamic extracts. A. Intracellular peptides from Qp60filt mouse. B. Secretory pathway peptides from Qp60filt mouse. C. Intracellular peptides from prohormone convertase 2 KO mice. D. Secretory pathway peptides from prohormone convertase KO mice.

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**Table S1** Data for individual peptides found in WT, pcd, and other mutant mice discussed in the paper. Peptide sequences are indicated along with information on the protein precursor (name, subcellular location). The ratio of peptide levels in mutant versus WT tissue are indicated for each biological replicate. Different worksheets in the table correspond to the various mutant mouse studies.

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**Author Contributions**

Conceived and designed the experiments: IB JJS JW. Analyzed the data: IB JJS JW LDF. Contributed reagents/materials/analysis tools: RCP NFB BKY. Wrote the paper: IB LDF.
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