Review: Insights into molecular mechanisms of disease in neurodegeneration with brain iron accumulation: unifying theories

C. E. Arber*, A. Li†, H. Houlden* and S. Wray*

*Department of Molecular Neuroscience, †Reta Lila Weston Institute, Institute of Neurology, University College London, London, UK

Neurodegeneration with brain iron accumulation (NBIA) is a group of disorders characterized by dystonia, parkinsonism and spasticity. Iron accumulates in the basal ganglia and may be accompanied by Lewy bodies, axonal swellings and hyperphosphorylated tau depending on NBIA subtype. Mutations in 10 genes have been associated with NBIA that include Ceruloplasmin (Cp) and ferritin light chain (FTL), both directly involved in iron homeostasis, as well as Pantothenate Kinase 2 (PANK2), Phospholipase A2 group 6 (PLA2G6), Fatty acid hydroxylase 2 (FA2H), Coenzyme A synthase (COASY), C19orf12, WDR45 and DCAF17. These genes are involved in seemingly unrelated cellular pathways, such as lipid metabolism, Coenzyme A synthesis and autophagy. A greater understanding of the cellular pathways that link these genes and the disease mechanisms leading to iron dyshomeostasis is needed. Additionally, the major overlap seen between NBIA and more common neurodegenerative diseases may highlight conserved disease processes. In this review, we will discuss clinical and pathological findings for each NBIA-related gene, discuss proposed disease mechanisms such as mitochondrial health, oxidative damage, autophagy/mitophagy and iron homeostasis, and speculate the potential overlap between NBIA subtypes.

Keywords: autophagy, mitochondria, NBIA, neurodegeneration, Tau, α-synuclein

Introduction

Neurodegeneration with brain iron accumulation (NBIA) is a group of neurodegenerative diseases characterized by iron accumulation in the basal ganglia. Specifically, excess iron accumulates in the globus pallidus (GP) and the substantia nigra (SN) and can be visualized with MRI. The cortex and the cerebellum can be affected, and cerebellar involvement correlates with the most severe NBIA subtypes. The SN and the GP naturally contain high iron concentrations [1,2] and also have a high metabolic requirement, potentially predisposing these areas to iron-related damage.

The accumulation of iron has neither been proven to be cause nor symptom in NBIA. Iron can shuttle between two redox states and so is utilized by the cell for electron donation. ‘Free’ iron [the so called labile iron pool (LIP)] is highly reactive and can be destructive to the cell via formation of reactive oxygen species (ROS) [3]. For this reason, there are precise homeostatic mechanisms to tightly control iron levels in the cell (for central nervous system review, see Rouault [4]; for peripheral systems, see Andrews and Schmidt [5]). In the central nervous system, extracellular iron is bound by Transferrin and internalized into the cell via Transferrin receptor-based endocytosis and the channel DMT1. Cytosolic iron is stored in Ferritin structures or transferred to organelles that are iron rich,
such as mitochondria. Mitochondria have devoted iron importers and storage proteins. Cellular iron is exported via Ferroportin with the support of ferroxidases: Ceruloplasmin (Cp; mostly astrocytes) and Hephaestin (widespread). Oxidative state is controlled by ferroxidases through the pathway, such as DMT1, Ferritin and Cp. Transcription of these iron homeostasis genes is controlled closely via Iron Regulatory Protein 1/2, which control expression based on cellular iron concentration (reviewed [6]).

As well as iron accumulation, patients exhibit dystonia, parkinsonism and spasticity. Genetically confirmed pathological studies have identified protein aggregates and axonal swellings that are reminiscent of other common neurodegenerative disorders [7]. NBIA was previously known as Hallervorden-Spatz disease if onset was after the first decade of life and infantile-neuroaxonal dystrophy with early onset; however, genetic findings are redefining the disease.

Genetic screening over the last decade has identified 10 disease-associated genes which lead to NBIA; however, around 20% of NBIA cases remain genetically undefined [8]. At first glance, these 10 genes appear to be unrelated and are involved in diverse cellular pathways (Figure 1). Only two genes are directly associated with iron homeostasis.

A greater understanding of the NBIA genes and any shared cellular function will ultimately help to link common clinical presentation, MRI findings and disease processes. Additionally, elucidating disease mechanisms that lead to NBIA may be relevant to other diseases such as frontotemporal dementia (FTD), Parkinson’s disease (PD), Alzheimer’s disease (AD), Friedreich’s ataxia and amyotrophic lateral sclerosis (ALS), which display similar aspects of disease [9]. Thus, utilising the monogenetic orphan diseases is of great use to define disease processes that are involved in common and genetically undefined diseases.

The clinical features associated with the various genetic causes of NBIA are outlined in Table 1, and the pathology of NBIA is reviewed in Table 2. In this review, we will discuss potential disease mechanisms for each gene defect and finally propose potential cellular processes that link the entire NBIA spectrum.

**PANK2 and PKAN**

Mutations in the Pantothenate Kinase 2 gene (PANK2) lead to Pantothenate Kinase-associated neurodegeneration (PKAN, NBIA type 1). PKAN represents the most common genetic cause of NBIA and occurs in around two-thirds of NBIA patients [10,32].

PANK2 is a seven exon gene which is alternatively spliced to form two transcript variants [33]. Mutations have been described throughout the gene leading to autosomal recessive PKAN. There is a strong correlation between loss of enzymatic activity and disease onset, for example, two null mutations will present with an early-onset phenotype [34,35]. Despite the fact that PANK2 proteins form homodimers, it is unlikely that mutant PANK2 has a dominant negative effect, as co-expression of mutant PANK2 and wild-type PANK2 shows similar activity to wild-type PANK2 alone [33]. The most frequent mutation is G521R which is thought to contribute to 25% of the disease alleles [10], and this mutant protein fails to fold properly, exhibiting no enzymatic activity [33].

PANK2 is one of four human pantothenate kinase proteins and is specifically located in the mitochondria [36,37]. PANK4 is proposed to be nonfunctional while PANK1 and PANK3 are active in the cytosol. The PANK2 polypeptide contains a mitochondrial localization signal. There is a catalytic ATP binding domain located at the N-terminus of the protein and a domain of unknown function at the C-terminus of the protein. PANK2 homodimers are located in the mitochondrial intermembrane space.

**Pathology**

Nine confirmed PKAN patients have been studied post mortem, exhibiting predominant GP pathology (Table 2) [20,21]. In all cases, the GP was discoloured and showed excess iron accumulation. Iron was distributed in degenerating neurones, macrophages, astrocytes, microglia and also in perivascular regions. Neuronal cell loss was evident and largely restricted to the GP, together with associated axonal loss. Astrogliosis of the GP was a common feature.

Cortical and subcortical brain regions show axonal swellings and spheroid bodies. These represent degenerating neurones and dystrophic axons. Swellings are immunoreactive at varying degrees for ubiquitin, amyloid precursor protein and phosphorylated neurofilament [20,21]. A loss of myelin and vacuolization has been described in the pallidum [21], potentially linking PKAN disease processes to other demyelinating conditions.
Tau pathology has been noted to a minor extent in all cases, with tangles and threads present in the cortex [21]. Historically, NBIA was thought to be an α-synucleinopathy; however, genetic screening methods have been distinguished from PKAN, which shows no α-synuclein accumulation, with other NBIA subtypes that do, for example MPAN and PLAN (see below).

PANK2 mechanisms

The primary function of PANK is to catalyse the ATP-dependent phosphorylation of dietary pantothenate (vitamin b5) to 4-phosphopantethenate, the first step in Coenzyme A (CoA) biosynthesis. CoA is central to metabolism and is required for β-oxidation, the citric acid cycle and the metabolism of amino acids and ketone bodies. CoA is also required for synthesis of fatty acids and amino acids.

A prominent hypothesis for neurodegeneration caused by PANK2 deficiency is that the lack of active enzyme leads to build-up of substrates in the CoA biosynthetic pathway. This leads to an accumulation of N-pantothenyl cysteine and free cysteine. Cysteine build-up can chelate...
| Gene       | NBIA subtype                                      | % [8] | Associated diseases            | Clinical symptoms                                      | Onset                          | MRI characteristics                                                                 | Reference       |
|------------|--------------------------------------------------|-------|--------------------------------|--------------------------------------------------------|-------------------------------|--------------------------------------------------------------------------------------|-----------------|
| PANK2      | Pantothenate kinase-associated neurodegeneration (PKAN) (NBIA1) | 35–50 | HARP syndrome                  | Dystonia, spasticity and Parkinsonism                  | Juvenile and adult onset (3 years or approximately 20 years) | Hypointensity with central hyperintensity of the GP, referred to as eye of the tiger | Hayflick et al. [10] |
| COASY     | COASY protein-associated neurodegeneration (CoPAN) (NBIA2) | <1    | Spasticity, dystonia, dysarthria and Parkinsonism | Cognitive decline                                      | Juvenile onset (2.5 years) | Hypointensity with central hyperintensity of the GP | Dusi et al. [11] |
| C19orf12   | Mitochondrial membrane-associated neurodegeneration (MPAN) (NBIA4) | 6–10  | SPG43                          | Spasticity, dystonia, dysarthria and Parkinsonism      | Childhood onset (11 years) | Hypointensity of the GP and SN plus hyperintensity in the GP | Hartig et al. [12] |
| PLA2G6     | PLA2G6-associated neurodegeneration (PLAN) (NBIA2) | 20    | INAD, Dystonia, Parkinsonism   | Hypotonia, spasticity, dystonia, Parkinsonism and cerebellar ataxia | Infantile, juvenile and late onset (1 year or 4 years or >18 years) | Hypointensity of the GP, hypointensity of the basal ganglia, thalamus, and dentate | Morgan et al. [13] |
| FA2H       | FA2H-associated neurodegeneration (FAN) (NBIA5)     | <1    | SPG35, Leukodystrophy          | Parkinonsim, dystonia and dementia                     | Juvenile to late onset (<20 years) | Hypointensity of the GP, hypointensity of the basal ganglia, thalamus, and dentate | Kruer et al. [14] |
| WDR45      | β-propeller-associated neurodegeneration (BPAN) (NBIA8) | 1–2   | SENDA                          | Parkinsonism, dementia and some pyramidal signs       | Childhood onset              | Hypointensity of the GP/SN with central hyperintense line | Hayflick et al. [15] |
| ATP13A2    | Kufor-Rakeb syndrome                              | <1    | Juvenile and adult onset (5 years or <40 years) | Parkinsonism, dementia, and some pyramidal signs     | Juvenile to adult onset      | Hypointensity of the GP/SN with central hyperintense line | Schneider et al. [16] |
| DCAF17     | Woodhouse Sakati syndrome                         | <1    | Diabetes, alopecia, hypogonadism, deafness | Dystonia, dystonia and mental retardation              | Juvenile to adult onset      | Hypointensity of the GP/SN with central hyperintense line | Alazami et al. [17] |
| CP         | Aceruloplasminaemia (NBIA3)                        | <1    | Diabetes and anaemia           | Dystonia, dyskinesia and cerebellar ataxia            | Adult onset (51 years)       | Hypointensity of the GP/SN with central hyperintense line | Yoshida et al. [18] |
| FTL        | Neuroferritinopathy (NBIA1)                       | <1    | Juvenile onset                 | Dystonia, spasticity, rigidity and Parkinsonism       | Adult onset (39 years)       | Hypointensity of the GP/SN with central hyperintense line | Curtis et al. [19] |

© 2015 The Authors. *Neuropathology and Applied Neurobiology* published by John Wiley & Sons Ltd on behalf of British Neuropathological Society.
### Table 2. Pathological findings from gene-confirmed NBIA cases

| Gene confirmed pathology cases | Gross morphology findings | Iron | Axonal spheroids | Lewy body pathology | Tau pathology | Gliosis | Refs |
|-------------------------------|---------------------------|------|------------------|---------------------|---------------|---------|------|
| PANK2                         | Neuronal loss in GP. Reduced myelin | GP neurones, glia, microglia, macrophages, perivascular and iron shunting | GP | No | Occasional tangles and threads | GP and widespread | Kruep et al. and Li et al. [20,21] |
| COASY                         | Proposed similar to PANK2 | GP neurones, macrophages, less in astrocytes | GP | Severe Lewy bodies and Lewy neurites | Rare hyperphosphorylated Tau inclusions | Widespread | Hartig et al. and Hogarth et al. [12,22] |
| PLA2G6                        | Cerebellar, cortical, GP and brain stem atrophy | GP and sparse in SNr | Severe p-NF, Ubq, APP, α-syn | Severe, Lewy bodies and Lewy neurites | Early onset hyperphosphorylated Tau inclusions, threads and tangles | Variable | Gregory et al. and Paisán-Ruiz et al. [23–25] |
| FLH                           | Proposed brainstem atrophy and demyelination | GP, SN > GP neuronal loss | Strongest in SN | No | Tau tangles, hippocampus, cortex, putamen, few in atrophied SN and GP | Putamen and thalamus | Hayflick et al. [15] |
| ATP13A2                       | Peripheral biopsies show demyelination and cytoplasmic inclusions in nerve and muscle tissue | Similar to neuron loss, cerebellum, GP > SN, cortex | Unknown | Unknown | Yes | Gonzalez-Cuyar et al., Kaneko et al., Martín et al., Olde et al. [26–29] |
| DCAF17                        | Peripheral biopsies show denervation of muscle tissue | Unknown | Unknown | Unknown | Yes | But some atrophy too | Curtis et al., Vidal et al., Mancuso et al. [19, 30, 31] |
| FTL                           | Mild atrophy in the cerebellum, cortex, putamen, GPe and SN mildly affected | Cerebellum and putamen | Yes, GP Ubq and NF | Unknown | Few | | |

**NAN 2016: 42: 220–241**

© 2015 The Authors. Neuropathology and Applied Neurobiology published by John Wiley & Sons Ltd.

**APP, amyloid precursor protein; GP, globus pallidus externa; GPe, globus pallidus interna; NF, neurofilament; SNc, substantia nigra pars compacta; SNr, substantia nigra pars reticulata; Ubq, ubiquitin; α-syn, alpha synuclein.**
iron and lead to its accumulation. Free cysteine can auto-
oxidize in the presence of iron and produce ROS, resulting
in widespread oxidative damage and cell death, perhaps
through lipid peroxidation [32,38]. Iron content is natu-
really high in the GP and SN [1], potentially sparking
degeneration synergistically with excess cysteine, followed
by a negative feedback of iron accumulation and ROS-
derived damage.

The localization of PANK2 at the mitochondria (Figure 1) prompted several groups to investigate mitochondrial dysfunction in PKAN pathogenesis. PANK2 mutant fibroblasts [39] as well as mouse and fly pank knockout models [40–42] have hinted to mitochondrial deficiencies, including reduced mitochondrial membrane potential, swollen mitochondria with altered cristae and ruptured membranes. ROS damage has been described with varying degrees.

The loss of mitochondrial membrane potential can affect the rate of mitophagy, and it is feasible that altered mitochondrial fission and/or trafficking, together with membrane remodelling deficiencies due to lower levels of CoA and acyl-CoA, might contribute to axonal swellings [43,44].

Drosophila have one pantothenate kinase, fumbi. The fumbi hypomorphic model displays locomotor defects and lipid dyshomeostasis which are fully rescued by human PANK2 [40]. This highlights the high degree of protein conservation between species [45]. Importantly, human PANK3 and PANK4, which are cytosolic proteins, cannot fully rescue drosophila mutants, hinting to a cellular dependence on specific PANK2 activity [45].

There is a rate limiting feedback inhibition step by acyl-
CoAs on the ATP binding pocket of PANK2 [34]. The negative regulation of PANK2 by acetyl-CoA is so tight that mechanisms are in place to relieve this inhibition. For example, palmitoylcarnitine activates PANK2 in physiological conditions [46]. Enzymatic regulatory deficien-
cies could thus be involved in disease. However, three mutant proteins tested in vitro display similar acetyl-CoA inhibition [33].

PANK2 may have a role in sensing CoA concentration and the metabolic cross-talk between the mitochondria and the cytoplasm [46]. This could explain why genetic mutations not affecting enzymatic activity could lead to a disease phenotype. In addition, pantothenate kinase-derived CoA is responsible for acetylation of proteins [47]. Protein acetylation and CoA levels have been shown to directly link cellular metabolism and autophagic flux
[48,49]. This implicates a new pathway that is distinct from the well-documented processes controlling autophagy, including AMPK and mTOR1.

PANK2 mutations could disrupt a putative mitochondrial-specific fatty acid synthase pathway – critical for mitochondrial membrane assembly and function [37]. One study has confirmed lipid dyshomeostasis via a metabolomics approach on blood plasma of PKAN patients [50]. Furthermore, the clinical overlap described between PKAN and HARP syndrome (hypoprebetalipoproteinaemia, acanthocytosis, retinitis pigmentosa and pallidal degeneration), which both derive from mutations in PANK2, strongly links lipid production deficits in PANK2 mutants [51].

It is important to consider the role of excess iron in PKAN. PANK2 is targeted to the mitochondria, which are a major sink of iron within the cells, and multiple proteins involved in the electron transport chain contain iron-sulphur clusters. Knockdown of PANK2 in HELA cells and hepatocytes leads to a marked induction of ferroportin mRNA, the sole cellular exporter of iron [52]. Additionally, patient fibroblasts showed a reduced response to excess iron, with reduced expression of ferritin (the cellular iron storage protein) and an increased LIP [39]. Increased labile iron could lead to ROS directly, due to the Fenton reaction, whereby water is broken down in the presence of iron to form ROS.

In summary, PKAN patients display excess iron deposition and neuronal cell death, primarily in the GP. PANK2 mutations might affect metabolism and lipid turnover and ultimate affect energy production or mitochondrial fitness. Conversely, reduced CoA levels have not been proven in patient tissue, and so, disease mechanisms through alternate unknown PANK2 functions remain a possibility. Investigations into cellular mechanisms of cell death are required to understand the reasons for specific degeneration due to this gene defect.

COASY and CoPAN

Mutations in the CoA Synthase (COASY) gene were recently shown to lead to NBIA [11]. Clinically, COASY protein-associated neurodegeneration (CoPAN) patients display similar clinical features to PKAN patients [11].

The COASY protein contains a mitochondrial localization signal, a regulatory region and a domain for each of the two catalytic kinase domains: adenyl transferase and adenyl transferase...
COASY is localized to the mitochondrial matrix [11,53]. There are two COASY isoforms, produced from alternate splicing. The longer β isoform is brain specific and has an additional proline-rich protein interaction domain but has identical enzymatic activity to the ubiquitous α isoform [54].

Two CoPAN mutations described by Dusi and colleagues represent a premature stop codon (G49Stop) and a mutation affecting a conserved residue involved in ATP binding and dephospho-CoA kinase activity (R499C). These are proposed to completely abrogate the protein functionality [55].

Interestingly, yeast knockout models for the two proteins that are COASY orthologues lead to a lethal phenotype. Human wild-type COASY protein can rescue this lethality, and mutant COASY from patients can rescue knockout cells but lead to a higher requirement for pantothenate in the growth media [11].

COASY mechanisms

COASY is in the same metabolic pathway as PANK2 and is a bifunctional enzyme catalysing the final two steps of CoA synthesis [53,56]. Mutant COASY protein appears to have no activity in vitro, whereas CoA levels in patient and control fibroblasts appear normal [11]. This suggests CoA levels are maintained by either residual activity of the COASY protein, or via an alternative, as yet unknown, pathway for CoA synthesis.

The fact that COASY acts in the same pathway as PANK2 suggests common mechanisms may be shared between PKAN and COPAN, such as reduced acyl-CoA and lipid synthesis leading to mitochondrial insufficiencies.

Of the 10 NBIA-related genes, only COASY shows a putative iron response element (IRE) via the SIREs 2.0 prediction tool (http://ccbg.imppc.org/sires/). The sequence is a putative 3'-IRE and would lead to mRNA stabilization in the presence of iron. This could potentially link COASY expression to iron availability in the cell.

C19orf12 and MPAN

C19orf12 is a protein located on the outer mitochondrial membrane with unknown function (Figure 1). Mutations in C19orf12 lead to mitochondrial membrane protein-associated neurodegeneration (MPAN). MPAN accounts for around 30% of NBIA [22]. Other diseases associated with C19orf12 mutations include pallido-pyramidal syndrome [57], hereditary spastic paraplegia type 43 (SPG43) [58] and ALS [59].

C19orf12 is a two-pass transmembrane protein with two alternative start codons. Missense mutations have been described to add charged amino acids to putative hairpin domains, potentially disrupting the 3D structure [12,58]. MPAN has been successfully modelled in drosophila [60].

Post-mortem examination of two MPAN cases [12,22] showed iron accumulation mainly confined to the GP. Iron accumulation was accompanied by neuronal atrophy, and depositions were present in neurones, macrophages and, to a lesser extent, astrocytes (Table 2).

Lewy bodies and Lewy neurites were a characteristic feature of MPAN with an extremely high burden [22]. Lewy neurites in the GP were adjacent to the atrophied region and Lewy pathology extended to the SN where they were associated with near complete neuronal loss. Lewy bodies were also present in the cortex, striatum and hippocampus. Axonal swellings and hyperphosphorylated tau were also present in the cortex, GP, caudate/putamen, SN and brain stem. Demyelination was evident in the spinal pyramidal tracts and optic nerve.

C19orf12 mechanisms

C19orf12 may be expressed in the ER as well as the mitochondria [58], and pathogenic mutations that lead to SPG43 or MPAN alter the distribution of the protein, potentially hinting to a lack of proper protein folding and enzymatic activity. C19orf12 is expressed in neurones, white blood cells and adipocytes [60]. During in vitro differentiation of white blood cells, C19orf12 expression closely follows fatty acid metabolism, suggesting a link to lipid homeostasis [12]. It was further proposed that this function could be associated with CoA metabolism, linking MPAN to PKAN and CoPAN [12].

Therefore, C19orf12, PANK2 and COASY are linked through mitochondrial localization, lipid metabolism and ultimately iron deposition and Tau pathology in disease.

PLA2G6 and PLAN

Mutations in the gene encoding phospholipase A2 group VI calcium-independent (PLA2G6, also known as iPLA2β or iPLA2-VIA) have been associated with two diseases: phospholipase associated neurodegeneration
(PLAN, NBIA type 2) and dystonia-parkinsonism (Table 1). PLA2G6-associated disease may have previously represented infantile neuroaxonal dystrophy (INAD) before NBIA reclassification.

There are four groups of PLA2s containing more than 20 proteins: secreted PLA2s, calcium-dependent cytosolic PLA2s, platelet-activating acetylhydrolases and calcium-independent PLA2s (iPLA2) [61]. PLA2G6 is responsible for 70% of total PLA2 activity in the brain [62].

The PLA2G6 gene encodes an 806 amino acid protein with predicted size of 88 kDa. The mature protein is tetrameric and composed of an ankyrin repeat domain that is involved in protein interaction, a GXSG lipase motif, an ATP binding pocket and a Calmodulin binding domain [61]. There are at least five splice variants, three of which contain the lipase motif GXSG, whereas two contain only the ankyrin repeats and most likely act as competitive inhibitors within the tetramer [63].

PLA2G6 has been shown to be expressed in the mitochondria [64,65] and be protective to mitochondrial health [66]. Expression is also seen at the nuclear envelope and at axon terminals in primate brain tissue [67]. PLA2G6 is ubiquitously expressed [68], although disease symptoms are solely neurological.

Autosomal recessive mutations are found throughout PLA2G6 and lead to reduced enzyme activity [13]. The magnitude of effect on enzyme activity correlates with disease severity: two null copies of PLA2G6 have the most severe phenotype [23]. Dystonia-parkinsonism mutations do not affect the enzymatic ability of the protein but could alter protein interactions [69].

The enzyme can undergo autoacylation by oleoyl-CoA near the catalytic residue [70]. Acylation leads to a cytosolic to membrane relocation of PLA2G6 and may lead to precise subcellular targeting of the enzyme and potential catalytic activation (Figure 1).

Pathology

The pathology of seven confirmed PLAN cases have been studied (Table 2). The major pathological hallmark of PLAN is the presence of axonal swellings throughout the cortex, GP, striatum, cerebellum, brain stem and spinal cord. These are also present throughout the peripheral nervous system and can be diagnostic via peripheral nerve biopsies. There is evidence for PLA2G6 mutation-associated PLAN without spheroids [13]. Peripheral spheroids are not observed in dystonia-parkinsonism subset of PLAN patients [71].

Spheroids stain positive for phosphorylated neurofilament, ubiquitin and APP. α-Synuclein positive Lewy bodies are found throughout the degenerating brain areas to a very high degree, and a proportion of the axonal swellings also stained positive for α-Syn. Lewy bodies were specifically seen in the SN pars compacta. Cases also exhibited advanced tau pathology in early adulthood [23–25], whereby braak stage V/VI neurofibrillary tangles and neuropil threads were evident in cortical areas at the age of 18 and 32 [25]. Granules within spheroids of pla2g6 mutant mice were positive for Periodic-acid Schiff (PAS) staining. This is likely to represent polysaccharides, glycated proteins or aldehyde products of ROS-damaged lipids [72].

Tubulovesicular structures, another characteristic of PLAN, are membrane-rich inclusions and thought to contain mitochondria, lysosomal components, vacuoles, endoplasmic reticulum as well as occasionally a protein component [73].

Atrophy is evident in the cortex, GP, white matter, the cerebellar granule cells and cerebellar purkinje cells and was shown to be associated with gliosis in the cortex and the striatum [25]. Excess iron deposition was specifically observed in the GP and the SN pars reticulata [23]. There is extensive cortical involvement in disease, and Lewy bodies are seen in all cases whereas tau deposition is rarer [24,25].

PLA2G6 mechanisms

PLA2G6 hydrolyses glycerophospholipids at the sn-2 position of acyl chains to produce lysophospholipids and free fatty acids, such as arachidonic acid. The downstream metabolites of free fatty acids such as leukotrienes and prostaglandin perform essential cellular functions and contribute to a variety of signalling events including membrane remodelling, fatty acid oxidation, cell signalling and growth, and apoptosis [61]. The 2-lysophospholipid may also play a role in signalling such as in the production of platelet activating factor. Phospholipase activity is crucial for membrane integrity via phospholipid recycling and homeostasis.

PLA2G6-mediated neurodegeneration may occur via the inability to remodel oxidized and damaged phospholipids. Polyunsaturated mitochondrial components, such as cardiolipin, are extremely sensitive to ROS. PLA2G6 has an increased affinity to membranes in hydrogen peroxide-treated cells resulting in a raised activity and increased...
free fatty acid release [74]. Cells that aberrantly produce ROS sequester PLA2G6 to the mitochondria, and this is protective against apoptosis [66]. Patient cells were shown to display faulty mitochondrial at the ultrastructural level [75,76].

Uncoupling of the mitochondrial respiratory chain and associated depolarization (e.g. in de-energized states or after calcium ion influx) can lead to activation of PLA2G6 within the mitochondria and therefore free fatty acid accumulation [77]. This can initiate apoptosis through stimulating the permeability transition pore and cytochrome C release leading to further lipid damage and apoptotic signals [65,78]. Reduced PLA2G6 activity may lead to dysregulation of this process.

Related to this, ER stress can increase the expression and activity of PLA2G6, bringing about the loss of mitochondrial membrane potential and increasing the likelihood of apoptosis [79].

PLA2G6 plays an important role in membrane homeostasis. The lipid profile of spinal cords of Pla2g6 knockout mice is greatly altered suggesting a defective remodelling ability. This may be causal for degenerative inner mitochondrial membranes and axonal termini [80]. Altered axonal and/or organelar membrane integrity may lead to improper axonal transport and accumulation of cellular components at distal axon locations, subsequently leading to axonal blockage and degeneration [81].

Both products of PLA2G6 catalysis, lysophospholipids and free fatty acids, may be able to increase membrane fusion via increased fluidity and nonbilayer structures in the local environment [61]. This could have important roles in autophagy, mitophagy and other vesicle-based processes that have been shown to be involved in neurodegeneration.

Using pla2g6 knockout mice, Beck and colleagues proposed that the disease progresses via two concurrent mechanisms [80]. Firstly, mitochondria exhibit degenerative inner membranes, and a large component of swellings was shown to be mitochondria. Faulty mitochondria transported through the axons may release ROS and proapoptotic factors that damage neighbouring membranes, producing axonal swellings. Degenerative mitochondria are accompanied by local cytoskeletal disappearance, exacerbating mitochondrial transport deficiencies. Secondly, at axon terminals, membranes and synaptic vesicles degenerate and form swellings and tubulovesicular structures. Together, the lack of membrane remodelling is proposed as the cause of the disease phenotypes seen within axons and at axon terminals, two sites where PLA2G6 is localized [66,67].

Finally, there is evidence for a noncell autonomous effect of PLA2G6 disruption. Docosahexaenoic acid (DHA) is an essential fatty acid that cannot be synthesized in neurones. Astrocytes have been shown to produce and release DHA due to PLA2G6 action [82]. Knockdown of the enzyme in astrocytes leads to a reduction of arachidonic acid and DHA in neurones and increased prostaglandin production [83], which could lead to increased apoptosis [84].

Calcium signalling was shown to be defective in astrocytes from pla2g6 mutant mice and in astrocytes treated with a PLA2G6 inhibitor [85]. This suggests that cross-talk between neurones and glia may be disrupted in PLAN/INAD.

PLA2G6 is classified as a PD gene (PARK14), and also, a reduction in PLA2G6 protein levels has been shown in the brains of Alzheimer’s patients [86]. Intriguingly, the SN shows low endogenous levels of PLA2G6, possibly showing a vulnerability to lipid oxidation [87]. Mitochondrial involvement, lipid turnover and Tau pathology are again implicated in this NBIA subtype.

FA2H and FAHN

Fatty acid 2 hydroxylase (FA2H) mutations are linked to leukodystrophy, hereditary spastic paraplegia (SPG35) and NBIA [14,88,89].

FAHN is a recessive disease, and it has been shown for SPG35-associated mutations that the enzyme is nonfunctional [88]. FA2H is a 43 kDa membrane-bound protein, residing in the ER [90]. The polypeptide has a C terminal sterol desaturase domain, which contains an iron binding histidine motif and is responsible for catalytic activity, and an N terminal cytochrome b, haem-binding domain, involved in redox activity and electron donation to the C terminus [91,92].

Mouse knockout models provide a hint towards FAHN pathology, whereby the CNS exhibits enlarged axons and demyelination [93]; however, no human post-mortem studies have been reported.

FA2H mechanisms

FA2H is an enzyme that catalyses the hydroxylation of fatty acids at position 2 of the N-acyl chain (Figure 1). 2-Hydroxy-fatty acids are a precursor for Ceramide
synthesis, a critical component of myelin sheaths [90]. Inactivating mutations in FA2H is likely to affect normal myelin production due to loss of the hydroxylase activity of the enzyme. The late onset and slow progression are consistent with this idea, analogous to demyelinating diseases such as multiple sclerosis (MS).

Altered ceramide signalling may have roles in Lewy body formation, and the role of ceramide in apoptosis and neurodegeneration has been implicated [94,95]. Myelin formation via FA2H is dependent on lysosomal acid ceramidase and also fatty acid oxidation in peroxisomes, potentially linking cellular compartments that are common between NBIA genetic disorders.

FA2H also has a role in lipid signalling pathways that can affect cell cycle and apoptotic pathways [92]. Iron storing Ferritin has been shown to associate with myelin [96], and it was postulated that iron accumulation in disease could be associated with faulty myelin affecting Ferritin dynamics [14].

Axonal myelination may be a common factor between NBIA subtypes. PANK2 and COASY are both involved in CoA synthesis. CoA has many diverse cellular roles but is critical for the generation of sphingolipids, another principle component of myelin [50]. Defective myelination may contribute to neuronal dysfunction and apoptosis and is shared between several neurodegenerative diseases such as MS.

WDR45 and BPAN

Mutations in the WD repeat domain 45 gene (WDR45) were linked to a form of NBIA and termed β-propeller-associated neurodegeneration (BPAN) [97].

Despite WDR45 being on the X chromosome, BPAN does not follow normal X-linked dominant inheritance. Both genders have similar clinical features, and the disease is always sporadic. Disease-associated mutations are predicted to lead to nonfunctional proteins, and are presumed to be lethal for male embryos. Male BPAN sufferers are presumed to have de novo mutations and were shown to have somatic or germline mosaicism [97].

One confirmed BPAN patient has been studied post mortem (Table 2) [15]. Iron deposition was strongest in the SN but also evident in the GP, concurrent with cerebral and cerebellar atrophy (purkinje and granule cells). Axonal spheroids were present in the basal ganglia (especially the GP and the SN), and gliosis was prominent. Tau-positive neurofibrillary tangles were common in the neocortex, putamen and the hippocampus, and less common in the atrophied GP and SN. Amyloid-β plaques and Lewy bodies were not evident [15]. Pathological changes appear to affect the SN preferentially over the GP.

WDR45 mechanisms

WDR45 (also known as WIPI4) is a β-propeller scaffold protein that has been predicted to have a role in autophagy. WDR45 is a member of the WD40 protein family, which provide a basis for protein–protein interactions and perform cellular functions such as autophagy, cell cycle progression and transcriptional control. WDR45 binds to phospholipids and autophagy-related proteins [98].

The functions of WDR45 have been investigated using several models. Yeast have over 30 autophagy-related genes (atg), of which many mammalian homologues have been found. WDR45 is one of 4 atg18 homologues, critical for autophagosome formation (WIPI1-4) [99]. atg18 binds the ER membrane via a phosphotidylinositide-3-phosphate binding motif and facilitates downstream protein complex formation [100] (Figure 1).

Phosphatidylinositol-3-phosphate (PI3P) is a critical factor in autophagy and is a major component of autophagosome membranes. Upstream signalling (e.g. mTOR) controls PI3P and autophagic flux. PI3P may be involved in early membrane curvature and autophagosome size. Therefore, WDR45 could regulate autophagosome size and maturation [99]. The Caenorhabditis elegans homologue of WDR45, epg-6, was shown to be required for regulating early autophagosome size and interacts with several other ATG-like genes [98]. Epg-6 recruits atg9 which is thought to supply lipids for newly forming autophagosomes [101].

Lymphoid cells from five patients with truncated or destabilized WDR45 protein show a blockage in autophagic flux when exposed to inhibitors or activators of autophagy [102].

atg Molecules are transiently present on mitochondrial outer membranes suggesting a link between WDR45 autophagosomes and mitochondrial function [103]. H2O2 and ROS generated in mitochondria perform an essential role in oxidising and inactivating atg4 so that the autophagosome may form [104], leading Scherz-Shouval and Elazar to hypothesize a signalling gradient that allows autophagy biogenesis in the vicinity of mitochondria.

© 2015 The Authors. Neuropathology and Applied Neurobiology published by John Wiley & Sons Ltd on behalf of British Neuropathological Society.
MRI scans, from patients with early disease stage presentation, suggest that iron accumulation is observed in the SN early in disease progression, and GP iron accumulation is a later event [15]. This is in contrast to other NBIA subtypes.

Finally, genes involved in autophagy are disrupted in other disease conditions. PD, Crohn’s disease, cancer and spastic paraparesis have all been shown to have disrupted autophagy [105]. The pathology of BPAN and the strong involvement of the SN potentially support a shared mechanism to PD.

**ATP13A2 and Kufor-Rakeb Syndrome**

Mutations in ATP13A2 lead to Kufor-Rakeb Syndrome, which is a disease exhibiting juvenile onset parkinsonism and dementia (PARK9) [106], neuronal Ceroid-Lipofuscinosis [107] and NBIA [16].

Mutations in ATP13A2 are autosomal recessive and cluster on the cytoplasmic domains, interfering with catalytic activity [108]. Transmembrane stretches may also be affected and consequently the protein architecture.

There are no post-mortem studies of ATP13A2 mutation-associated NBIA. However, studies of sporadic PD patients have shown that ATP13A2 staining is increased in cortical and nigral neurones [108,109], and ATP13A2 protein is found associated with Lewy bodies [110]. Post-mortem studies of Kufor-Rakeb Syndrome patients confirm the atrophy seen with MRI and also lipofuscinosis in neurones and glia of the cortex, basal ganglia and cerebellum [107].

**ATP13A2 mechanisms**

ATP13A2 is a lysosomal P-type ATPase that functions as a divergent cation transporter. P-type ATPases are a large family of transporters that also include calcium pumps, proton pumps and phospholipid flipases. This superfamily consists of highly conserved, 10-pass transmembrane proteins, which utilize the energy stored in ATP to transport ions across membranes [111].

ATP13A2 was shown to be associated with membranes of the lysosome [108] (Figure 1), although it has also been linked with mitochondrial and synaptic membranes [109]. Knocking down the expression of ATP13A2 affects the size and number of autophagosomes [109]. Mutations in ATP13A2 may cause a mislocalization of the protein to the ER [112].

The overexpression of ATP13A2 in model organisms has been shown to protect against potentially cytotoxic environments such as α-synuclein overexpression [112,113] and heavy metal ions including cadmium, manganese, nickel and selenium [114]. Indeed, patient fibroblasts from ATP13A2 mutants displayed lysosomal deficiencies that lead to cytotoxic effects in the presence of α-synuclein [115] and zinc [116]. Intracellular manganese levels were shown to be higher in cells with mutant ATP13A2 compared with wild type, hinting to a reduced secretion ability [112]. This may directly lead to cytochrome C release from mitochondria and apoptosis.

The increased cytosolic heavy metal status of the cell may be linked with the characteristic fragmented mitochondrial phenotype witnessed in mutant cells [109,117]. Patient-derived fibroblasts and olfactory neurones have exhibited fragmented mitochondria, reduced ATP production, oxidative stress and mitochondrial DNA lesions [118] that may be in part due to reduced intracellular free zinc levels and a sensitivity to environmental zinc [117]. Transcription of zinc homeostatic genes is upregulated in a compensatory manner.

Metal ions are enriched in acidic lysosomal compartments and could be involved in destructive conditions required for degradation and recycling events [119]. Additionally, divergent metal ion transporters are unfaithful and appear to function to transport a range of metal ions. Although iron has not been shown to be directly dysregulated in ATP13A2 deficient cells, it will be interesting to investigate this further due to the accumulation seen in the putamen of Kufor-Rakeb patients.

Another P-type ATPase, ATP7B, is mutated in Wilson’s disease. Upon elevation of cytoplasmic copper levels, wild-type ATP7B translocates from the golgi to the lysosome whereby it aids the loading of copper into lysosomes. Copper-rich lysosomes are then secreted from the cell via lysosomal exocytosis [120]. Wilson’s disease results from dysfunctional ATP7B, increased copper levels and altered redox state. Therefore, the faulty ATP13A2 protein may reduce a noncanonical cellular excretion mechanism for heavy metals (Figure 2).

Post-mortem studies of NBIA-associated ATP13A2 patient brains have not been described but will be informative with regard to pathological hallmarks. The fact that ATP13A2 is a lysosomal disorder means that there are tantalising links to lysosomal storage diseases. Additionally, patients exhibit dementia and cortical atrophy, and iron accumulation in the caudate and the putamen,
hinting to ties with FTD and HD. Similar to other NBIA subtypes, there is an effect on mitochondrial health.

DCAF17

DCAF17 (also called C2orf37) is a protein of unknown function that is expressed in the nucleolus (Figure 1). Mutations in this gene lead to Woodhouse-Sakati syndrome (WSS), and a subset of patients displays brain iron accumulation [17]. Many patients display no iron accumulation; however, the GP and the SN are affected in some patients [8].

WSS manifests during adolescence and presents with extrapyramidal symptoms, dystonia and cognitive decline. Typically, patients exhibit hypogonadism, alopecia and diabetes mellitus.

Very little is known about the function of DCAF17; however, the clinical phenotype of the patients and the expression data are very similar to the ribosomal synthesis deficiencies of patients with RBM28 mutations [121].

DCAF17 is a multipass transmembrane protein that is named due to protein–protein interactions, Ddb1 and Cul4-associated factor (damaged DNA binding protein 1 and cullin 4 ubiquitin ligase complex) [122]. This association links DCAF17 to protein ubiquitination involved in DNA damage and cell cycle control. The nucleolar localization leads to intriguing questions about specific function and any links to other NBIA-linked mutations.
Iron

Many transition metal ions are critical cellular components. Iron is an essential dietary component and acts as a crucial cofactor for enzymes and proteins. Iron is synthesized into iron-sulphur clusters and haem groups to perform catalytic events, and for example, 12 iron-sulphur clusters and seven haem groups are required for the mitochondrial respiratory chain.

Iron is present in two oxidative forms, ferrous Fe(II) and ferric Fe(III), enabling effective electron transport. Iron that is not bound into iron-sulphur clusters or haem groups is termed the theoretical LIP. Free ferrous iron can be extremely reactive, producing hydroxyl radicals and leading to oxidative damage. Therefore, there exists tight control over iron metabolism within the cell, whereby entry, exit, redox state and total iron content are tightly controlled (for review, see Rouault [4]). Haem consists of an iron atom at the centre of an organic porphyrin ring and has a distinct cellular metabolic pathway.

Two forms of NBIA are a result of mutations to proteins directly involved in cellular iron homeostasis, suggesting that iron metabolism defects might be causative for NBIA. Importantly, the GP and the SN are naturally rich in iron, and iron content increases with age [1,2]. This may predispose these brain regions to iron-induced damage in NBIA disorders.

Cp and aceruloplasminaemia

Cp is involved in cellular iron export. Cp is a glycoprotein containing many copper atoms (and potentially iron atoms) that acts as a ferroxidase to facilitate ferroportin-mediated cellular iron export (Figure 1). Oxidation of Fe(II) to Fe(III) enables binding to transferrin for transport in the extracellular environment. Aceruloplasminaemia, leading to brain iron accumulation, has been shown to be associated with a lack of ferroxidase activity (reviewed [123]); however, mutant Cp protein accumulation has also been described [124].

Disease is thought to progress as a lack of Cp-based ferroxidase activity leads to reduced cellular export of iron as well as an accumulation of extracellular transferrin-free Fe(II) [125]. Astrocytic sequestration of nontransferrin bound extracellular iron could lead to a dearth of iron available for neurones and subsequent cell death [126]. Importantly, other cell types such as astrocytes contain an alternative export-associated ferroxidase, Hephaestin, and may explain specific regional degeneration in aceruloplasminaemia [127].

Aceruloplasminaemia patients display increased serum nontransferrin-bound iron, MRI-based evidence of iron accumulation in the basal ganglia, movement disorders and dementia. Pathology includes iron deposition in neurones and glia within the basal ganglia and dentate nucleus (Table 2) [26–28]. The retina and cerebellum may also show iron accumulation as well as the cortex in late stage disease. Astrocytes show spheroid-like globular structures and iron-rich inclusions in their processes [26]. Oxidative damage was shown to be increased in patient tissue, possibly due to increased reactive ferrous iron [128].

FTL and neuroferritinopathy

Ferritin is the major storage protein for cellular iron (Figure 1). Twenty-four monomers consisting of the heavy and light chain subunits form a proteinaceous shell that stores iron. Ferritin can store up to 4500 iron atoms. The heavy chain has a ferroxidase activity, and the light chain aids mineralization within the ferritin structure. Mutations that affect the ferritin light chain (FTL) lead to the autosomal dominant disease neuroferritinopathy. The most common insertion mutation may lead to a poisoning of the holo-ferritin structure, potentially at the iron entry pore [129], leading to an iron-porous ferritin structure [130].

Iron deposition has been described throughout the basal ganglia from MRI studies [131]. Importantly, evidence of iron deposition has been seen in presymptomatic familial carriers of the disease, leading to the hypothesis that iron accumulation begins in childhood and worsens until symptoms begin, in the fourth decade of life [132]. Pathological investigations have shown ferritin inclusions are present in glia and neurones, as well as confirming iron deposition [19,30,31]. Neurones and glia exhibit Ferritin inclusions in the putamen, GP, thalamus and cerebellum. In the cortex, inclusion-positive glia were perivascular and perineural. Aggregates stain positive for ferritin heavy chain, FTL, iron and ubiquitin. Neuroaxonal swellings have been described, displaying ubiquitin reactivity.

Mitochondrial abnormalities have been highlighted, and an increased oxidative stress of the cells, possibly due to iron, was shown via peroxidation and nitrosylation [31]. Indeed, several animal and cell systems have
confirmed an increase in oxidative stress in FTL mutant models: through mitochondrial and nuclear DNA damage, proteasomal insufficiencies, and damage to proteins and lipid via reactive species [133–135]. Iron chelators were able to reverse cell sensitivity, promoting iron as the main cause of disease [133].

Unifying theories

Despite the fact that mutations in a diverse range of genes lead to NBIA, there are several emerging themes that hint of a mechanistic overlap across the NBIA spectrum: mitochondrial involvement, ROS damage, lipid metabolism, iron and autophagy.

Iron

One major question that remains unanswered is whether iron accumulation is causative or symptomatic of NBIA. Iron accumulation is not specific to NBIA but is observed in a range of neurodegenerative disorders including AD, HD and PD. This excess iron leads to increased oxidative stress and neuronal toxicity.

Cp and FTL have a direct role in the cellular iron homeostatic pathway. It seems rational that mutations in these genes will lead to iron accumulation: for example, mutant Cp leads to increased transferrin-free iron and uncontrolled cellular uptake [125].

Other NBIA genes seem more obscure with regard to iron homeostasis, but similar neuropathology and gross clinical symptoms could argue that mechanisms are conserved between all NBIA subtypes. Contrary to this is the argument that single gene defects such as PLA2G6 lead to distinct diseases with variable iron accumulation. Indeed, iron chelation therapy was able to reduce iron deposition on MRI in PKAN patients but did not lead to clinical benefit [136], suggesting that iron accumulation is not causative of symptoms in NBIA. One explanation for this is that iron accumulation could be indirectly associated with disease. Divalent metal ions have similar properties in cells, and iron dysregulation may affect the steady state of other metal ions – such as zinc or copper leading to neurodegeneration [137].

Iron dyshomeostasis could be explained from a faulty mitochondrial component, as many NBIA-related genes implicate mitochondria. Mitochondria can act as a cellular iron sink with a dedicated mitochondrial ferritin and a series of enzymes requiring iron and haem for function. However, there has been no discovery of a mitochondrial iron exporter. Therefore, it appears the major mechanism for iron recycling is via mitophagy and lysosomal degradation of iron-containing proteins (Figure 2) [138]. Indeed, techniques for staining iron primarily depict iron in lysosomal compartments [139]. Altered mitochondrial fitness may therefore alter mitophagic rates and so iron turnover.

Lipid metabolism

Several of the NBIA genes suggest altered lipid metabolism as a potential disease mechanism: PANK2, COASY, PLA2G6, FA2H and potentially C19orf12. Altered lipid metabolism would affect synthesis and remodelling of lipid bilayers, and Kotzbauer et al. hypothesized a mitochondrial specific fatty acid synthase pathway, which could employ these genes for membrane remodelling and mitochondrial function [37].

Distortion of mitochondrial cristae structure is evident in some NBIA mutant conditions, and it is has been theorized that respiratory chain super structures might be disrupted as a result of lipid insufficiencies [41]. Mitochondrial membrane abnormalities could explain the respiratory deficiencies and ROS damage seen in many NBIA subtypes, and gradually lead to neuronal death.

Lipid metabolism may have a central role in myelin production, and alterations could lead to neurodegeneration. FAH2 is involved in ceramide production, and both PANK2 and COASY are required for sphingomyelin production, demonstrated as PANK2 mutation carriers have decreased sphingosine and cholesterol [50], two critical components of myelin.

Autophagy and mitophagy

Lipid metabolism can be linked with autophagy-associated WDR45 and ATP13A2 via the endosome-autophagolysosome pathways. It is conceivable that genes involved in lipid metabolism are required to construct lipid bilayers for autophagosomal vesicle formation. Mitochondria may represent an intersection of lipid metabolism and autophagy. Not only have mitochondria been shown to donate membranes to autophagic vesicles but also autophagy-associated proteins are transiently seen on mitochondrial outer membranes [103]. Mitochondrial-ER contact sites may be critical for this process [140], although this theory is still controversial [141].
In yeast, mitochondrial health requires one of either mitochondrial-ER connections, or mitochondrial-vacuole connections (the yeast lysosomal orthologue) [142]. These contacts are critical for lipid, calcium and amino acid transfer, yeast health, and metabolic status [143]. This again hints that mitochondria and component recycling are critical for cellular health.

It is tempting to speculate iron-laden autophagolysosomes can also link into an exocytosis pathway (Figure 2). Mitochondria were recently shown to be exocytosed for astrocytic degradation [144]. This could explain neuronal and glial susceptibility for iron deposition in affected brain regions. For example, defective mitochondria (PANK2, COASY, PLA2G6, C19orf12) may be packaged into early autophagosomes via WDR45. ATP13A2 might load autophagic vesicles rich in mitochondria with excess iron from the cytoplasm. These could then be secreted and recycled in neighbouring astrocytes.

Unconventional modes of exocytosis are used in several mammalian systems, for example, interleukin secretion [145], and yeast unconventional exocytosis requires autophagic vesicles and acyl-CoAs: yet another role for CoA [146,147].

**Unfolded protein response**

Tau, α-synuclein, Cp and FTL have all been shown to produce cellular inclusions in NBIA tissue. Taken together with the presence of Ubiquitin in the axonal swellings described in NBIA, the unfolded protein response could be central to cell death in NBIA [148].

Axonal swellings and tubulovesicular structures are another common theme. Components of these pathological hallmarks include mitochondrial components, ubiquitinated proteins and cytoskeletal alterations. Potential mechanisms of cellular disturbance include local sites of oxidative damage, for example, due to axonal transport defects, local mitochondrial ROS production and membrane damage [80,81].

**Energetics**

Finally, the energetic state is implicated throughout NBIA. Mitochondrial health, together with CoA metabolism and lipid homeostasis all point towards the metabolism of fatty acids and amino acids. Insufficiencies of mitochondrial energy production and generation of ROS could point to defective metabolism and subsequent oxidative damage [149]. Acetyl-CoA concentration and COASY protein complexes have recently been shown to be cellular sensors of energy status and can directly bridge metabolic state and autophagic flux through CoA concentration [48,49], further linking PANK2 and COASY to WDR45 and ATP13A2.

Overlap between NBIA disorders and other neurodegenerative disorders is seen via neuropathological evidence of tau and synuclein aggregates, through clinical manifestations and also iron dyshomeostasis. It is tempting to consider shared mechanisms between NBIA, PD, FTD and ALS. The fact that NBIA is often child-onset may highlight specific mechanisms that go awry in late-onset diseases due to a reduction of putative ‘coping mechanisms’ for natural stressors. Therefore, mechanisms linking NBIA neurodegeneration among subtypes may bring to light important insights into neuronal health and ageing.

**Conclusions**

Using rare, penetrant and genetically defined disorders to understand disease mechanisms can help to make more general conclusions. Cellular processes that lead to neurodegeneration may be shared between a range of neurodegenerative diseases, reinforced by similar pathological and clinical findings.

The specific susceptibility of the GP to iron accumulation leading to movement disorders is intriguing due to the ubiquitous expression of the 10 NBIA genes. This may point to a specific high iron content of the GP [2], a reliance on iron-rich glial support cells or due to the tonic activity of the GP, a very high metabolic demand. Retinal cells also have a high metabolic rate and are often affected in NBIA patients.

The involvement of the cerebellum correlates with disease severity, e.g. PLAN and FAHN. Additionally, the early involvement of the SN (e.g. BPAN) might stratify the NBIA genes into Parkinsonian-like and dementia-like categories, helping to explain the Lewy body and tau-positive and negative NBIA subtypes.

Future experiments will explain why cellular coping mechanisms are devoid in NBIA mutations, and these perturbations will be of huge interest to more common neurodegenerative studies.

**Acknowledgements**

The authors would like to acknowledge funding from the Wellcome Trust, the Medical Research Council, the
Mechanisms of neurodegeneration with brain iron accumulation

Dystonia Coalition, the Bachmann-Strauss Dystonia and Parkinson Foundation and Eli Lilly. The authors are funded by the National Institute for Health Research (NIHR) Biomedical Research Unit in Dementia based at University College London Hospitals (UCLH), University College London (UCL). The views expressed are those of the author(s) and not necessarily those of the NHS, the NIHR or the Department of Health.

Author contributions

CA drafted the manuscript. AL, HH and SW provided critical reading and final editing.

References

1 Hill JM, Switzer RC. The regional distribution and cellular localization of iron in the rat brain. Neuroscience 1984; 11: 595–603
2 Hallgren B, Sourander P. The effect of age on the non-phaem iron in the human brain. J Neuroch 1958; 3: 41–51
3 Crichton RR, Wilmet S, Legssyer R, Ward RJ. Molecular and cellular mechanisms of iron homeostasis and toxicity in mammalian cells. J Inorg Biochem 2002; 91: 9–18
4 Rouault TA. Iron metabolism in the CNS: implications for neurodegenerative diseases. Nat Rev Neurosci Nature Publishing Group 2013; 14: 551–64.
5 Andrews NC, Schmidt PJ. Iron homeostasis. Annu Rev Physiol Ann Rev 2007; 69: 69–85
6 Rouault TA. The role of iron regulatory proteins in mammalian iron homeostasis and disease. Nat Chem Biol Nat Publishing Group 2006; 2: 406–14
7 Schneider SA, Hardy J. Bhatia KP. Syndromes of neurodegeneration with brain iron accumulation (NBIA): an update on clinical presentations, histological and genetic underpinnings, and treatment considerations. Mov Disord 2012; 27: 42–53
8 Gregory A, Hayflick Sj. Neurodegeneration with Brain Iron Accumulation Disorders Overview. Gene Reviews. Seattle: University of Washington, 2014
9 Kruer MC, Keogh MJ, Chinnery PF. Current concepts and controversies in neurodegeneration with brain iron accumulation. Semin Pediatr Neurol 2012; 19: 51–6
10 Hayflick Sj, Westaway SK, Levinson B, Zhou B, Johnson MA, Ching KHL, Gitschier J. Genetic, clinical, and radiographic delineation of Hallervorden-Spatz syndrome. N Engl J Med 2003; 348: 33–40
11 Dusi S, Valletta L, Haack TB, Tsuchiya Y, Venco P, Pasqualato S, Goffrini P, Tigano M, Demchenko N, Wieland T, Schwarzmayr T, Strom TM, Invernizzi F, Garavaglia B, Gregory A, Sanford L, Hamada J, Bettencourt C, Houlden H, Chiapparini L, Zorzi G, Kurian MA, Nardocci N, Prokisch H, Hayflick S, Gout I, Tiranti V. Exome sequence reveals mutations in CoA synthase as a cause of neurodegeneration with brain iron accumulation. Am J Hum Genet 2014; 94: 11–22
12 Hartig MB, Iuso A, Haack T, Kmiec T, Jurkiewicz E, Heim K, Roebor S, Tarabin V, Dusi S, Krajewska-Walasek M, Jozwiak S, Hempel M, Winkelmann J, Elstner M, Oexle K, Klopopstock T, Mueller-Felber W, Gasser T, Trenkwalder C, Tiranti V, Kretzschmar H, Schmitz G, Strom TM, Meitinger T, Prokisch H. Absence of an orphan mitochondrial protein, c19orf12, causes a distinct clinical subtype of neurodegeneration with brain iron accumulation. Am J Hum Genet 2011; 89: 543–50
13 Morgan NV, Westaway SK, Morton JEV, Gregory A, Gissen P, Sonek S, Cangul H, Corgyll J, Canham N, Nardocci N, Zorzi G, Pasha S, Rodriguez D, Desguerre I, Mubaidin A, Bertini E, Trembath RC, Simonati A, Schanen C, Johnson CA, Levinson B, Woods CG, Wilmot B, Kramer P, Gitschier J, Mahg ER, Hayflick Sj, PLA2G6, encoding a phospholipase A2, is mutated in neurodegenerative disorders with high brain iron. Nat Genet 2006; 38: 752–4
14 Kruer MC, Paisan-Ruiz C, Boddarett N, Yoon MY, Hama H, Gregory A, Malandrini A, Wolter RL, Munnich A, Gobin S, Polster BJ, Palmeri S, Edmundson S, Hardy J, Houlden H, Hayflick Sj. Defective FA2H leads to a novel form of neurodegeneration with brain iron accumulation (NBIA). Anu Neurol 2010; 68: 611–18
15 Hayflick Sj, Krue MC, Gregory A, Haack TB, Kurian MA, Houlden HH, Anderson J, Boddarett N, Sanford L, Harik SI, Danduy VH, Nardocci N, Zorzi G, Dunaway T, Tarnopoltsky M, Skinner S, Holden KR, Frucht S, Hanspal E, Schrandt-Stumpel C, Mignot C, Héron D, Saunders DE, Kaminska M, Pin J-P, Lascelles K, Cuno SM, Meyer E, Garavaglia B, Bhatia K, de Silva R, Crisp S, Lunt P, Carey M, Hardy J, Meitinger T, Prokisch H, Hogarth P. β-Propeller protein-associated neurodegeneration: a new X-linked dominant disorder with brain iron accumulation. Brain 2013; 136 (Pt 6): 1708–17
16 Schneider SA, Paison-Ruiz C, Quinn NP, Lees AJ, Houlden H, Hardy J, Bhatia KP. ATP13A2 mutations (PARK9) cause neurodegeneration with brain iron accumulation. Mov Disord 2010; 25: 979–84
17 Alazami AM, Al-Saif A, Al-Semari A, Bohlela S, Zlitni S, Alzahrani F, Bavi P, Kaya N, Colak D, Khalak H, Baltus A, Peterlin B, Danda S, Bhatia KP, Schneider SA, Sakati N, Walsh CA, Al-Mohanna F, Meyer B, Alkuraya FS. Mutations in C2orf37, encoding a nucleolar protein, c19orf12, cause hypogonadism, alopecia, diabetes mellitus, mental retardation, and extrapyramidal syndrome. Am J Hum Genet 2008; 83: 684–91
18 Yoshioka K, Furuhata K, Takeda S, Nakamura A, Yamamoto K, Morita H, Shimizu S, Ikeda S, Shimizu N, Yanagisawa N. A mutation in the ceruloplasmin © 2015 The Authors. Neuropathology and Applied Neurobiology published by John Wiley & Sons Ltd on behalf of British Neuropathological Society.
gene is associated with systemic hemosiderosis in humans. Nat Genet 1995; 9: 267–72
19 Curtis AR, Fey C, Morris CM, Bindoff LA, Ince PG, Chinnery PF, Coullard A, Jackson MJ, Jackson AP, McHale DP, Hay D, Barker WA, Markham AF, Bates D, Curtis A, Burn J. Mutation in the gene encoding ferritin light polypeptide causes dominant adult-onset basal ganglia disease. Nat Genet 2001; 28: 350–4
20 Krueer MC, Hiken M, Gregory A, Malandrini A, Clark D, Hogarth P, Grafe M, Hayflick SJ, Woltjer RL. Novel histopathologic findings in molecularly-confirmed pantothenate kinase-associated neurodegeneration. Brain 2011; 134 (Pt 4): 947–58
21 Li A, Paudel R, Johnson R, Courtney R, Lees AJ, Holton JL, Hardy J, Revesz T, Houlden H. Pantothenate kinase-associated neurodegeneration is not a synucleinopathy. Neuropathol Appl Neurobiol 2012; 39 (2): 121–31
22 Hogarth P, Gregory A, Krueer MC, Sanford L, Wagoner W, Natowicz MR, Egel RT, Subramony SH, Goldman JG, Berry-Kravis E, Foulds NC, Hammans SR, Desguerre I, Rodriguez D, Wilson C, Diedrich A, Green A, Tran H, Reese L, Woltjer RL, Hayflick SJ. New NBIA subtype: genetic, clinical, pathologic, and radiographic features of MPAN. Neurology 2013; 80: 268–75
23 Gregory A, Westaway SK, Holm IE, Kotzbauer PT, Hogarth P, Sonek S, Coryell FC, Nguyen TM, Nardocci N, Zorzi G, Rodriguez D, Desguerre I, Bertini E, Simonati A, Levinson B, Dias C, Barbot C, Carrilho I, Santos M, Malik I, Gitschier J, Hayflick SJ. Neurodegeneration associated with genetic defects in phospholipase A2 (A2). Neurology 2008; 71: 1402–9
24 Riku Y, Ikuchi T, Yoshino H, Mimuro M, Mano K, Goto Y, Hattori N, Sobue G, Yoshida M. Extensive aggregation of α-synuclein and tau in juvenile-onset neuroaxonal dystrophy: an autopsied individual with a novel mutation in the PLA2G6 gene-sPLICing site. Acta Neuropathol Commun. BioMed Central Ltd; 2013; 1: 12
25 Paisán-Ruiz C, Li A, Schneider SA, Holton JL, Johnson R, Kidd D, Chataway J, Bhatia KP, Lees AJ, Hardy J, Revesz T, Houlden H. Widespread Lewy body and tau accumulation in childhood and adult onset dystonia-parkinsonism cases with PLA2G6 mutations. Neurobiol Aging 2012; 33: 814–23
26 Gonzalez-Cuyar LF, Perry G, Miyajima H, Atwood CS, Riveros-Angel M, Lyons PF, Siedlak SL, Smith MA, Castellani RJ, Redox active iron accumulation in aceruloplasminemia. Neuropathology 2008; 28: 466–71
27 Kaneko K, Hineno A, Yoshida K, Ohara S, Morita H, Ikeda S. Extensive brain pathology in a patient with aceruloplasminemia with a prolonged duration of illness. Hum Pathol 2012; 43: 451–6
28 Morita H, Ikeda S, Yamamoto K, Morita S, Yoshida K, Nomoto S, Kato M, Yanagisawa N. Hereditary ceruloplasmin deficiency with hemosiderosis: a clinicopathological study of a Japanese family. Ann Neurol 1995; 37: 646–56
29 Oide T, Yoshida K, Kaneko K, Ohta M, Arima K. Iron overload and antioxidative role of perivascular astrocytes in aceruloplasminemia. Neuropathol Appl Neurobiol 2006; 32: 170–6
30 Vidal RP, Ghetti BM, Takao MM, Brefel-Courbon CM, Uro-Coste EM, Glazier BS, Siana VM, Benson MDM, Calvas PM, Miravalle LP, Rascol O, Delisle MBM. Intracellular ferritin accumulation in neural and extraneural tissue characterizes a neurodegenerative disease associated with a mutation in the ferritin light polypeptide gene. J Neuropath Exp Neurol 2004; 63: 363–80
31 Mancuso MM, Davidzon GM, Kurlan RMM, Tawil RM, Bonilla EM, Di Mauro SM, Powers JMM. Hereditary ferritinopathy: a novel mutation, its cellular pathology, and pathogenetic insights. J Neuropath Exp Neurol 2005; 64: 280–94
32 Zhou B, Westaway SK, Levinson B, Johnson MA, Gitschier J, Hayflick SJ. A novel pantothenate kinase gene (PANK2) is defective in Hallervorden-Spatz syndrome. Nat Genet 2001; 28: 345–9
33 Zhang Y-M, Rock CO. Jackowski S. Biochemical properties of human pantothenate kinase 2 isoforms and mutations linked to pantothenate kinase-associated neurodegeneration. J Biol Chem 2006; 281: 107–14
34 Hong BS, Senisterra G, Rabeh WM, Vedadi M, Leonardi R, Zhang Y-M, Rock CO, Jackowski S, Park H-W. Crystal structures of human pantothenate kinases. Insights into allosteric regulation and mutations linked to a neurodegeneration disorder. J Biol Chem 2007; 282: 27984–93
35 Hartig MB, Hörtngael K, Garavaglia B, Zorzi G, Kmiec T, Klopstock T, Rostasy K, Svetel M, Kostic VS, Schuelke M, Botz E, Weindl A, Novakovic I, Nardocci N, Prokisch H, Mettinger T. Genotypic and phenotypic spectrum of PANK2 mutations in patients with neurodegeneration with brain iron accumulation. Ann Neurol 2006; 59: 248–56
36 Hortnagel K. An isoform of hPANK2, deficient in pantothenate kinase-associated neurodegeneration, localizes to mitochondria. Hum Mol Genet 2003; 12: 321–7
37 Kotzbauer PT, Truax AC, Trojanowski JQ, Lee VM-Y. Altered neuronal mitochondrial coenzyme A synthesis in neurodegeneration with brain iron accumulation caused by abnormal processing, stability, and catalytic activity of mutant pantothenate kinase 2. J Neurosci 2005; 25: 689–98
38 Perry TL, Norman MG, Yong VW, Whiting S, Crichton JU, Hansen S, Kish SJ. Hallervorden-Spatz disease: cysteine accumulation and cysteine dioxygenase deficiency in the globus pallidus. Ann Neurol 1985; 18: 482–9
39 Campanella A, Privitera D, Guaraldo M, Rovelli E, Barzaghi C, Garavaglia B, Santambrogio P, Cozzi A, Levi S. Skin fibroblasts from pantothenate kinase-associated
neurodegeneration patients show altered cellular oxidative status and have defective iron-handling properties. *Hum Mol Genet* 2012; 21: 4049–59

40 Bosveld F, Rana A, van der Wouden PE, Lemstra W, Ritsema M, Kampinga HH, Sibon OCM. De novo CoA biosynthesis is required to maintain DNA integrity during development of the Drosophila nervous system. *Hum Mol Genet* 2008; 17: 2058–69

41 Brunetti D, Dusi S, Morbin M, Uggetti A, Moda F, Amato D, Giordano I, d C, Amati G, Cozzi A, Levi S, Hayflick S, Tiranti V. Pantothenate kinase-associated neurodegeneration: altered mitochondria membrane potential and defective respiration in Pank2 knock-out mouse model. *Hum Mol Genet* 2012; 21: 5294–305

42 Rana A, Seinen E, Sludeja K, Muntendam R, Srinivasan B, van der Want JJ, Hayflick S, Reijnoudt D-J, Kayser O, Sibon OCM. Pantethine rescues a Drosophila model for pantothenate kinase-associated neurodegeneration. *Proc Natl Acad Sci USA* 2010; 107: 6988–93

43 Deas E, Wood NW, Plun-Favreau H. Mitophagy and Parkinson’s disease: the PINK1-parkin link. *Biochim Biophys Acta* 2011; 1813: 623–33

44 Chen H, Chan DC. Mitochondrial dynamics – fusion, fission, movement, and mitophagy – in neurodegenerative diseases. *Hum Mol Genet*. Oxford University Press 2009; 18 (R2): R169–76

45 Wu Z, Li C, Lv S, Zhour B. Pantothenate kinase-associated neurodegeneration: insights from a Drosophila model. *Hum Mol Genet* 2009; 18: 3659–72

46 Leonard R, Rock CO, Jackowski S, Zhang Y-M. Activation of human mitochondrial pantothenate kinase 2 by palmitoylcarnitine. *Proc Natl Acad Sci USA* 2007; 104: 1494–9

47 Sludeja K, Srinivasan B, Xu L, Rana A, de Jong J, Nollen EAA, Jackowski S, Sanford L, Hayflick S, Sibon OCM. Impaired Coenzyme A metabolism affects histone and tubulin acetylation in Drosophila and human cell models of pantothenate kinase associated neurodegeneration. *EMBO Mol Med* 2011; 3: 755–66

48 Breus O, Panasyuk G, Gout IT, Filonenko V, Nemazanyy I. CoA synthase is in complex with p85alphaPI3K and affects PI3K signaling pathway. *Biochem Biophys Res Commun* 2009; 385: 581–5

49 Mariño G, Pietroccola F, Eisenberg T, Kong Y, Malik SA, Andryushkova A, Schroeder T, Morbidelli M, Niso-Santano M, Nocenzi M, Durand S, Enot DP, Fernández AF, Martins I, Kepp O, Senovilla L, Bricceno K V, Rinaldi C, Meilleur KG, Sangaré M, Diallo WA, Traynor BJ, Marques W, Züchner S, Blackstone C, Fischbeck KH, Johnson JO, Houthen H, Johnston J, Mooney C, Alzahrani J, Elmaliak E, Van der Want JJ, Hayflick S, van der Want JJ, Houthen H, Azzedine C, Lalso-button K, Van der Want JJ, Hayflick S, Van der Want JJ, Houthen H, Azzedine C, Lalso-button K, Van der Want JJ, Hayflick S, van der Want JJ, Houthen H, Azzedine C, Lalso-button K, Van der Want JJ, Hayflick S, van der Want JJ, Houthen H, Azzedine C, Lalso-button K, Van der Want JJ, Hayflick S, van der Want JJ, Houthen H, Azzedine C, Lalso-button K, Van der Want JJ, Hayflick S, van der Want JJ, Houthen H, Azzedine C, Lalso-button K, Van der Want JJ, Hayflick S, van der Want JJ, Houthen H, Azzedine C, Lalso-button K, Van der Want JJ, Hayflick S, van der Want JJ, Houthen H, Azzedine C, Lalso-button K, Van der Want JJ, Hayflick S, van der Want JJ, Houthen H, Azzedine C, Lalso-button K, Van der Want JJ, Hayflick S, van der Want JJ, Houthen H, Azzedine C, Lalso-button K, Van der Want JJ, Hayflick S, van der Want JJ, Houthen H, Azzedine C, Lalso-button K, Van der Want JJ, Hayflick S, van der Want JJ, Houthen H, Azzedine C, Lalso-button K, Van der Want JJ, Hayflick S, van der Want JJ, Houthen H, Azzedine C, Lalso-button K, Van der Want JJ, Hayflick S, van der Want JJ, Houthen H, Azzedine C, Lalso-button K, Van der Want JJ, Hayflick S, van der Want JJ, Houthen H, Azzedine C, Lalso-button K, Van der Want JJ, Hayflick S, van der Want JJ, Houthen H, Azzedine C, Lalso-button K, Van der Want JJ, Hayflick S, van der Want JJ, Houthen H, Azzedine C, Lalso-button K, Van der Want JJ, Hayflick S, van der Want JJ, Houthen H, Azzedine C, Lalso-button K, Van der Want JJ, Hayflick S, van der Want JJ, Houthen H, Azzedine C, Lalso-button K, Van der Want JJ, Hayflick S, van der Want JJ, Houthen H, Azzedine C, Lalso-button K, Van der Want JJ, Hayflick S, van der Want JJ, Houthen H, Azzedine C, Lalso-button K, Van der Want JJ, Hayflick S, van der Want JJ, Houthen H, Azzedine C, Lalso-button K, Van der Want JJ, Hayflick S, van der Want JJ, Houthen H, Azzedine C, Lalso-button K, Van der Want JJ, Hayflick S, van der Want JJ, Houthen H, Azzedine C, Lalso-button K, Van der Want JJ, Hayflick S, van der Want JJ, Houthen H, Azzedine C, Lalso-button K, Van der Want JJ, Hayflick S, van der Want JJ, Houthen H, Azzedine C, Lalso-button K, Van der Want JJ, Hayflick S, van der Want JJ, Houthen H, Azzedine C, Lalso-button K, Van der Want JJ, Hayflick S, van der Want JJ, Houthen H, Azzedine C, Lalso-button K, Van der Want JJ, Hayflick S, van der Want JJ, Houthen H, Azzedine C, Lalso-button K, Van der Want JJ, Hayflick S, van der Want JJ, Houthen H, Azzedine C, Lalso-button K, Van der Want JJ, Hayflick S, van der Want JJ, Houthen H, Azzedine C, Lalso-button K, Van der Want JJ, Hayflick S, van der Want JJ, Houthen H, Azzedine C, Lalso-button K, Van der Want JJ, Hayflick S, van der Want JJ, Houthen H, Azzedine C, Lalso-button K, Van der Want JJ, Hayflick S, van der Want JJ, Houthen H, Azzedine C, Lalso-button K, Van der Want JJ, Hayflick S, van der Want JJ, Houthen H, Azzedine C, Lalso-button K, Van der Want JJ, Hayflick S, van der Want JJ, Houthen H, Azzedine C, Lalso-button K, Van der Want JJ, Hayflick S, van der Want JJ, Houthen H, Azzedine C, Lalso-button K, Van der Want JJ, Hayflick S, van der Want JJ, Houthen H, Azzedine C, Lalso-button K, Van der Want JJ, Hayflick S, van der Want JJ, Houthen H, Azzedine C, Lalso-button K, Van der Want JJ, Hayflick S, van der Want JJ, Houthen H, Azzedine C, Lalso-button K, Van der Want JJ, Hayflick S, van der Want JJ, Houthen H, Azzedine C, Lalso-button K, Van der Want JJ, Hayflick S, van der Want JJ, Houthen H, Azzedine C, Lalso-button K, Van der Want JJ, Hayflick S, van der Want JJ, Houthen H, Azzedine C, Lalso-button K, Van der Want JJ, Hayflick S, van der Want JJ, Houthen H, Azzedine C, Lalso-button K, Van der Want JJ, Hayflick S, van der Want JJ, Houthen H, Azzedine C, Lalso-button K, Van der Want JJ, Hayflick S, van der Want JJ, Houthen H, Azzedine C, Lalso-button K, Van der Want JJ, Hayflick S, van der Want JJ, Houthen H, Azzedine C, Lalso-button K, Van der Want JJ, Hayflick S, van der Want JJ, Houthen H, Azzedine C, Lalso-button K, Van der Want JJ, Hayflick S, van der Want JJ, Houthen H, Azzedine C, Lalso-button K, Van de...
phospholipase A2 in activated cells. Cell Signal 2005; 17: 1052–62.
62 Yang HC, Mosior M, Johnson CA, Chen Y, Dennis EA. Group-specific assays that distinguish between the four major types of mammalian phospholipase A2. Ann Biochem 1999; 269: 278–88.
63 Larsson PKA, Claesson H-E, Kennedy BP. Multiple splice variants of the human calcium-independent phospholipase A2 and their effect on enzyme activity. J Biol Chem 1998; 273: 207–14.
64 Liu J-Y, Aleksic N, Chen S-F, Han T-J, Shyue S-K, Wu KK. Mitochondrial localization of cyclooxygenase-2 and calcium-independent phospholipase A2 in human cancer cells: implication in apoptosis resistance. Exp Cell Res 2005; 306: 75–84.
65 Williams SD, Gottlieb RA. Inhibition of mitochondrial calcium-independent phospholipase A2 (iPLA2) attenuates mitochondrial phospholipid loss and is cardioprotective. Biochem J Portland Press Ltd. 2002; 362 (Pt 1): 23–32.
66 Selezniev K, Zhao C, Zhang XH, Song K, Ma ZA. Calcium-independent phospholipase A2 localizes in and protects mitochondria during apoptotic induction by staurosporine. J Biol Chem 2006; 281: 22275–88.
67 Ong W-Y, Yeo J-F, Ling S-F, Faroqui AA. Distribution of calcium-independent phospholipase A2 (iPLA2) in monkey brain. J Neurocytol 2005; 34: 447–58.
68 Song H, Bao S, Lei X, Jin C, Zhang S, Turk J, Ramanadham S. Evidence for proteolytic processing and stimulated organelle redistribution of iPLA2(2)beta. Biochim Biophys Acta Elsevier B.V., 2010; 1801 (5): 547–58.
69 Engel LA, Jing Z, O’Brien DE, Sun M, Kotzbauer PT. Catalytic function of PLA2G6 is impaired by mutations associated with infantile neuroaxonal dystrophy but not dystonia-parkinsonism. PLoS ONE 2010; 5: e12897.
70 Jenkins CM, Yan W, Mancuso DJ, Gross RW. Highly selective hydrolysis of fatty acyl-CoA by calcium-independent phospholipase A2beta. Enzyme auto-acylation and acyl-CoA-mediated reversal of calcium-independent phospholipase A2 activity. J Biol Chem 2006; 281: 15615–24.
71 Paisan-Ruiz C, Bhattia KP, Li A, Hernandez D, Davis M, Wood NW, Hardy J, Houlden H, Singleton A, Schneider SA. Characterization of PLA2G6 as a locus for dystonia-parkinsonism. Ann Neurol 2009; 65: 19–23.
72 Shinzawa K, Sumi H, Ikawa M, Matsuoka Y, Okabe M, Sakoda S, Tsujimoto Y. Neuroaxonal dystrophy caused by group VIA phospholipase A2 deficiency in mice: a model of human neurodegenerative disease. J Neurosci 2008; 28: 2212–20.
73 Malik I, Turk J, Mancuso DJ, Montier L, Wohltmann M, Wozniak DF, Schmidt RE, Gross RW, Kotzbauer PT. Disrupted membrane homeostasis and accumulation of ubiquitinated proteins in a mouse model of infantile neuroaxonal dystrophy caused by PLA2G6 mutations. Am J Pathol. American Society for Investigative Pathology 2008; 172: 406–16.
74 Balboa MA, Balsinde J. Involvement of calcium-independent phospholipase A2 in hydrogen peroxide-induced accumulation of free fatty acids in human U937 cells. J Biol Chem 2002; 277: 40384–9.
75 Itoh K, Negishi H, Obayashi C, Hayashi Y, Hanika K, Imai Y, Itoh H. Infantile neuroaxonal dystrophy – immunohistochemical and ultrastructural studies on the central and peripheral nervous systems in infantile neuroaxonal dystrophy. Kobe J Med Sci 1993; 39: 133–46.
76 Mahadevan A, Santosh V, Gayatri N, Ratnavalli E, NandanGopal R, Vasanth A, Roy AK, Shankar SK. Infantile neuroaxonal dystrophy and giant axonal neuropathy – overlap diseases of neuronal cytoskeletal elements in childhood? Clin Neuropathol 2000; 19: 221–9.
77 Broekemeier KM, Iben JR, LeVan EG, Crouser ED, Pfeiffer DR. Pore Formation and Uncoupling Initiate a Ca2+-Independent Degradation of Mitochondrial Phospholipids†. Biochemistry. American Chemical Society 2002; 41: 7771–80.
78 Gadd ME, Broekemeier KM, Crouser ED, Kumar J, Graff G, Pfeiffer DR. Mitochondrial iPLA2 activity modulates the release of cytochrome c from mitochondria and influences the permeability transition. J Biol Chem 2006; 281: 6931–9.
79 Lei X, Zhang S, Bohrer A, Barbour SE, Ramanadham S. Role of calcium-independent phospholipase A2(2)beta in human pancreatic islet β-cell apoptosis. Am J Physiol Endocrinol Metab 2012; 303: E1386–95.
80 Beck G, Sugiuira Y, Shiznawa K, Kato S, Setou M, Tsujimoto Y, Sakoda S, Sumi-Akamaru H. Neuroaxonal dystrophy in calcium-independent phospholipase A2β deficiency results from insufficient remodeling and degeneration of mitochondrial and presynaptic membranes. J Neurosci 2011; 31: 11411–20.
81 Roy S, Zhang B, Lee VM-Y, Trojanowski JQ. Axonal transport defects: a common theme in neurodegenerative diseases. Acta Neuropathol 2005; 109: 5–13.
82 Green JT, Orr SK, Bazinet RP. The emerging role of group VI calcium-independent phospholipase A2 in releasing docosahexaenoic acid from brain phospholipids. J Lipid Res 2008; 49: 939–44.
83 Strokin M, Sergeeva M, Reiser G. Prostaglandin synthesis in rat brain astrocytes is under the control of the n-3 docosahexaenoic acid, released by group VII calcium-independent phospholipase A2. J Neurochem 2007; 102: 1771–82.
84 Kondo M, Shibata T, Kumatagi T, Osawa T, Shibata N, Kobayashi M, Sasaki S, Iwata M, Noguchi N, Uchida K. 15-Deoxy-Delta(12,14)-prostaglandin J(2): the endogenous electrophile that induces neuronal apoptosis. Proc Natl Acad Sci U S A 2002; 99: 7367–72.
85 Strokin M, Seburn KL, Cox GA, Martens KA, Reiser G. Severe disturbance in the Ca2+ signaling in astrocytes.
from mouse models of human infantile neuroaxonal dystrophy with mutated Pld2g6. *Hum Mol Genet* 2012; 21: 2807–14

86 Ross BM, Moszcynska A, Erlich J, Kish SJ. Phospholipid-metabolizing enzymes in Alzheimer’s disease: increased lysophospholipid acyltransferase activity and decreased phospholipase A2 activity. *J Neurochem* 2002; 70: 786–93

87 Ross BM, Moszcynska A, Erlich J, Kish SJ. Low activity of key phospholipid catabolic and anabolic enzymes in human substantia nigra: possible implications for Parkinson’s disease. *Neuroscience* 1998; 83: 791–8

88 Edvardsson S, Hama H, Shaag A, Gomori JM, Berger I, Soffer D, Korman SH, Tautein I, Saada A, Elpeleg O. Mutations in the fatty acid 2-hydroxylase gene are associated with leukodystrophy with spastic paraparesis and dystonia. *Am J Hum Genet* 2008; 83: 643–8

89 Dick KJ, Eckhardt M, Paisan-Ruiz C, Alshehhi AA, Proukakis C, Sibtain NA, Maier H, Crosby AH. Mutation of the FA2H gene underlies a complex form of hereditary spastic paraplegia (SPG35). *Hum Mutat* 2010; 31: E1251–60

90 Eckhardt M, Yaghoofifam A, Fewou SN, Zöller I, Gieselmann V. A mammalian fatty acid hydroxylase responsible for the formation of alpha-hydroxylated galactosylceramide in myelin. *Biochem J* 2005; 388 (Pt 1): 245–54

91 Alderson NL, Rembiesa BM, Walla MD, Bielawska A, Bielawski J, Hama H. The human FA2H gene encodes a fatty acid 2-hydroxylase. *J Biol Chem* 2004; 279: 48562–8

92 Hama H. Fatty acid 2-Hydroxylation in mammalian sphingolipid biology. *Biochim Biophys Acta* 2010; 1801: 405–14

93 Potter KA, Kern MJ, Fullbright G, Bielawski J, Scherer SS, Yum SW, Li JJ, Cheng H, Han X, Venkata JK, Khan PAA, Rohrer B, Hama H. Central nervous system dysfunction in a mouse model of FA2H deficiency. *Glia* 2011; 59: 1009–21

94 Bras J, Singleton A, Cookson MR, Hardy J. Emerging pathways in genetic Parkinson’s disease: potential role of ceramide metabolism in Lewy body disease. *FEBS J* 2008; 275: 5767–73

95 Jana A, Hogan EL, Pahan K. Ceramide and neurodegeneration: susceptibility of neurons and oligodendrocytes to cell damage and death. *J Neurosci* 2009; 29(20): 515

96 Fukunaga M, Li T-Q, van Gelderen P, de Zwart JA, Shmueli K, Yao B, Lee J, Marie D, Aironova MA, Zhang G, Leapman RD, Schenck JF, Merkle H, Duyn JH. Layerspecific variation of iron content in cerebral cortex as a source of MRI contrast. *Proc Natl Acad Sci U S A* 2010; 107: 3834–9

97 Haack TB, Hogarth P, Krueer MC, Gregory A, Wieland T, Schwarzmuay T, Graf E, Sanford L, Meyer E, Kara E, Cuno SM, Harik SI, Danuaw VH, Nardoci N, Zorzzi G, Dunaway T, Tarnopolsky M, Skinner S, Frucht S, Hanspal E, Schrander-Stumpel C, Héron D, Mignot C, Garavaglia B, Bhatia K, Hardy J, Strom TM, Bodaert N, Houlden HH, Kurian MA, Meitinger T, Prokisch H, Hayflick SJ. Exome sequencing reveals de novo WDR45 mutations causing a phenotypically distinct, X-linked dominant form of NBIA. *Am J Hum Genet* 2012; 91: 1144–9

98 Lu Q, Yang P, Huang X, Hu W, Guo B, Wu F, Lin L, Kovács AL, Yu L, Zhang H. The WD40 repeat PtdIns(3)P-binding protein EPG-6 regulates progression of omegasomes to autophagosomes. *Dev Cell* 2011; 21: 343–57

99 Dall’Armi C, Devereaux KA, Di Paolo G. The role of lipids in the control of autophagy. *Curr Biol* 2013; 23: R33–45

100 Prokias-Cezanne T, Waddell S, Gaugel A, Fricke T, Lupas A, Nordheim E. WIPI-1alpha (WIPI49), a member of the novel 7-bladed WIPI protein family, is aberrantly expressed in human cancer and is linked to starvation-induced autophagy. *Oncogene*. Nature Publishing Group 2004; 23: 9314–25

101 Tooze SA, Yoshimori T. The origin of the autophagosomal membrane. *Nat Cell Biol* 2010; 12: 831–5

102 Saitsu H, Nishimura T, Muramatsu K, Kodera H, Kumada S, Sugai K, Kasai-Yoshida E, Sawaura N, Nishida H, Hoshino A, Ryujin F, Yoshioka S, Nishiyama K, Kondo Y, Tsurusaki Y, Nakashima M, Miyake N, Arakawa H, Kato M, Mizushima N, Matsumoto N. De novo mutations in the autophagy gene WDR45 cause static encephalopathy of childhood with neurodegeneration in adulthood. *Nat Genet* Nature Publishing Group, a division of Macmillan Publishers Limited. All Rights Reserved 2013; 45: 445–9, 449e1.

103 Hailey DW, Rambold AS, Satpute-Krishnan P, Mitra K, Sougrat R, Kim PK, Lippincott-Schwartz J. Mitochondria supply membranes for autophagosome biogenesis during starvation. *Cell* 2010; 141: 656–67

104 Scherz-Shouval R, Elazar Z. ROS, mitochondria and the regulation of autophagy. *Trends Cell Biol* 2007; 17: 422–7

105 Jiang P, Mizushima N. Autophagy and human diseases. *Cell Res* Shanghai Institutes for Biological Sciences, Chinese Academy of Sciences 2014; 24: 69–79.

106 Di Fonzo A, Chien HF, Socal M, Giraudo S, Tassorelli C, Iliceto G, Fabbrini G, Marconi R, Fincati E, Abbruzzese G, Marini P, Squirrier F, Horstink MW, Montagna P, Libera AD, Stocchi F, Goldwurm S, Ferreira JJ, Meco G, Martignoni E, Lopiano L, Jardim LB, Oostra BA, Barbosa ER, Bonifati V. ATP13A2 missense mutations in juvenile parkinsonism and young onset Parkinson disease. *Neurology* 2007; 68: 1557–62

107 Bras J, Verloes A, Schneider SA, Mole SE, Guerreiro RJ. Mutation of the parkinsonism gene ATP13A2 causes...
neuronal ceroid-lipofuscinosis. *Hum Mol Genet* 2012; 21: 2646–50

108 Ramirez A, Heimbach A, Gründemann J, Stillier B, Hampshire D, Ced LP, Goebel I, Mubaidin AF, Wriekat A-L, Roeppe J, Al-Din A, Hillmer AM, Karsak M, Liss B, Woods CG, Behrens M, Kubisch C. Hereditary parkinsonism with dementia is caused by mutations in ATP13A2, encoding a lysosomal type 5 P-type ATPase. *Nat Genet* 2006; 38: 1184–91

109 Ramonet D, Podjaiska A, Stafa K, Sonnay S, Trancikova A, Tsika E, Pletnikova O, Troncoso JC, Glauers L, Moore DJ. PARK9-associated ATP13A2 localizes to intracellular acidic vesicles and regulates cation homeostasis and neuronal integrity. *Hum Mol Genet* 2012; 21: 1725–43

110 Dehay B, Ramirez A, Martinez-Vicente M, Perier C, Canron M-H, Doudnikoff E, Vital A, Vila M, Klein C, Bezard E. Loss of P-type ATPase ATP13A2/PARK9 function induces general lysosomal deficiency and leads to Parkinson disease neurodegeneration. *Proc Natl Acad Sci USA* 2012; 109: 9611–16

111 Kühlbrandt W. Biology, structure and mechanism of P-type ATPases. *Nat Rev Mol Cell Biol* Nature Publishing Group 2004; 5: 282–95.

112 Tan J, Zhang T, Chi J, Hu D, Pan Q, Wang D, Zhang Z. Regulation of intracellular manganese homeostasis by Kufor-Rakeb syndrome-associated ATP13A2 protein. *J Biol Chem* 2011; 286: 29654–62

113 Gitter AD, Chesi A, Geddie ML, Strathern KE, Hamamichi S, Hill KJ, Caldwell KA, Caldwell GA, Cooper AA, Rochet J-C, Lindquist S. Alpha-synuclein is part of a diverse and highly conserved interaction network that includes PARK9 and manganese toxicity. *Nat Genet*. Nature Publishing Group 2009; 41: 308–15

114 Schmidt K, Wolfe DM, Stilller B, Pearce DA. Cd2+, Mn2+, Ni2+ and Se2+ toxicity to Saccharomyces cerevisiae lacking YPK9p the orthologue of human ATP13A2. *Biochem Biophys Res Commun* 2009; 383: 198–202

115 Usenovic M, Tresse E, Mazzulli JR, Taylor JP, Krainic D. Deficiency of ATP13A2 leads to lysosomal dysfunction, α-synuclein accumulation, and neurotoxicity. *J Neurosci* 2012; 32: 4240–6

116 Tsunemi T, Krainic D. Zn2+ dyshomeostasis caused by loss of ATP13A2/PARK9 leads to lysosomal dysfunction and alpha-synuclein accumulation. *Hum Mol Genet* 2014; 23: 2791–801

117 Park J-S, Koentjoro B, Veivers D, Mackay-Sim A, Sue CM. Parkinson’s disease-associated human ATP13A2 (PARK9) deficiency causes zinc dyshomeostasis and mitochondrial dysfunction. *Hum Mol Genet* 2014; 23: 2802–15

118 Grünewald A, Arns B, Seibler P, Rakovic A, Münchau A, Ramirez A, Sue CM, Klein C. ATP13A2 mutations impair mitochondrial function in fibroblasts from patients with Kufor-Rakeb syndrome. *Neurobiol Aging* 2012; 33: 1843, e1–7

119 Pivtoraiko VN, Stone SL, Roth KA, Shacka JJ. Oxidative stress and autophagy in the regulation of lysosome-dependent neuron death. *Antioxid Redox Signal* 2009; 11: 481–96

120 Polishchuk EV, Concilli M, Iacobacci S, Chesi G, Pastore N, Piccolo P, Paladino D, Baldantoni D, van Ijzendoorn SCD, Chan J, Chang CJ, Amoresano A, Pane F, Pucci P, Tarallo A, Parenti G, Brunetti-Pierri N, Settembre C, Ballabio A, Polishchuk RS. Wilson disease protein ATP7B utilizes lysosomal exocytosis to maintain copper homeostasis. *Dev Cell* Elsevier 2014; 29: 686–700.

121 Nousheek J, Spiegel R, Ishida-Yamamoto A, Indelman M, Shani-Adir A, Adir N, Lipkin E, Bercovici S, Geiger D, van Steensel MA, Steijlen PM, Bergman R, Bindereif A, Choder M, Shaley S, Sprecher E, Allopecia, neurological defects, and endocrinopathy syndrome caused by decreased expression of RBM28, a nuclear protein associated with ribosome biogenesis. *Am J Hum Genet* 2008; 82: 1114–21

122 Jin J, Arias EE, Chen J, Harper JW, Walter JC. A family of diverse Cul4-Ddb1-interacting proteins includes Cdt2, which is required for S phase destruction of the replication factor Cdt1. *Mol Cell* 2006; 23: 709–21

123 Levi S, Finazzi D. Neurodegeneration with brain iron accumulation: update on pathogenic mechanisms. *Front Pharmacol* 2014; 5 (May): 99

124 Kono S, Suzuki H, Oda T, Miyajima H, Takahashi Y, Shirakawa K, Ishikawa K, Kitagawa M. Biochemical features of ceruloplasmin gene mutations linked to ceruloplasminemia. *Neuromolecular Med* 2006; 8: 361–74

125 Brissot P, Ropert M, Le Can C, Léoréal O. Non-transferrin bound iron: a key role in iron overload and iron toxicity. *Biochim Biophys Acta* 2012; 1820: 403–10

126 Jeong SY, David S. Age-related changes in iron homeostasis and cell death in the cerebellum of ceruloplasmin-deficient mice. *J Neurosci* 2006; 26: 9810–19

127 Cui R, Duan X-L, Andersson GJ, Qiao Y-T, Yu P, Qian Z-M, Yoshida K, Takeda S, Guo P, Yang Z-L, Chang Y-Z. Age-dependent expression of hephaestin in the brain of ceruloplasmin-deficient mice. *J Trace Elem Med Biol* 2009; 23: 290–9

128 Yoshida K, Kaneko K, Miyajima H, Tokuhisa T, Nakamura A, Kato M, Ikeda S. Increased lipid peroxidation in the brains of aceruloplasminemia patients. *J Neurol Sci* 2000; 175: 91–5

129 Burn J, Chinnery PF. Neuroferritinopathy. *Semin Pediatr Neurol* 2006; 13: 176–81

130 Friedman A, Arosio P, Finazzi D, Koziorowski D, Galazka-Friedman J. Ferritin as an important player in neurodegeneration. *Parkinsonism Relat Disord* 2011; 17: 423–30

131 McNeill A, Birchall D, Hayflick SJ, Gregory A, Schenk JF, Zimmerman EA, Shang H, Miyajima H, Chinnery PF. T2* and FSE MRI distinguishes four subtypes of neuro-
Mechanisms of neurodegeneration with brain iron accumulation

132 Keogh MJ, Jonas P, Coultard A, Chinnery PF, Burn J. Neuroferritinopathy: a new inborn error of iron metabolism. Neurogenetics 2012; 13: 93–6
133 Cozzi A, Rovelli E, Friziale G, Campanella A, Amendola M, Arosio P, Levi S. Oxidative stress and cell death in cells expressing L-ferritin variants causing neuroferritinopathy. Neurobiol Dis 2010; 37: 77–85
134 Barbeito AG, Garringer HJ, Baraibar MA, Gao X, Arredondo M, Núñez MT, Smith MA, Ghetti B, Vidal R. Abnormal iron metabolism and oxidative stress in mice expressing a mutant form of the ferritin light polypeptide gene. J Neurochem 2009; 109: 1067–78
135 Deng X, Vidal R, Englander EW. Accumulation of oxidative DNA damage in brain mitochondria in mouse model of hereditary ferritinopathy. Neurosci Lett 2010; 479: 44–8
136 Zorzi G, Zibordi F, Chiapparini L, Bertini E, Russo L, Piga A, Longo F, Garavaglia B, Aquino D, Savoiaardo M, Solari A, Nardocci N. Iron-related MRI images in patients with pantothenate kinase-associated neurodegeneration (PKAN) treated with deferiprone: results of a phase II pilot trial. Mov Disord 2011; 26: 1756–9
137 Yadrick M, Kenney M, Winterfeldt E. Iron, copper, and zinc status: response to supplementation with zinc or zinc and iron in adult females. Am J Clin Nutr 1989; 49: 145–50
138 Xu W, Barrientos T, Andrews NC. Iron and copper in mitochondrial diseases. Cell Metab 2013; 17: 319–28
139 Kurz T, Eaton JW, Brunk UT. The role of lysosomes in iron metabolism and recycling. Int J Biochem Cell Biol 2011; 43: 1686–97
140 Hamasaki M, Furuta N, Matsuda A, Nezu A, Yamamoto A, Fujita N, Oomori H, Noda T, Haraguchi T, Hiraoka Y, Amano A, Yoshimori T. Autophagosomes form at ER-mitochondria contact sites. Nature Publishing Group, a division of Macmillan Publishers Limited. All Rights Reserved 2013; 495: 389–93.
141 Maday S, Holzbaur ELF. Autophagosome biogenesis in primary neurons follows an ordered and spatially regulated pathway. Dev Cell 2014; 30: 71–85
142 Elbaz-Alon Y, Rosenfeld-Gur E, Shinder V, Futerer AH, Geiger T, Schuldiner M. A dynamic interface between vacuoles and mitochondria in yeast. Dev Cell 2014; 30: 95–102
143 Hönscher C, Mari M, Auffarth K, Bohnert M, Griffith J, Geerts W, van der Laan M, Cabrera M, Reggiori F, Unger mann C. Cellular metabolism regulates contact sites between vacuoles and mitochondria. Dev Cell 2014; 30: 86–94
144 Davis C-HO, Kim K-Y, Bushong EA, Mills EA, Boassa D, Shih T, Kinebuchi M, Phan S, Zhou Y, Bihlmeyer NA, Nguyen JV, Jin Y, Ellisman MH, Marsh-Armstrong N. Transcellular degradation of axonal mitochondria. Proc Natl Acad Sci USA 2014; 111: 9633–8
145 Keller M, Rüegg A, Werner S, Beer H-D. Active caspase-1 is a regulator of unconventional protein secretion. Cell 2008; 132: 818–31
146 Duran JM, Anjard C, Stefan C, Loomis WF, Malhotra V. Unconventional secretion of Acb1 is mediated by autophagosomes. J Cell Biol 2010; 188: 527–36
147 Manjithaya R, Anjard C, Loomis WF, Subramani S. Unconventional secretion of Pichia pastoris Acb1 is dependent on GRASP protein, peroxisomal functions, and autophagosome formation. J Cell Biol 2010; 188: 537–46
148 Moreno JA, Halliday M, Molloy C, Radford H, Verity N, Axten JM, Ortori CA, Willis AE, Fischer PM, Barrett IA, Mallucci GR. Oral treatment targeting the unfolded protein response prevents neurodegeneration and clinical disease in prion-infected mice. Sci Transl Med. 2013; 5: 206ra138
149 Adibhatla R, Hatcher J. Lipid oxidation and peroxidation in CNS health and disease: from molecular mechanisms to therapeutic opportunities. Antioxid Redox Signal 2010; 12 (1): 125–169

Received 4 November 2014
Accepted after revision 18 March 2015
Published online Article Accepted on 14 April 2015