A novel neuroferritinopathy mouse model (FTL 498InsTC) shows progressive brain iron dysregulation, morphological signs of early neurodegeneration and motor coordination deficits

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A B S T R A C T

Neuroferritinopathy is a rare genetic disease with a dominant autosomal transmission caused by mutations of the ferritin light chain gene (FTL). It belongs to Neurodegeneration with Brain Iron Accumulation, a group of disorders where iron dysregulation is tightly associated with neurodegeneration. We studied the 498–499insTC mutation which causes the substitution of the last 9 amino acids and an elongation of extra 16 amino acids at the C-terminus of L-ferritin peptide. An analysis with cyclic voltammetry on the purified protein showed that this structural modification severely reduces the ability of the protein to store iron. In order to analyze the impact of the mutation in vivo, we generated mouse models for the some pathogenic human FTL gene in FVB and C57BL/6J strains.

Transgenic mice in the FVB background showed high accumulation of the mutated ferritin in brain where it correlated with increased iron deposition with age, as scored by magnetic resonance imaging. Notably, the accumulation of iron–ferritin bodies was accompanied by signs of oxidative damage. In the C57BL/6 background, both the expression of the mutant ferritin and the iron levels were lower than in the FVB strain. Nevertheless, also these mice showed oxidative alterations in the brain. Furthermore, post-natal hippocampal neurons obtained from these mice experienced a marked increased cell death in response to chronic iron overload and/or acute oxidative stress, in comparison to wild-type neurons. Ultrastructural analyses revealed an accumulation of lipofuscin granules associated with iron deposits, particularly enriched in the cerebellum and striatum of our transgenic mice. Finally, experimental subjects were tested throughout development and aging at 2-, 8- and 18-months for behavioral phenotype. Rotarod test revealed a progressive impaired motor coordination building up with age, FTL mutant old mice showing a shorter latency to fall from the apparatus, according to higher accumulation of iron aggregates in the striatum. Our data show that our 498–499insTC mouse models recapitulate early pathological and clinical traits of the human neuroferritinopathy, thus providing a valuable model for the study of the disease. Finally, we propose a mechanistic model of lipofuscin formation that can account for the etiopathogenesis of human neuroferritinopathy.

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Introduction

Iron dyshomeostasis is often associated with neurodegenerative diseases (Rouault, 2013; Vidal et al., 2008), however whether it is a primary cause of the neurodegenerative process or a secondary effect is still debated (Levi and Finazzi, 2014). An opportunity to investigate the iron involvement in neurodegenerative processes is the group of diseases, referred to as “Neurodegeneration with Brain Iron Accumulation”
Ferritin is an iron storage protein able to detoxify cells from the redox-active free iron, that exists in a prevalent cytosolic form and in a minor mitochondrial one (Arosio and Levi, 2010). The cytosolic ferritin is composed of 24 subunits that co-assemble in heteropolymers of H- and L-chains. The H chain has a ferroxidase activity that is necessary for ferric iron incorporation, while the L chain assists and facilitates the nucleation of the ferric iron inside the cavity (Arosio and Levi, 2010). Until now, seven mutations responsible for NF were identified. Only one is a missense mutation in exon 3 of FTL found in a single subject (Maciel et al., 2005) and its causative role is dubious; all the other ones are nucleotide insertion that strongly modify the C-terminus of the L-chain (Curtis et al., 2001; Vidal et al., 2004; Mancuso et al., 2005; Devos et al., 2009; Ohta and Takiyama, 2012; Kubota et al., 2009; Nishida et al., 2014). The C-terminus is rich of inter-chains contacts connecting four subunits at the 4-fold symmetry axis that participate to protein assembly (Luscieti et al., 2010). It has been demonstrated that NF mutations reduce ferritin stability as well as its ability to retain iron within the cavity, and that they act in a dominant negative manner even when present in a low proportions, 2–3 subunits in the 24-mer shells (Luscieti et al., 2010; Cozzi et al., 2010). Actually, the nucleotide insertions/duplications alter the terminal part of the ferritin, the extent depending on the site of insertion. Notably, it was indicated that onset and severity of the disease correlate with the degree of alteration of the protein (Kubota et al., 2009). The mutants are ubiquitously expressed but the main pathological effects are observed in the brain and more specifically in the basal ganglia, where neurons seem to have a higher susceptibility. The 460InsA mutation was the first to be identified in a large number of families in West Anglia and, clinically, is the best characterized form of NF (Curtis et al., 2001). However, most of the experimental research on in vitro and animal models has been concentrated on the insertion of TC nucleotides in FTL gene (498–499InsTC mutation, analogous to the 497–498dupTC) that coding for a variant protein with the substitution of the last 9 amino acids and an extension of further 16 amino acids, which is biochemically more stable than the 460InsA mutant (Muhoberac and Vidal, 2013). These studies reported a reduction in the iron storage capacity of ferritin as well as the formation of ferritin aggregates, suggesting that these features might concur to the pathogenic events of the disease (Baralbar et al., 2008). Other data support and extend these observations by providing evidence of oxidative damage in cells expressing two pathogenic L-ferritin variants in cell lines (460InsA and 498–499InsTC) (Cozzi et al., 2010, 2006) and in fibroblasts derived from a NF patient with 498–499InsTC mutation (Barbeito et al., 2010).

The only animal model of the disorder produced so far is a transgenic mouse overexpressing the 498–499InsTC human ferritin mutant (Vidal et al., 2008). These mice were reported to show nuclear and cytoplasmic aggregates of ferritin throughout the CNS and in other organs, a progressive neurological phenotype, a decreased mobility and a reduced life expectancy as well as an increase in the amount of iron in brain, with altered levels of the iron-related proteins (Vidal et al., 2008). Analysis of brain tissues from these mice indicated an accumulation of oxidized DNA in the mitochondria but no significant damage to the nuclear DNA (Deng et al., 2010); moreover, oxidative stress markers such as protein carbonylation and lipid peroxidation were reported (Barbeito et al., 2009).

Here we report a comprehensive study of the 498–499InsTC human ferritin mutants that includes: i) an electrochemical analysis of the recombinant mutated ferritin heteropolymer, which precisely defines functional alterations of the protein expressed in our mutant mice; ii) a thorough characterization of our transgenic model, which re-evaluates and extends biochemical, morphological and physiological defects; iii) a behavioral investigation of these mice, which formerly proves the pathologic effect of the expression of the mutated ferritin in aged subjects. Overall, our data further extend and redefine the knowledge of the effects exerted by mutated ferritin on cellular iron handling and, more specifically, on the neuronal susceptibility to oxidative stress and motor coordination deficit.

Material and methods

Generation of FTL-498InsTC transgenic mice

The human FTL cDNA bearing the 498–499InsTC mutation (Cozzi et al., 2006) was subcloned into the BamHI–Xba I site between the PGK promoter and the SV40 polyadenylation signal of the pCCL.sin.cPPT.hPGK vector (Follenzi et al., 2000). The correct assembly of the construct was firstly verified by restriction mapping and then confirmed by DNA sequencing (data not shown). The transgene was excised from the vector by SfiI–Hpal restriction digestion and purified by gel extraction with the QIAEX II Gel Extraction Kit (Qiagen, Hilden, Germany). Transgenic mice were generated by pronuclear microinjection of the linearized fragment (shown in Fig. 1S panel A) into fertilized mouse eggs from the FVB/N mouse strain. Microinjection and egg implantation procedures were carried out at the Core Facility for Conditional Mutagenesis (CFCM) Unit at the San Raffaele Scientific Institute, Milano, Italy. Screening for the correct insertion of the transgene was carried out by PCR amplification of the FTL1 gene. To this purpose, tail DNA was purified from offspring of matings of the heterozygous founders with wild-type mice and amplified with the following combination of primers which were specifically designed to amplify a fragment of the human FTL1 cDNA: forward primer (5′–GTCTCCGAAAGTACGAAAC CAG–3′) and reverse primer (5′–AACCTCCACACGGCTGTGAT–3′). The amplification product for construct insertion was of 275 bps (Fig. 15 B). One independent line of the FTL-498InsTC FVB mice was chosen for stable expression of the transgene, as assayed by Western blotting of the non-denaturated protein with the anti-human FTL mouse monoclonal-Ab (LF03, not shown). This FTL-498InsTC line was backcrossed with C57BL/6J (C57) mice (purchased from Harlan Laboratories Srl, Udine, Italy) for >6 generations for generating mice suitable for behavioral studies. Transgenic mice were bred and housed under a 12 h light–dark cycle with free access to food and water.

Tissue preparations

Biochemical analysis was performed on FVB strain at different ages: 3, 6, 12, 18, 22 months-old. For each age were analyzed WT (n = 2 male and 1 female) and Tg (n = 2 male and 1 female). C57 mice were tested at 6,12,18 months-old (n = 2 male and 2 female) and Tg (n = 2 male and 2 female). Mice were anesthetized with Avertine (23 μl/g mouse; Sigma) and perfused with a physiological saline solution containing 2% heparin; tissue samples were then removed and: i) immediately frozen for biochemical analysis; ii) immersed in PBS containing 4% paraformaldehyde 18 h for ex vivo MRI, immunofluorescence and Prussian Blue staining; iii) or immersed in PBS containing 4% parafomaldehyde and 2.5% glutaraldehyde for electron microscopy. For biochemical analysis tissues were lysed in ice-cold lysis buffer (20 mM Tris buffer, pH 7.4, 1% Triton X-100, 1 mM Na azide, 1 mM PMSE, 10 mM leupeptin, 1 mM pepstatin) using a Potter homogenizer. The homogenate was clarified by centrifugation at 10,000 g for 10 min
at 4 °C, and the supernatant was used in Western blotting or in ELISA experiments. For histochemistry and immunofluorescence experiments brains were incubated in PBS containing 4% of paraformaldehyde for 18 h, then, were washed with PBS and included in agarose (4% in PBS) for sectioning. Coronal slices of 40 μm were done with a Cryostat and, then, cryopreserved in a solution composed of 30% glycol ethylene, 30% glycerol and 40% PBS at –20 °C.

For electron microscopy analysis small pieces of fixed brains were dehydrated and embedded in Epon 812. Sections were cut with a diamond knife by an Ultracut UCT (Leica Microsystems, Wetzlar, Germany) and double stained with uranyl acetate and lead citrate.

Ferritin quantification and protein immunodetection

Transgenic human ferritin levels were determined on Architect equipment of Abbot with reagents for the immunological determination of serum ferritin. We verified that the ELISA assay is specific for human L ferritin with no cross reaction with mouse ferritins. Endogenous L- and H-ferritins were quantified by ELISA using specific rabbit polyclonal antibodies, E17 and 217 respectively calibrated on the corresponding recombinant homopolymer (Santambrogio et al., 2007). Protein concentration was evaluated by the bicinchoninic acid (BCA) method (Pierce) calibrated on BSA.

The expression of the transgene was also confirmed by western blotting. 20 μg of soluble proteins were heated at 70 °C for 10 min, centrifuged at 13,000 rpm at 4 °C and the supernatants were separated by non-denaturing-PAGE (native-PAGE). Immunoblotting was performed using a specific antibody for human ferritin (mouse monoclonal LP03) (Cozzi et al., 2006). To study iron incorporation capacity, equal amounts of the 70 °C heated samples were separated on native-PAGE and then the gels were stained with Prussian Blue and enhanced with 0.025% 3,3′-diaminobenzidine (DAB) and 0.05% H2O2 for 15 min. The expression of other proteins was assayed by analyzing 20 μg of homogenate soluble proteins separated by sodium dodecyl sulfate -PAGE (SDS-PAGE) and immunoblotting was performed using specific antibodies for Catalase (EMD, Millipore), TR1 (Zymed), Ubiquitin (Santa Cruz Biotechnology) and β-actin (Sigma), followed by peroxidase-labeled secondary antibodies (Sigma-Aldrich). Band intensity was revealed by enhanced chemiluminescence (ECL; GE Healthcare).

Detection of protein, lipid and DNA oxidation

The following kits were used according to manufacturer’s instructions:

- Oxidized proteins were revealed by the Oxyblot Protein Oxidation Detection Kit (EMD Millipore). In brief, the soluble protein extracts were derivatized to 2,4-dinitrophenylhydrazone (DNP), and 1 μg was loaded on 12% SDS-PAGE, blotted and incubated with an anti-DNP antibody, followed by an HRP conjugated secondary antibody. The bound activity was revealed by ECL advance (GE Healthcare).

- Lipid peroxidation was revealed by the OxiSelect MDA Immunoblot Kit (Cell Biosols, Inc). In brief, 10 μg of the soluble protein extracts was loaded on 12% SDS-PAGE, blotted and incubated with an anti-MDA antibody and then with an anti-Rabbit IgG, HRP conjugate. The bound activity was revealed as above.

- Oxidative DNA damage, was detected in terms of 8-OHdG, by the OxiSelect Oxidative DNA Damage ELISA Kit (Cell Biosols, Inc). Extract DNA from tissues treated with the nuclease P1 was used in ELISA assay: the samples were first added to the 8-OHdG/BSA preabsorbed plates, then incubated with an anti-8-OHdG antibody, followed by an HRP conjugated secondary antibody. The absorbances were detected at 450 nm on a microplate reader.

Histochemistry

DAB-enhanced Prussian Blue staining was performed to identify iron deposition in serial 40 μm coronal mouse brain sections. Briefly, brain sections were washed twice with PBS, permeabilized with PBS/Triton 0.5% and then incubated for 1 h with 4% potassium ferrocyanide in 6% hydrochloric acid. Then slices were reacted with activated 0.014% DAB containing 0.03 H2O2 for 10 min and then washed three times with PBS. DAB staining was acquired with the Nuance Multispectral Tissue Imaging Systems (PerkinElmer) assembled on Zeiss Axioplan2 microscope and quantified with the Advanced Image Analysis Software inForm (PerkinElmer).

Immunofluorescence

Brain slices were washed twice in PBS and incubated with blocking solution (PBS with 10% PBS and 0.3% Triton X-100) for 1 h at room temperature. Then, tissues were incubated overnight at 4 °C with anti-ferritin primary antibody (LF03 5 μg/ml) diluted in blocking solution. Following 3 washes in PBS, cells were incubated with secondary antibody in PBS (donkey anti-mouse Alexa 488; Molecular Probes) for 90 min at room temperature. Tissues were also stained for NeuroTrace 530/615 (a neuronal marker, diluted 1:100 in PBS; Molecular Probes) and TO-PRO-3 (a nuclear marker diluted 1:1000 in PBS; Molecular Probes). All brain slices were mounted on glass slide using Fluorsave (Calbiochem) and acquired on confocal microscope (BioRad confocal).

Magnetic resonance imaging (MRI)

For in vivo MRI, mice were anesthetized with gas (2% isoflurane in oxygen) and brain acquisitions were done with a 7 T MR scanner (Bruker Biospec 30/70, Ettlingen, Germany) and a 4 channel dedicated mouse brain phased-array coil.

2D Fast Low Angle SHot (FLASH) Susceptibility Weighted Imaging (SWI) images [repetition time (TR) = 474.5 ms; echo time (TE) = 15 ms; flip angle (FA) = 30°; number of excitations (NEX) = 10; field of view (FOV) = 2.5 cm; slice thickness/inter-slice distance = 0.6 mm; acquisition matrix = 512 × 384] and multiecho Rapid Acquisition with Relaxation Enhancement (RARE) T2 sequence (TR = 3851.8 ms; 16 echoes with TE ranging from 11 to 176 ms; NEX = 2; FOV = 2.5 cm; slice thickness/inter-slice distance = 0.6 mm; acquisition matrix = 198 × 198) were acquired. T2 maps were calculated using the manufacturer’s software (Paravision 5.1). In order to evaluate the local relaxation times, circular ROIs of the same size (4 × 105 μm2) over different areas of the brain (cortex, striatum, hippocampus, pons and cerebellum) were drawn by the same operator on T2 images and then transferred on T2 maps.

For ex vivo imaging, brain acquisitions were performed on a human grade 3 T MRI scanner (Achieva 3 T, Philips Medical Systems, the Netherland) equipped with a mouse dedicated volumetric coil. The following sequences were: 3D Turbo Spin Echo T2 (TR = 2500 ms; TE = 90 ms; Turbo Spin Echo Factor = 4; NEX = 4; FOV = 2.5 × 1.2 × 1.2 cm; slice thickness = 0.5 × 0.5 × 0.5 mm; acquisition matrix = 256 × 128) and SWI (TR = 61 ms; TE 40 ms; flip angle = 30°; NEX = 28; FOV = 3.5 × 1.9 × 1.4 cm; slice thickness = 0.1 mm; acquisition matrix = 336 × 184).

Electron spectroscopic imaging

The thin sections (~25 nm, gray tone) analyzed by Electron Spectroscopic Imaging (ESI) were first examined at 250 eV (i.e. at an energy loss where scattered electrons of most elements contribute to the image),
thus providing a general view of the ultrastructural organization (Pezzati et al., 1997). The pattern of net iron distribution was then obtained by computer-assisted processing of two images collected below (651 and 683 eV) and one beyond the Fe-L3 absorption edge at 719 eV. The typical energy width of the imaging electrons was 5 eV. A threshold of iron values was routinely set to exclude random noise from the final representation. The map thus obtained, represented according to a pseudocolor scale (from dark red to yellow), was superimposed to the corresponding 250-eV image. The nature of iron signals in the maps was confirmed by EELS spectra.

**Autofluorescence spectra**

Since lipofuscin is a native autofluorescent material we tested its possible presence by exposing brain sections to excitation at two wavelengths: 458 and 514 nm in a Leica SP2 confocal microscope. The lambda scan mode was selected and the detection window width was set to 10 nm. The PMT offset was left at 0. The detection window was moved to the different laser lines and the PMT gain adjusted so that the reflection image at each line was below saturation. AOTF settings for each laser line were adjusted to obtain similar peak intensities. The lambda scan was performed in 10 nm intervals (exc. 458, em. 480–700 nm; exc. 514, em. 540–740).

**Recombinant ferritins**

The recombinant human ferritin homopolymers and heteropolymers were produced in Escherichia coli as described in Lusci et al., 2010. The heteropolymers were produced by a bicistronic expression vector in which the sequence of the Ferritin H chain was followed by a ribosome binding site and by the sequence of the L-chain, wild type or mutated. This construct produced in *E. coli* heteropolymers composed by 20–22 H chain and 4–2 L-chains, which are expected to mimic the subunit composition of the brain ferritin. The proteins were purified and loaded with 1000 Fe atoms per molecule before cyclic voltammetry studies. PfL167fsx26 corresponds to the protein encoded by the 498-499insTC mutation.

**Cyclic voltammetry**

Cyclic voltammetry investigations were performed to determine the electrochemical properties of recombinant ferritins (in Tris 30 mM pH 7.4) and accomplished using a CHI 210B bipotentiosstat (CHI Instrument Inc., Texas, USA). All potentials are referred to Ag/AgCl reference electrode.

We used screen printed gold electrodes (DROPSENS, Llanera, Spain) designed for electrochemical investigation of small sample volumes in order to enable the direct characterization of the dissolved proteins. Using the aforementioned strategy, additional surface adsorption procedures were avoided, that could potentially interfere with the protein functioning and with the iron release/uptake kinetics (Bard and Faulkner, 2001).

**Primary culture of hippocampal neurons**

The hippocampal cultures were prepared from 1–2 d-old transgenic (Tg)-C57 and wild-type (WT)-C57 mice, according to Codazzi et al. (2006). Briefly, the hippocampal sections were digested into Hank’s solution containing 3.5 mg/ml trypsin type IX (Sigma, St. Louis, MO) and 0.5 mg/ml DNase type IV (Calbiochem, La Jolla, CA) for 5 min and then mechanically dissociated in a Hank’s solution supplemented with 12 mM MgSO4 and 0.5 mg/ml DNase IV. After centrifugation, the pool of the neurons obtained from control and transgenic mice were plated onto polyornithine-coated 96-multi wells and maintained in MEM supplemented with 0.3% glucose, B27 supplement, 2 mM glutamax, 5% fetal calf serum, and 3 μM Ara-C (1-D-cytosine-arabinofuranoside) (Sigma). Cultures were maintained at 37 °C in a 5%CO2 humidified incubator, and used from 1 to 2 weeks after plating.

**Cell death analysis**

Hippocampal neurons were maintained in culture for one to two weeks. Before the experiment, cells were washed with Krebs Ringer Henes buffer (KRH, containing 5 mM KCl, 125 mM NaCl, 2 mM CaCl2, 1.2 mM MgSO4, 1.2 mM KH2PO4 and 6 mM glucose, 20 mM Hepes, pH 7.4). Cell death and vital analyses were performed by exposing the cells in Systox Orange and Hoechst 33342, respectively (5 μM in KRH, 5 min incubation at room temperature). The analysis was performed by a High Throughput Microscopy (HTM) system, the IN Cell Analyzer 1000 (GE Healthcare, Grandview Blvd, Waukesha, WI, USA).

**Behavioral test**

Motor coordination and balance were tested using an accelerating rotarod (Accelerating Rotarod, Ugo Basile, Gemonio, Italy). The rotarod test was specifically designed for automated measurements of neurological deficits in rodents; it allows studying motor coordination using the natural fear of falling response in mice (Brooks and Dunnett, 2009). Latency to fall is a more stringent measure of ability to maintain balance, since the requirements for running forward are gradually increasing. Mice with deficits in motor coordination or balance fall off the rotarod before the end of the 5-minute test session (Crawley, 1999). Deficits in the rotarod test can be caused by pathology in the cortex, cerebellum, and spinal cord, in addition to pathology in dopaminergic regions such as the substantia nigra and striatum. More in detail, GABAergic interneurons in the dorsal striatum are seemingly crucial for facilitating fine movement, as GABAergic dysfunction in the striatum has been linked with hyperkinetic movements and motor coordination (Plenz and Kitai, 1998; Do et al., 2013).

Mice were tested at different ages: a) young mice, 2 months-old (n = 7 male and 7 female WT; n = 8 male and 8 female Tg); b) adult mice, 8 months-old (n = 18 male and 12 female WT; n = 15 male and 13 female Tg); c) aged mice, 18 months-old (n = 6 male and 5 female WT; n = 5 male and 5 female Tg). The test was performed by placing a mouse on a rotating rod with a diameter of 3 cm and measuring the latency to fall (in seconds), through a switch placed on the floor below. Mice were given three trials each with a maximum time of 300 s (5 min), during which time the rotating rod underwent a linear acceleration from 4 to 40 rpm over 5 min. Animals were rested a minimum of 10 min between trials to avoid fatigue and exhaustion (Paylor et al., 2006).

**Statistical analyses**

Data, except where otherwise indicated, are reported as the mean ± SD values or as a representative of at least three independent experiments with similar results. Statistically significant differences between CTR and Tg mice were determined by the Student’s *t* test, at three different levels: *p* ≤ 0.05; **p** ≤ 0.01; ***p*** ≤ 0.001.

Behavioral data were analyzed using parametric analysis of variance (ANOVA) with “genotype” (WT and Tg) and “gender” (male and female) as between-subjects factors and “trial” as within-subject repeated measure. Post hoc comparisons were performed by the Tukey’s test. Statistical analysis was performed using Statview II (Abacus Concepts, CA, USA). Data are expressed as mean ± SEM. The level of significance was set at 0.05.

**Experimental procedures and animal care approval**

Animal handling and experimental procedures were performed in accordance with the EC guidelines (EC Council Directive 86/609/1987) and with the Italian legislation on animal experimentation.
The brain of Tg-FVB mice shows increased expression of mutated human FTL and oxidative damage

Groups of three WT-FVB and Tg-FVB mice were sacrificed, the brains collected and the protein extracts analyzed by ELISA for the expression of human FTL as well as of the endogenous FTL and FTH at different ages (3, 6, 12 and ≥18 months). Accumulation of transgenic human FTL increased progressively with age and reached a maximum of about 4 μg/mg of total proteins at 18 months of age (Fig. 2A, upper panel). Also the accumulation of the soluble endogenous murine FTL and FTH increased with age, but they reached a maximum at 12 months. At this
time point, Tg-FVB brains showed 2.5 and 3.1 fold higher levels of FtL and FtH than WT-FVB, respectively (Fig. 2A, middle and lower panels). Immunoblotting analysis of the Transferrin Receptor 1 (TfR1) showed in the brain of Tg-FVB mice a significant reduction (~58%, **p ≤ 0.001; Fig. 2B) that was paralleled by that of mRNA level (~50% reduction, *p ≤ 0.05; obtained by qRT-PCR, not shown). The increase in endogenous ferritin and the concomitant decrease in TfR1 indicate the high availability of intracellular iron, a condition that is line with our experimental evidence that iron is not properly retained inside the cavity of variant ferritins.

We analyzed total ferritin iron and ferritin protein on the same brain extracts. This was done by staining non-denaturing PAGE with enhanced Prussian Blue and with Coomassie blue (Fig. 2C). The analysis of the band densities clearly indicates that the iron/protein ratio

![Image](https://via.placeholder.com/150)

**Fig. 2.** Increased expression of mutated human FTL induces elevation of cytosolic iron and oxidative damage in Tg-FVB mouse brains. A. The soluble brain homogenate fractions of WT-FVB and Tg-FVB mice at different ages were analyzed for human L-ferritin (upper panel) and endogenous L- and H-ferritins (middle and bottom panel, respectively) content by specific ELISA. The values are the means of three experiments performed in triplicate. B 20 μg of brain homogenates of 12-month-old mice were loaded on 12% SDS-PAGE and blotted with an anti-TfR1 antibody (the arrow points to the TfR1). Coomassie blue staining (C.B.) of total protein as loading control. Band densities are reported in the lower panel. C Enhanced Prussian Blue (P.B.) staining of brain soluble homogenates of 12-month-old Tg-FVB mice and WT-FVB siblings (native-PAGE). Coomassie blue staining (C.B.) of ferritin protein as loading control. The histogram shows the iron incorporation capacity of the ferritin in the two groups of animals as the ratio between the total ferritin iron and the ferritin protein content in the same brain extract. D The carbonylated proteins of brain homogenate of 12-month-old WT-FVB and Tg-FVB were revealed by treating with DNPH and blotting with anti-DNP antibody as described in Material and methods. Band densities are expressed as values relative to the mean of the controls of three independent experiments *p ≤ 0.05.
(Fig. 2C) was lower in Tg-FVB than in WT-FVB mice, indicating a reduced capacity of the ferritin in the Tg mice to incorporate iron in vivo. Since this condition is expected to promote oxidative damage, we evaluated the oxidative status of brain tissues by OxyBlot. The transgenic mice showed a higher level of carbonylated proteins already at 6 months (not shown) and reached a two fold increase at 12 months (Fig. 2D, *p ≤ 0.05).

Tg-FVB mice show iron and ferritin accumulation in the brain

The pathologic hallmark of the human disease is the accumulation of iron and ferritin bodies in the brain and other tissues, with late onset neurological symptoms. Based on this premise, brain from old mice were investigated for iron and ferritin content. First, we evaluated iron accumulation in wild-type and transgenic mice in vivo by MRI. The MRI changes associated with iron deposition were investigated in Tg-FVB mice at 18 and 22 months of age as modification of the signal intensity in T2-weighted images (Fig. 3A, left panel, upper row) and in SWI images (Fig. 3A, left panel, lower row). In all Tg-FVB mice, an overall darker appearance could be observed at the level of the cortex and hippocampus in all sequences suggesting diffuse neuronal iron deposition. Further, several hypointense spots highly evocative of iron aggregation could be detected on SWI images in the striatum and Hippocampal cortex of 18 and 22 month-old Tg-FVB mice, but not in age-related WT mice. Using T2 relaxometry, in vivo MRI T2 maps were generated and showed a significant decrease in the T2 decay, a parameter that is well known to correlate with iron deposition, in the frontal cortex, striatum, hippocampus, pons and cerebellum of Tg-FVB mice (Fig. 3B, *p ≤ 0.05; **p ≤ 0.01; ***p ≤ 0.001). As expected, these observations were corroborated by ex vivo MRI analyses which confirmed the presence of the above described hypointense areas on high resolution 3D T2 and SWI sequences (Fig. 3A, right panel). These data most likely indicate the increased level of iron in the brain of living mice expressing mutant FTL.

To verify that iron accumulates in mutant brains, we performed histochemical analysis of iron by DAB-enhanced prussian blue stain. DAB-stained sections of 18 and 22 month-old mutant brains showed a large increase in the number of brown positive precipitates over control mice, with no significant variations between the two ages (Fig. 4A). This result indicates iron accumulation when the human transgene is expressed.

Fig. 3. Iron accumulation in whole brain of FVB mutated mice. A MRI brain examination of 18 and 22 month-old controls and transgenic FVB mice. Left panel: RARE T2 (upper row) and SWI (lower row) coronal images are shown. Areas of brain iron accumulation are detected in vivo as multiple small dark hypointense areas. Right panel: the brain from the same animals were analyzed ex vivo, Fast Spin Echo T2 (upper row) and SWI (lower row) images confirm the results obtained in vivo and show the presence of iron accumulation at the level of the hippocampus, brain cortex and basal ganglia. B In vivo T2 relaxation times of WT- and Tg-FVB mice at 18 and 22 months of age measured on T2 maps. These measurements revealed a significant reduction of the T2 decay in the brains of Tg mice with respect to the control, in particular in striatum, hippocampus, cerebellum and pons, highly suggestive of iron accumulation.
Brain sections were also decorated with a monoclonal antibody for hFtL, Neurotrace as a neuronal marker and TO-PRO-3 for nuclei. The images show the presence of aggregates of human ferritin inside neurons in Tg-FVB mice at both 18 and 22 months (Fig. 4B).

Transgenic mice in the C57BL/6J background show lower accumulation of iron and ferritin but maintain oxidative damage

To perform behavioral analysis, we backcrossed transgenic FVB strain with C57BL/6J (C57) mice for >6 generations (see Material and methods) to obtain an incipient congenic strain where the mutant FTL is stably expressed. The C57 strain is widely used in behavioral studies to investigate motor and higher brain functions. We initially found that the transgene ferritin was expressed at detectable level in the brain of the 12-month-old Tg-C57 mice (n = 4) (Fig. 1S, C), but, surprisingly, its accumulation was much lower than that found in the FVB strain with the same transgene (Fig. 2A upper panel). This occurred at both the transcript (not shown) and the protein level (-90% and -93% decrease, respectively) compared to Tg-FVB brains. MRI was performed in vivo and no hypointense spots could be observed in SWI images of the Tg-C57 brains (not shown). However, MRI T2 maps showed a decrease in T2 relaxation times and an increase in the R2 in the brains of older transgenic mice, suggestive of diffuse neuronal iron deposition: a significant T2 reduction was measured in striatum and pons of 12 and 18-month-old mice compared to age matched controls (Fig. 5, *p ≤ 0.05; **p ≤ 0.01; ***p ≤ 0.001). Of note, the serum endogenous feritin levels halved in transgenic mice (from 775 ± 160 vs 351 ± 139; n = 4), in a way similar to what observed in neuroferritinopathy patients (McNeill et al., 2008). We then evaluated signs of increased oxidative damage. We compared brain tissues from Tg-C57 with those of WT-C57 (n = 4 + 4). We observed higher levels of oxidized proteins (OxyBlot; *p ≤ 0.001, Fig. 6A); higher levels of protein ubiquitination (immunoblotting; *p ≤ 0.05, Fig. 6B); higher lipidic peroxidation (oxy-MDA, *p ≤ 0.05, Fig. 6C) in Tg-C57 mice respect to the WT-C57. Furthermore, the evaluation of 8-OHdG resulted in higher levels of DNA oxidation (1.68 fold, *p ≤ 0.05) respect to the control mice (not shown). The altered oxidative status was also confirmed by a 1.5-fold increase (*p ≤ 0.05) in Tg-C57 mice of the expression of catalase, one of the early defense response to altered oxidative conditions (Fig. 6D).

Primary cultures of hippocampal neurons from Tg mice show increased susceptibility to oxidative stress

Based on the correlation of iron accumulation and oxidative stress observed in Tg-FVB and Tg-C57 brains, hippocampal cultures were obtained from both WT- and Tg-C57 mice. These preparations were subjected to different oxidative stress conditions to mimic, in a very short period of time, what it might occur to neurons during their lifespan. In particular, the susceptibility of neurons to the oxidative insults was evaluated in conditions of chronic iron overload (100 μM ferric iron, administered as ferric ammonium citrate, for 24 h) and/or after acute exposure to hydrogen peroxide (300 μM, added immediately before the experiment). The kinetic analysis of cell death was performed by high throughput microscopy (Pelizzoni et al., 2011), an approach that allowed to partially overcome the high variability of the measurements, largely due to the fact that in the neuronal preparations several hippocampi from littermates with mixed genotype were pooled due to technical restrictions. For all oxidative conditions, a toxic effect over time is demonstrated by the increasing percentage of Sytox-positive nuclei (a marker of cell death); however, this effect was clearly higher in
the neurons from Tg-C57 than in those from WT-C57 matings (Fig. 7A). The statistical analysis of the number of dead cells at the end of the experiments (90 min) shows a significant difference between the two populations, analyzed after both 7 and 14 days in culture (Fig. 7B), indicating an increased susceptibility for oxidative damage of neural populations enriched for mutant FtL in respect with pure WT neurons.

**Tg-C57 mice show iron and lipofuscin accumulation**

Brain tissues were investigated at the ultrastructural level for possible histologic hallmarks of neurodegeneration as early signs of a developing neuroferritinopathy; the presence of these signs is essential to assess the suitability of our animal model to investigate early
pathogenetic events of the disease. The presence of relatively small, membrane bounded and vacuolated, osmophilic bodies was detected in 12 month-old mice (Fig. 8A). However, they were more abundant in Tg-C57 than in WT-C57 mice (about one order of magnitude in the average). Moreover, while their regional distribution was homogeneous in WT brains, a higher density (about 3 folds) was observed in the striatum of Tg mice. The ultrastructural localization of these osmophilic bodies was cytoplasmatic with no evidence of their presence in either the nucleus or extracellularly. They were typically composed of a granular matrix, with irregularly distributed dark-black dots, and a vesicle with a clear inner matrix.

The presence of these osmophilic bodies has long been reported in anatomical and histochemical studies and referred to as lipofuscin granules, i.e. pigment granules containing lipid residues of the lysosomal digestion and metals. Accordingly, we employed electron spectroscopic imaging (ESI), a technique which allows high resolution elemental analysis of thin sections, to look for the presence of iron. In Fig. 8B, a pair of pictures are presented where the image to the left illustrates the ultrastructural organization at 240 eV (i.e., below the carbon K-edge), and that to the right shows the same image with superimposed the ESI iron distribution (coded from dark to light red, low to high values, respectively). The ESI analysis of brain tissue revealed high iron signals within the granular matrix and the dense dots. No comparable signals were observed in other subcellular compartments.

To further confirm the presence of lipofuscin in these osmophilic bodies, we studied their autofluorescence properties. The emission of fluorescence from putative lipofuscin granules was excited at three wavelengths (458, 514 and 543 nm). With increasing excitation wavelength, the emission maximum shifted towards longer wavelengths and spectral width (Fig. 8C) was decreased as previously reported for lipofuscin accumulations in the retina (Sparrow, 2010).

### Tg-C57 mice develop a motor coordination impairment with aging

As a final characterization of our model of neuroferritinopathy, we evaluated the motor coordination by using the natural fear of falling response. The rotarod test revealed a significant effect of trials in young (main effect of trial F(2,52) = 16.884, ***p ≤ 0.0001), adult (main effect of trial F(2,108) = 36.772, ***p ≤ 0.0001) and old (main effect of trial F(2,34) = 4.730, *p ≤ 0.05) mice showing a typical training curve. Analysis of motor performance of transgenic mice on the rotating rod revealed no difference in the latency to fall either at the earliest age tested, (2 months) (no main effect of genotype F(1,26) = 1.803, p ≤ 0.1909) or in the adult group (no main effect of genotype F(1,54) = 0.406, p ≤ 0.5267). Regardless of gender, at 18 months of age Tg-C57 mice fell earlier from the apparatus compared to wild-type subjects (main effect of genotype F(1,17) = 15.317, **p ≤ 0.001 and interaction between genotype and trials F(2,34) = 3.255, *p ≤ 0.05), an effect suggesting impaired motor coordination building up with age (Fig. 8D). Interestingly, female Tg-C57 subjects in trial 3 performed better than males, although not significantly, indicating a less penetrant phenotype in females at old age (interaction among trial, genotype and gender F(1,34) = 2.934, p ≤ 0.0668).

### Discussion

Alteration of cellular oxidative status is a main determinant in neurodegeneration (Rouault, 2013), and iron excess is one of the principal
causes of oxidative stress (Halliwell and Gutteridge, 1992). However, the interplay among iron, oxidative damage and neurodegeneration is not fully understood. Neuroferritinopathy constitutes an interesting pathogenic model to study the interplay among all these factors. Previous studies on patients’ autopic samples (Curtis et al., 2001; Baraibar et al., 2012) as well as on animal (Vidal et al., 2008; Barbeito et al., 2009) and cellular models (Barbeito et al., 2010; Cozzi et al., 2006, 2010) have suggested a few elements as key factors involved in the etiopathogenesis of the disease: i) alteration of iron metabolism, ii) increased oxidative stress and iii) formation of aggregates of iron and ferritin. To prove that these events are interrelated and that their concomitant action leads to neuronal death, we generated two new transgenic mouse models overexpressing the 498–499InsTC human ferritin mutant, under the PGK promoter, in FVB and in C57 strains. This latter mouse model is analogous to that described by Vidal (Vidal et al., 2008), but the transgene is under the PGK promoter instead of the Prion protein one used by Vidal (Vidal et al., 2008). Notably, the prion protein promoter has a much more neuron-selective and powerful expression than the PGK promoter (Borchelt et al., 1996). Hence our C57 mouse model, characterized by a more attenuated and, thus, a more physiological expression of the mutant FTL gene, might better recapitulate the human disease.

Although the transgene was expressed in all tissues tested, we focused our analysis on the brain, the organ in which neuroferritinopathy is pathogenic. The expression of the transgene product progressively increased with age and was paralleled by an elevation of the endogenous ferritins and down-regulation of TIR1. This response is consistent with an elevation of free cytotoxic iron, a condition that modulate, in an opposite way, ferritins and TIR1 translation, via the IRE/IRPs machinery. Therefore these data strongly suggest that the increase in ferritin levels leads to a decrease in iron buffering capacity and not in an elevation, as it would be expected in conditions of overexpression of the native protein (Cozzi et al., 2000).

Based on this premise, we hypothesized that the biophysical properties of the transgenic ferritin in storing iron were altered. To address this point, we investigated recombinant ferritins at the molecular level. A critical issue might be the composition of different subunits in the shell. As previously demonstrated in vitro (Lusci et al., 2010) and as expected by the dominant transmission of the disease, a low percentage of the variant peptide is sufficient to strongly reduce the capacity of ferritin to incorporate iron. Accordingly, we produced heteropolymer ferritins characterized by a high proportion of H chains (21–22 per 24-mer) and a low proportion of the FTL WT or the FTL-HS167SfsX26.

Cyclic voltammetry analysis revealed that the variant ferritin facilitates the release of iron from the shell thereby reducing the overall capacity to retain iron within the inner cavity.

The lower ability of the mutated protein to buffer iron can account for the elevation of the iron reported in patients brains and documented by the MRI and by the presence of granules of ferritin and iron in the autopic samples (Curtis et al., 2001; Baraibar et al., 2012). A very similar condition was observed in the brain of the aged transgenic mice, which showed hypodense areas at the MRI, typical signs of an accumulation of hemosiderin and ferritin within tissues (Robinson and Bhuta, 2011). In particular, in the FVB mutant mice, both T2 and SWI MRI sequences demonstrated the in vivo presence of multiple small dark hypointense areas in the basal ganglia and hippocampi highly suggestive for sites of iron accumulation. MRI T2-relaxometry is a sensitive technique for the in vivo assessment of brain iron content that has already been employed for iron load quantification in different neurological disorders, including PKAN, multiple sclerosis an Parkinson’s disease (Zorzzi et al., 2011; Khalil et al., 2009; Mondino et al., 2002) as well as in neuroferritinopathy (McNeill et al., 2008). A significant accumulation of iron within the striatum, hippocampus and pons was further confirmed, ex vivo, by the presence of histochemical positivity for iron. The number and size of the Fe3+ -positive granules increased with age, in parallel with the accumulation of the exogenous ferritin, mainly localized in neurons.

Additional information was provided by the elemental analysis at the ultrastructural level that revealed iron accumulation within the granular matrix of osmophilic bodies. These bodies were membrane bound and contained vacuolated areas with a clear inner matrix; fluorescence spectral analysis provided conclusive evidence that these granules were lipofuscin and not neuromelanin granules, although they share similar morphology (Double et al., 2008). Notably, although iron aggregates were typically localized close the nucleus, we never observed any nuclear accumulation of this ion. This is at variance with what was reported in the previous mouse model for the same mutation; a reason for this discrepancy may be related to the lower expression of the transgene in our mutant mice (Vidal et al., 2008). However, lack of elemental analysis in the article from the Vidal laboratory makes difficult to compare these morphological aspects.

All our findings point to an altered kinetics of iron uptake/release of the Fh/FTL167SfsX26, which are expected to result in higher availability of free iron. In turn, higher levels of cytoplasmic iron are expected to promote the Fenton reaction whose end-products are ROS. This dynamics is supported by molecular evidence in vitro, where the higher availability of iron around the electrodes coated with the mutated FTL protein caused, in the buffer solution, a potent increase in ROS. We substantiated this observation with conclusive evidence of increased ROS production and of oxidative damage in the brain tissue derived from our transgenic mice by dosing oxidation of proteins, lipids, and nucleic acids with a variety of techniques. Notably, primary neuronal cultures derived from our mouse model showed an increased susceptibility to iron-induced death, thus providing strong evidence that the basal oxidative stress is coupled with a reduced antioxidant cellular defense, a condition which actively cooperates in cell damage. Along this line is the observation of an the increased presence of lipofuscin granules. In fact, lipofuscin is a well-known hallmark of aging, hence its alternate name of “age pigment”, and the rate of its accumulation is reported to correlate negatively with longevity (Bunk and Terman, 2002). A number of experimental evidence have previously linked lipofuscin accumulation with oxidative stress. In particular, it has been reported that the oxidation of 26S proteasome by a variety of altered pathways (Reineke et al., 2000) promotes its disassembly (and consequent accumulation of polyubiquitinated proteins (Wang et al., 2010). This occurrence starts a cascade of events that ultimately leads to the formation of lipofuscin-containing aggresomes (Hohn and Grune, 2013). interestingly, iron has been described as an activator of lipofuscin formation and a major component of these aggresomes (Bunk and Terman, 2002). Lipofuscin accumulates within the lysosomes where the degradation of iron-containing proteins, such as cytchromes and ferritin, leads to the release of redox-active free iron. Then, lipofuscin is able to adsorb ferrous iron thus constituting a redox-active surface that catalyzes the Fenton reaction with generation
of hydroxyl radicals and amplification of the oxidative damage (Hohn and Grune, 2013). Within this framework, the concomitant overloading of the proteasome by all oxidized proteins might favor a diversion of all ferritins, including their oxidized products, towards the aggregosomes in a self-maintained circle of harmful events. The lysosome, overloaded by lipofuscin, may undergo rupture with direct cell damage by hydrolytic enzymes and indirect adverse effects of lipofuscin release into cytosol, where it may act as a catalyst for further iron precipitation by enhancing the formation of ferritin–iron aggregates (Fig. 9).

From a phenotypic point of view, our data indicate that FTL transgenic mice recapitulate not only some pathological signs of human neuroferritinopathy but also some initial neurological signs associated with the disease. In particular, higher accumulation of iron aggregates in the striatum, which is almost exclusively composed of GABAergic neurons, strictly correlates with the defects in motor coordination observed by the rotarod test. These defects were somewhat expected as a direct effect of a dysfunction of the basal nuclei. Furthermore, we scored a progressive worsening of the motor coordination parameters with age, with only 18-month-old Tg mice falling from the rotating rod before controls.

Conclusion

We can conclude that our transgenic neuroferritin model recapitulates some key pathological and clinical aspects of the initial phases of human neuroferritinopathy, including initial CNS pathological hallmarks and motor coordination deficits. Hence, the molecular characterization of this model provides new and relevant insights on the biochemical defects at the basis of the disease, pointing to the oxidative stress as the major molecular effector in the etiopathogenesis of neuroferritinopathy. Overall, our mutants represent a universal model for studying specific therapies aimed to counteract the adverse effects of oxidative stress in the brain, not only in NBIA disorders, but also in a vast group of disorders characterized by ROS production and neurodegenerative processes, including Parkinson and Alzheimer diseases.

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Fig. 9. Scheme of lipofuscin formation pathway. The mutation in ferritin structure induces higher levels of cytosolic free iron (cytLIP), due to its reduced ability to incorporate it, and faster degradation. Enhanced LIP promotes ferritin synthesis and ROS formation with consequent protein oxidation (including ferritin) and proteosomal impairment. Under these conditions, it is favored ferritin/iron aggregation with lysosomal degradation and lipofuscin accumulation. Following lysosome rupture may also cause lipofuscin release with further enhancing of ferritin/iron aggregation.
