



# Autolysis Affects the Iron Cargo of Ferritins in Neurons and Glial Cells at Different Rates in the Human Brain

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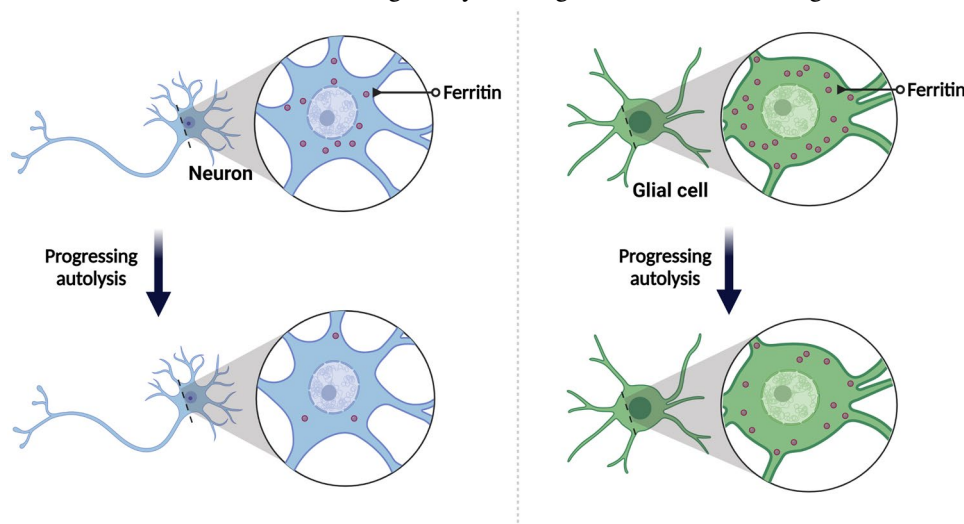
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## Abstract

Iron is known to accumulate in neurological disorders, so a careful balance of the iron concentration is essential for healthy brain functioning. An imbalance in iron homeostasis could arise due to the dysfunction of proteins involved in iron homeostasis. Here, we focus on ferritin—the primary iron storage protein of the brain. In this study, we aimed to improve a method to measure ferritin-bound iron in the human post-mortem brain, and to discern its distribution in particular cell types and brain regions. Though it is known that glial cells and neurons differ in their ferritin concentration, the change in the number and distribution of iron-filled ferritin cores between different cell types during autolysis has not been revealed yet. Here, we show the cellular and region-wide distribution of ferritin in the human brain using state-of-the-art analytical electron microscopy. We validated the concentration of iron-filled ferritin cores to the absolute iron concentration measured by quantitative MRI and inductively coupled plasma mass spectrometry. We show that ferritins lose iron from their cores with the progression of autolysis whereas the overall iron concentrations were unaffected. Although the highest concentration of ferritin was found in glial cells, as the total ferritin concentration increased in a patient, ferritin accumulated more in neurons than in glial cells. Summed up, our findings point out the unique behaviour of neurons in storing iron during autolysis and explain the differences between the absolute iron concentrations and iron-filled ferritin in a cell-type-dependent manner in the human brain.

## Graphical Abstract

The rate of loss of the iron-filled ferritin cores during autolysis is higher in neurons than in glial cells.



**Keywords** Ferritin · Human brain · Energy-filtered transmission electron microscopy · Quantitative magnetic resonance imaging · Inductively coupled plasma mass spectrometry · Autolysis

Extended author information available on the last page of the article

## Introduction

Non-heme iron accumulates in the normal ageing brain until the fourth decade of life (Hallgren and Sourander 1958). The rate of accumulation and the absolute iron concentration vary strongly across brain regions with the highest concentrations in the basal ganglia, including the putamen and the globus pallidus (GP; Hallgren and Sourander 1958; Krebs et al. 2014). The cause of iron accumulation is still unclear, in particular because much more iron is stored than metabolically relevant.

Basal levels of iron are essential in an array of functions in the brain: e.g. myelin synthesis, as cofactors for enzymatic reactions (Codazzi et al. 2015), and synaptic plasticity (Muñoz et al. 2011). Labile iron ions behave as free radicals, inducing reactive oxygen species (Halliwell 2006), and can lead to neuroferritinopathy followed by neurodegeneration (Friedman et al. 2011). In a normal functioning brain, excess iron is stored in its primary storage protein, ferritin (Wang and Pantopoulos 2011), shielding the brain from potential iron toxicity.

Ferritin is at the heart of iron regulation. Ferritin acts as a buffer that stores and releases iron upon cellular demand, which is crucial in preventing ferroptosis (apoptosis induced by iron overload; Jiang et al. 2021) and other detrimental effects caused by free iron. Ferritin contains between 500 and 5000 ferric iron atoms in its inner core (Harrison and Arosio 1996; Iancu 2011; Jiang et al. 2021). The size of the ferritin core in electron micrographs ranges from 5.3 to 11 nm (Iancu 2011) depending on various factors such as iron load, contrasting agent, hydration, or the focus of the micrograph. Despite the difference in the concentrations of total iron and ferritin-bound iron between brain areas, the size of ferritin cores remains largely uniform (Friedman et al. 2011).

Ferritin takes a unique position among iron-binding proteins, due to its size and cargo capacity. Other iron-containing proteins, transferrin and haemoglobin contain only 2 or 4 iron atoms, respectively (Giometto et al. 1993; Marengo-Rowe 2006; Wagner et al. 2003), and are below the detection threshold for energy-filtered transmission electron microscopy (EFTEM), our method of choice for ferritin detection. Hemosiderin is composed of conglomerates of clumped ferritin particles, denatured proteins, and lipids with a size of only 1–2  $\mu\text{m}$  (Wagner et al. 2003). Considering this, hemosiderin is easily distinguished from ferritin by size but is rarely detected in the normal brain.

*Post-mortem* studies have the advantage that tissue is available for a variety of measurements and procedures, including destructive ones. But autolysis, a cell-disintegration process through its internal enzymes (Tsokos and Byard 2016), sets off immediately after death, causing a

breakdown of cells, changing structural components, and hindering cell identification (Sele et al. 2019), reviewed in Lewis et al. (2019).

The total iron concentration in freeze-dried tissue samples can be determined precisely by inductively coupled plasma mass spectrometry (ICP-MS), e.g. in Krebs et al. (2014). However, this method is destructive. A non-invasive and thus non-destructive way to measure spatial iron concentrations is quantitative Magnetic Resonance Imaging (qMRI). qMRI allows mapping the local transverse relaxation rate  $R_2^*$ , which has been validated as a measure for iron content in brain tissue using ICP-MS (Langkammer et al. 2010). However, none of those methods provides information on the amount of iron stored in ferritins or its cellular distribution. We thus established a workflow using EFTEM for localisation and quantification of the iron-filled cores of ferritin. We combined the three methodological approaches in our study to obtain insights into the effects of autolysis on iron storage within ferritin cores.

For this, we investigated the concentrations of iron and ferritin cores filled with iron in the human brain samples *post-mortem* at the region and cellular level. We observed that autolysis influences the loss of iron-filled ferritin cores in a time-dependent manner. Moreover, our study reveals different rates of loss of iron-filled cores between neurons and glial cells during autolysis.

## Materials and Methods

### Human Brain Samples

The samples used here were collected from six deceased human subjects who had to undergo routine brain autopsy at the Diagnostic and Research Institute of Pathology of the University Hospital Graz (approved by the Ethics Committee of the Medical University of Graz, votum number 28-549 ex 15/16). The subjects had not suffered from a known neurological disorder. The sex of the subjects was not considered for this study. To ensure that the analyses were not confounded by age, the specimens were collected from patients with ages ranging from 61 to 86. At this age saturation in iron accumulation has already been reached (Hallgren and Sourander 1958). The age and *post-mortem* intervals (PMI) are given in Table 1. One-half of the brain was preserved for quantitative magnetic resonance imaging (qMRI) while four brain areas were dissected out of the other half and further divided into samples for EFTEM and ICP-MS. The brain regions selected included two regions with the lowest iron concentrations—the frontal grey matter (FGM), the superficial frontal white matter (FWM), and two regions with higher concentrations of

iron—the putamen, and the globus pallidus (GP; Bilgic et al. 2012; Ramos et al. 2014; Ward et al. 2014). Once the tissue was cleared for research purposes by the pathologists, samples were dissected out and either immersed in fixative for EM, or placed into a plastic tube, weighed, and frozen for ICP-MS. The brain hemisphere for qMRI remained unfixed.

## Ferritin Isolation

Ferritin was isolated from human *post-mortem* brain samples in a series of purification steps. In brief, the brain sample weighing approximately 10 g immersed in 10 mM Tris–HCL, 150 mM NaCl containing protease inhibitor was homogenized using Turrax homogenizer IKA T-10. Followed by centrifugation at 10,000g for 30–60 min at 4 °C and sonication for 2 min at 4 °C using a Branson Sonifier 250. The sample was centrifuged again at 10,000g for 30–60 min at 4 °C retaining the supernatant for further purification. The supernatant was heated up to 70–75 °C for 10 min under constant stirring and immediately cooled. 75% saturated ammonium sulfate was used to precipitate the proteins in the supernatant overnight at 4 °C. The precipitated proteins were collected the following day by centrifugation at 10,000g for 60 min at 4 °C. This was followed by size exclusion chromatography and density gradient centrifugation. Isolated ferritin was loaded onto a carbon-coated nickel grid and stained with 1% uranyl acetate to negatively contrast the sample before visualization by EM.

## Electron Microscopy

Following the autopsy, 6–10 sub-samples of approximate wet weight 0.1–0.2 g (maximum dimensions  $1 \times 1 \times 1 \text{ mm}^3$ ) were cut out of the samples and immersed in chemical fixative containing 2% formaldehyde, 2% glutaraldehyde in 0.1 M sodium cacodylate buffer. After fixation, the samples were rinsed in the same buffer, postfixed in 1% osmium tetroxide solution in the same buffer, dehydrated in a graded series of alcohol, immersed in propylene oxide,

and embedded in TAAB embedding resin (TAAB, Aldermaston, UK). After curing for 3 days at 60 °C, thin sections were cut using a Leica UC6 or a Leica UC7 ultramicrotome at a thickness of either 60 or 70 nm. The sections were contrasted with platinum blue and lead citrate solutions. One block was randomly selected from each brain area for imaging and elemental mapping.

EM was performed with a Zeiss EM 900 electron microscope at 80 kV for cell type identification and a Thermo Fisher Tecnai G2 20 electron microscope operated at 200 keV for EFTEM. An Ametek Gatan Quantum GIF energy filter with Ametek Gatan Ultrascan 1000 XP camera was used for EFTEM, and a bottom-mounted Ametek Gatan Ultrascan 1000 camera for bright field imaging. For elemental mapping, three windows—Pre-edge 1, Pre-edge 2, and Post-edge—were made at 80,000× magnification, a binning of 2, a slot width of 40 eV, and an exposure time of 30 s. The energy losses were: 718 eV for the iron L—post-edge window, and 633 eV and 673 eV for the two pre-edge windows. Drift correction was done manually, and both an iron L—elemental map and an iron L—jump-ratio were computed on Digital Micrograph (Gatan, Inc) using respectively the three-window method and the two-window method. The iron cores inside the ferritins are visible as bright spots using EFTEM.

An unbiased sampling protocol described earlier (JoVE Video Dataset; Wernitznig et al. 2019) automatically selects 60 random locations on each sample determined using a random number generator. The elemental maps were accompanied by overview micrographs taken from the same locations using the bright field mode at 3500× magnification.

## Ferritin Quantification and Cell Type Assessment

Four independent researchers (SS, SnR, SL, and GL) each quantified the number of iron-loaded ferritins and assessed the cell types. Ferritins were only counted if they were clearly visible as a bright spot on both the iron L—jump-ratio and the iron L—elemental maps and if at least three of the four researchers agreed on their number. In case of disagreement, the images were analysed again simultaneously. If the number of ferritins exceeded 20 in any map, the mean value of the counts from four researchers was considered as the final count. Cell types were identified on the bright field images and only noted if three of the four researchers agreed. For cell identification, the criteria described by Aten et al. (2022), Hermel et al. (2006), and Nahirney and Tremblay (2021) and the Boston University web atlas (Peters and Sethares 2022) were used. A total of 1440 iron L—elemental maps and 1440 iron L—jump-ratios were generated and used for this study.

**Table 1** Patient information

Patient code	Age (years)	PMI (h)
1	73	06:30
2	72	9
3	61	16
4	69	18
5	86	20
6	81	24

## Quantitative Magnetic Resonance Imaging

MR imaging of the extracted brain hemisphere was performed using a 3.0 Tesla scanner (MAGNETOM PRISMA, Siemens Healthineers, Erlangen, Germany). A phased array head coil with 20 elements was used. For acquiring  $R_2^*$  relaxation data, a two-dimensional radiofrequency-spoiled multi-echo gradient-echo sequence with 6 equally spaced echoes was applied (repetition time = 1150 ms, first echo time = 4.92 ms, echo spacing = 5.42 ms, flip angle = 15°; field of view = 256 × 256 mm<sup>2</sup>; in-plane resolution = 0.75 × 0.75 mm<sup>2</sup>; 2.4-mm-thick sections covering the entire hemisphere). Generalized auto-calibrating partially parallel acquisition (GRAPPA) was performed with an acceleration factor of 2.

To reduce phase dispersion effects and to increase B0 homogeneity, second-order shimming was applied.  $R_2^*$  maps were calculated with the relaxometry toolbox (ESM-RMB 2016, 33rd Annual Scientific Meeting, Vienna, AT, September 29–October 1: ePoster/Paper Poster/Clinical Review Poster/Software Exhibits 2016).

## Inductively Coupled Plasma Mass Spectrometry

All solutions were prepared with ultrapure water (18.2 MΩ cm, Merck Millipore, Bedford, USA). Nitric acid (> 65% p.a.) was further purified via sub-boiling. For quantification, single elements standards (ROTI®Star, Carl Roth GmbH+Co. KG, Karlsruhe, Germany) were used. Brain tissue was freeze-dried to constant mass with a Gamma 1–16 LSC freeze-dryer (Martin Christ GmbH, Osterode am Harz, Germany) and the wet/dry mass ratio was determined. Freeze-dried samples were weighed to 0.1 mg into the 12 mL quartz vessels of a microwave-heated autoclave, UltraCLAVE IV (EMLS, Leutkirch, Germany). After the addition of 3 mL subboiled nitric acid and 2 mL of water the quartz vessels were placed in the 40 positions rack of the autoclave and the autoclave was pressurized with Argon to a pressure of  $4 \times 10^6$  Pa. The following microwave heating program was applied: from room temperature to 80 °C in 5 min, from 80 to 150 °C in 15 min, from 150 to 250 °C in 20 min, and 250 °C for 30 min. After cooling to 80 °C, the pressure was released and the samples were transferred to 50 mL polypropylene tubes (CELLSTAR®, blue screw cap, Greiner Bio-One GmbH, Kremsmuenster, Austria) and filled to the mark.

For the quantification of iron (iron calibration curves were prepared from 0.0100 to 10.0 Fe. Germanium (Ge) at a concentration of 200 µg/L was added as an internal standard online before the nebulizer. The samples were pumped through a peristaltic pump tubing with an i.d. of 1.05 mm and the internal standard with a tubing of 0.19 mm. The elemental iron was quantified with an inductively coupled

plasma mass spectrometer (ICP-MS, Agilent 7700x, Agilent Technologies, Waldbronn, Germany) at a mass-to-charge ratio of Fe  $m/z = 56$ . All isotopes were measured in the collision mode using He at a flow rate of 5.0 mL/min. The determined concentrations were calculated for fresh tissue. The accuracy of the determined concentrations was confirmed with the certified material bovine muscle (RM 8414; NIST; now BOVM-1 NRCC).

## Statistical Analysis

Statistical analysis was performed using GraphPad Prism 9.2.0. One-way ANOVA with a Tukey's multiple comparisons test was used for statistical analysis: \* $p < 0.05$ ; \*\*\* $p < 0.001$ ; \*\*\*\* $p < 0.0001$  for Figs. 2, 3, 5 and 8. For all comparison plots of data from three methodological approaches in Figs. 6, 7, and 9, a simple linear correlation analysis was performed.

## Results

### EFTEM Elemental Mapping Allows Visualizing the Ferritins' Cores in the Human Brain

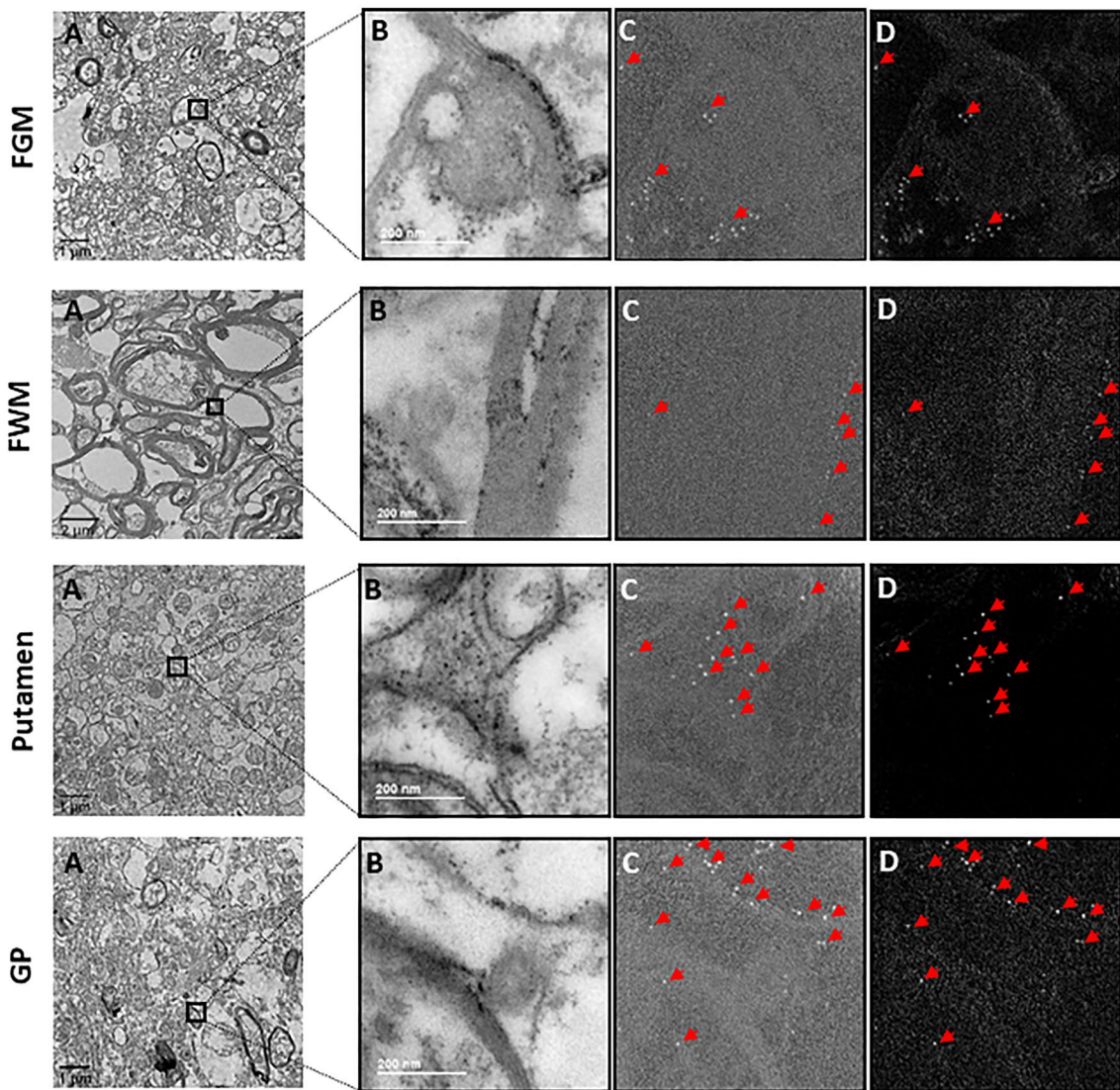
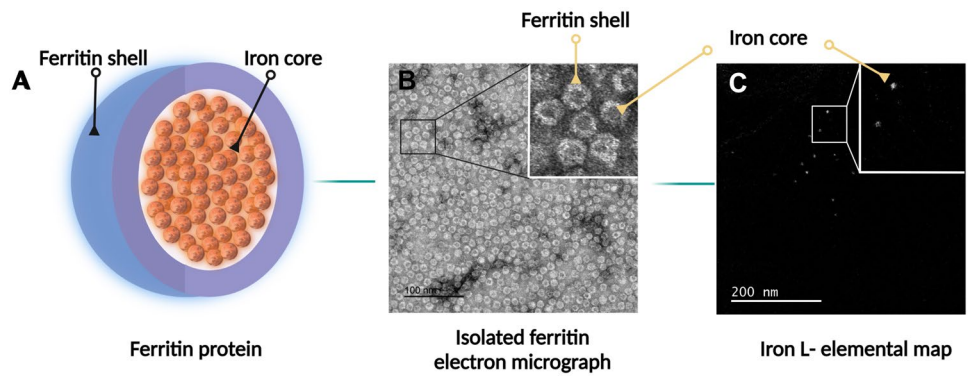
Ferritins consist of a shell and an iron-filled core, and the core is visible as an electron-dense body in negative contrast electron micrographs of isolated ferritins (Fig. 1b). The same core is visible as a bright spot in iron L—elemental maps (Fig. 1c).

To visualize the ferritin cores in electron micrographs, we generated iron L—elemental maps and iron L—jump-ratios of brain samples for four different brain areas in each of six different deceased subjects. The maps consistently exhibited ferritins as bright spots of 5–10 nm in diameter in all four examined brain regions (Fig. 2a–d). The size of the spots corresponded to the published sizes of ferritin cores (Iancu 2011), indicating that iron-filled ferritin cores could be visualized with EFTEM.

### Ferritin Cores Were Clustered in Distinct Cells

We next counted the number of ferritin cores in samples taken from FGM, FWM, putamen, and GP of each patient. The counts were made within 60 micrographs of 556 × 556 nm<sup>2</sup> dimensions and showed that the ferritins were not evenly distributed but clustered in certain micrographs (Fig. 3). We observed that the majority of ferritin accumulated as clusters as shown in Fig. 1, inside the cells. At a closer examination, it became apparent that certain cells accumulated ferritins, whereas others, often neighbouring cells, did not accumulate ferritins (Fig. 2). None of the ferritins appeared to be extracellular. We

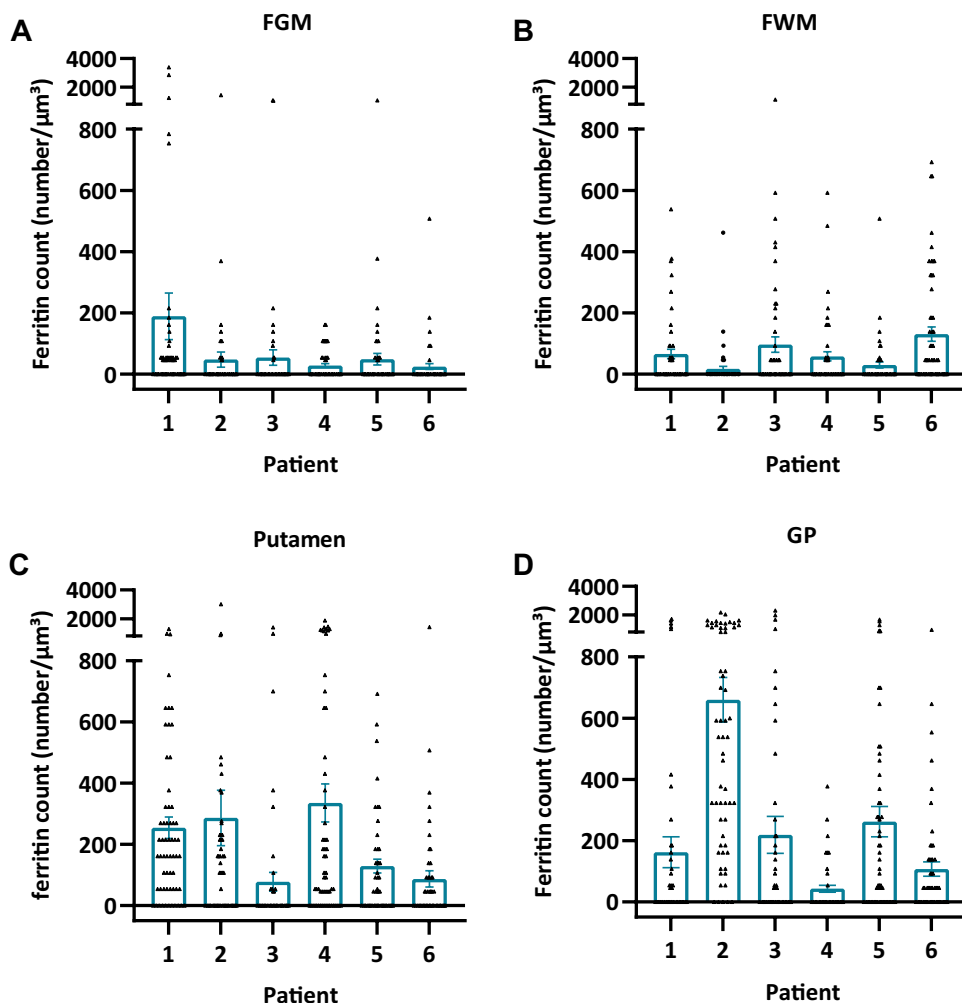
**Fig. 1** Visualization of the iron core of ferritin. **a** Schematic diagram of the molecular architecture of ferritin. **b** Negative contrasted electron micrograph of isolated ferritin from the human brain. The ferritin shell appears as white rings shielding the electron-dense iron core (inset). **c** Iron L—elemental map displaying the iron cores as bright spots



**Fig. 2** EFTEM mapping of iron in different regions in post-mortem human brain sample. **a** Bright-field electron micrographs of overview images of the FGM region, the FWM region, the putamen region, and the GP region. **b** Inverted post-edge micrographs. **c** iron L—jump-

ratios. **d** corresponding iron L—elemental maps of the same region as B. Ferritin filled with iron core appear as bright spots. Examples are marked by red arrows. Corresponding iron L—jump-ratios and iron L—elemental maps (**c**, **d**) show ferritin present in clusters in the cells

**Fig. 3** The ferritin concentration in different brain regions varies between patients 1–6. **a** FGM. **b** FWM. **c** Putamen. **d** GP. Data are shown as bar graphs with whiskers representing the mean  $\pm$  SEM. Each dot represents the ferritin concentration measured in one iron L—elemental map, ( $n = 60$  for each patient in each brain region). One-way ANOVA with Tukey’s multiple comparisons test was used



were able to assess the cell type (neuron, oligodendrocyte, astrocyte, or other unidentified glial cells) in 2195 (62%) of the 3537 ferritin cores that we had identified in five out of six samples. 471 additional ferritin cores were identified in the sixth sample with a *post-mortem* interval (PMI) of 24 h, but in this sample, autolysis had progressed considerably, preventing us from faithfully identifying cell types. Accordingly, this sample was excluded from cell type identification experiments.

### Ferritins Accumulate in the Putamen and Globus Pallidus

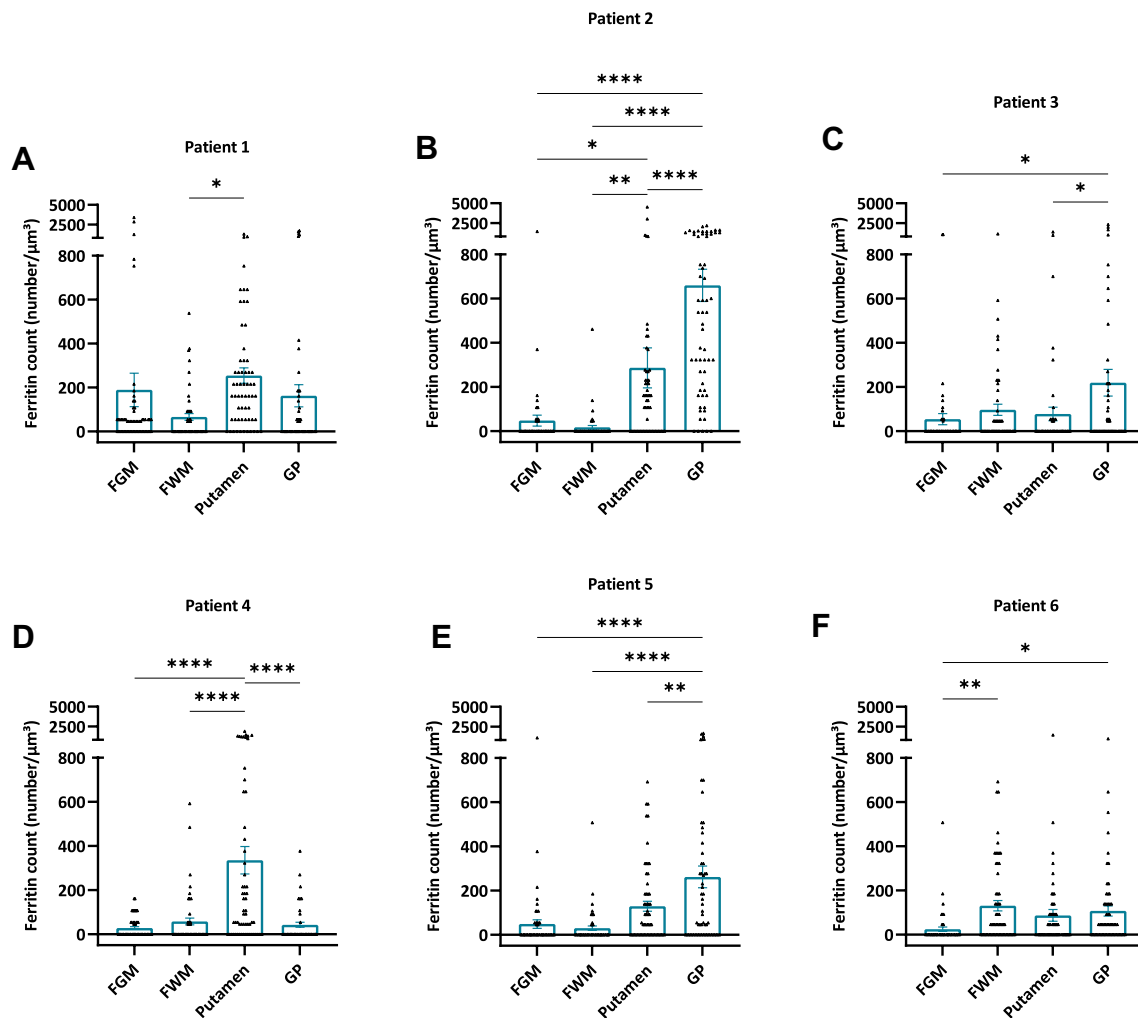
We next calculated the number of ferritins per  $\mu\text{m}^3$  from the micrographs’ 3D dimensions. Plotting ferritin concentration in each brain region for each patient revealed that either the putamen, GP, or both, had a significantly higher concentration of ferritins than either the FGM or FWM in five out

of six patients (Fig. 4). Further, between the putamen and GP (4 out of 6 patients) significant differences were found (Fig. 4). The GP had a significantly higher concentration compared to the former in three out of six patients and the putamen had a significantly higher ferritin concentration than the GP in one out of six patients (Fig. 4).

### qMRI and ICP-MS Data are in Line with the EFTEM Data

The ferritin concentrations counted from elemental maps were validated with the mean iron concentrations. The latter was determined by both mass spectrometry in adjacent tissue samples and by the proton transverse relaxation rate  $R_2^*$ —a qMRI measure proportional to the iron concentration—measured in the contralateral hemisphere (Fig. 5).

Each of the three methodological approaches provided higher mean values (of all the six samples) in the putamen and GP than in FGM and FWM (Fig. 6a–c). qMRI revealed

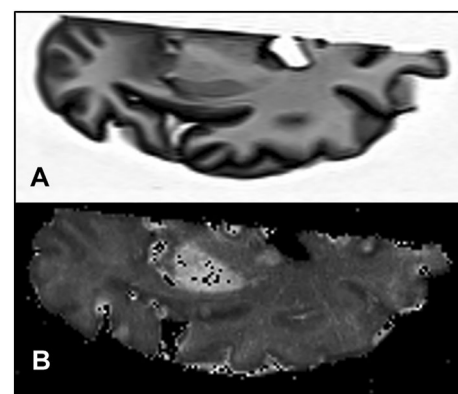


**Fig. 4** The ferritin concentration is often, but not always, higher in the basal ganglia than in the frontal gry or frontal white matter. Graphs show the ferritin concentration in the FGM, FWM, putamen, and GP in **a** patient 1, **b** patient 2, **c** patient 3, **d** patient 4, **e** patient 5, and **f** patient 6. Data are shown as bar graphs with whisk-

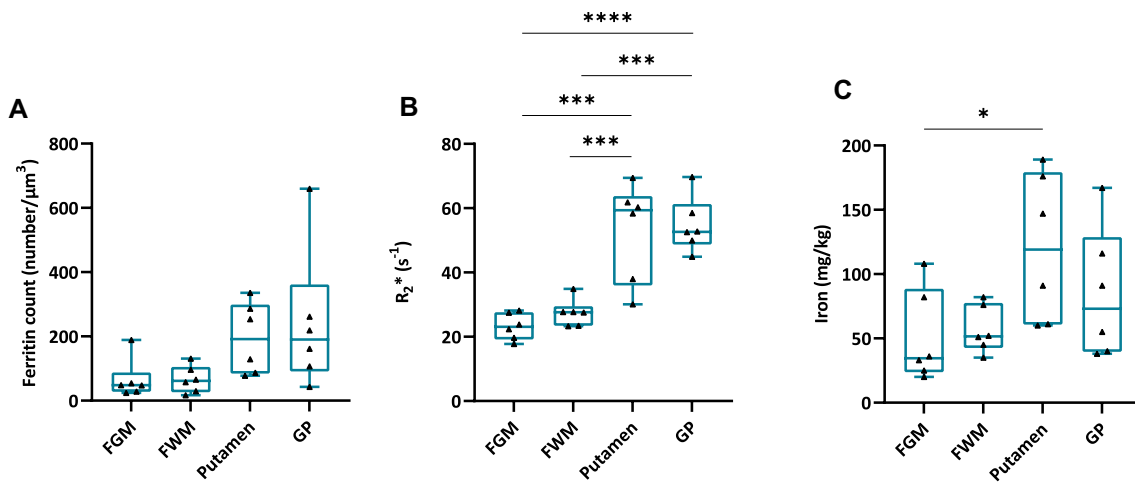
ers representing the mean  $\pm$  SEM. Each dot represents the ferritin concentration measured in one iron L—elemental map, respectively ( $n=60$  for each brain region and each patient). \* $p<0.05$ ; \*\* $p<0.01$ ; \*\*\*\* $p<0.0001$ , One-way ANOVA with a Tukey's multiple comparisons test

a significant  $R_2^*$  increase in putamen and GP compared to FGM and FWM (Fig. 6b). Significant differences were found in the iron concentrations between the putamen and FGM using mass spectrometry (Fig. 6c).

To compare the three different methodological approaches, the mean values of the measured brain regions from all six patients were correlated with each other. There was a high correlation between qMRI and ICP-MS results, each measuring the absolute iron concentration. The higher the  $R_2^*$  value, the higher the iron concentration (Fig. 7a, slope differs significantly from 0). However, we observed that the mean ferritin count obtained using EFTEM does not correlate with the iron concentration obtained with either qMRI (Fig. 7b) or ICP-MS (Fig. 7c). This apparent discrepancy prompted us to check whether the autolysis had an influence on the ferritin count or its cellular distribution.

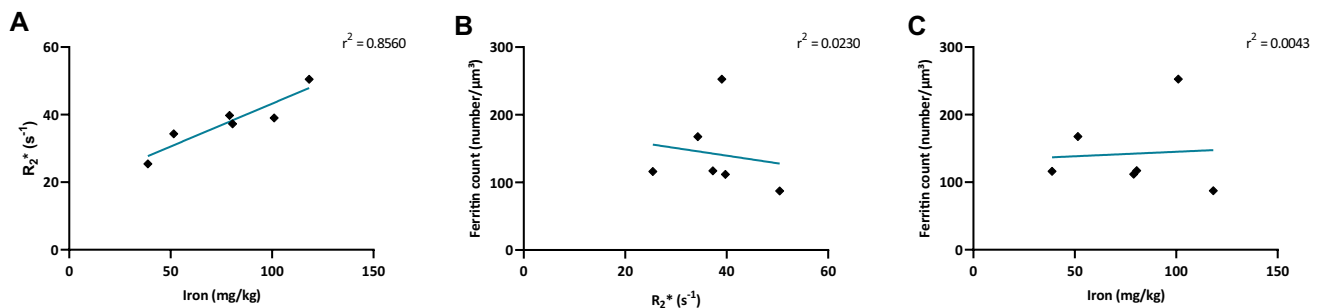


**Fig. 5** MRI scan showing an axial T1-weighted image of **a** one cerebral hemisphere, and **b** the corresponding  $R_2^*$  map. The bright areas in **b** are the putamen and the globus pallidus with higher  $R_2^*$  rates as a consequence of age-related iron accumulation



**Fig. 6** qMRI and ICP-MS data support EFTEM findings. **a** Ferritin concentration in the FGM, FWM, putamen, and GP measured by EFTEM. **b** Proton transverse relaxation rate ( $R_2^*$ ) in the FGM, FWM, putamen, and GP measured by qMRI. **c** Iron concentration in the FGM, FWM, putamen, and GP measured by ICP-MS. The differences in ferritin concentration determined by EFTEM between brain

areas follow the same trend as the sample from the counter brain half in qMRI data and adjacent samples in ICP-MS data. Data are shown as bar graphs with whiskers representing the mean  $\pm$  SEM. Each dot represents the mean ferritin concentration measured in one patient, respectively ( $n=6$ ). \* $p < 0.05$ ; \*\* $p < 0.01$ ; \*\*\*\* $p < 0.0001$ , One-way ANOVA with a Tukey's multiple comparisons test



**Fig. 7** Correlation between mean concentrations of ferritin and iron measured via EFTEM, qMRI, and ICP-MS, respectively. **a** Proton transverse relaxation rate (qMRI) plotted over the iron concentration (ICP-MS) for each patient. The mean proton transverse relaxation rate of each patient correlates significantly with the mean iron concentration,  $r^2=0.8560$ ,  $p=0.00082$ . **b** Mean ferritin concentration (EFTEM) plotted over the proton transverse relaxation rate ( $R_2^*$ ) (qMRI). **c** Mean ferritin concentration (EFTEM) plotted over the iron

concentration (ICP-MS). In contrast to **a**, the mean concentration of ferritin particles does not correlate with proton relaxation rate **b**,  $r^2=0.0230$ ,  $p=0.7742$  nor with the iron concentration **c**,  $r^2=0.0043$ ,  $p=0.9015$ . Each dot represents the mean ferritin concentration or the mean iron concentration measured in one patient, respectively ( $n=6$ ). All the plots are based on mean values of the FGM, FWM, the putamen, and the GP. \* $p < 0.05$ ; \*\* $p < 0.01$ ; \*\*\*\* $p < 0.0001$ , Linear correlation analysis

### Autolysis Enhances the Unloading of the Iron Cargo from Ferritin Cores

We next verified whether putative autolysis that progresses with increasing PMI influenced the iron concentration, the storage of iron in ferritins' core, or the cellular distribution of the ferritins.

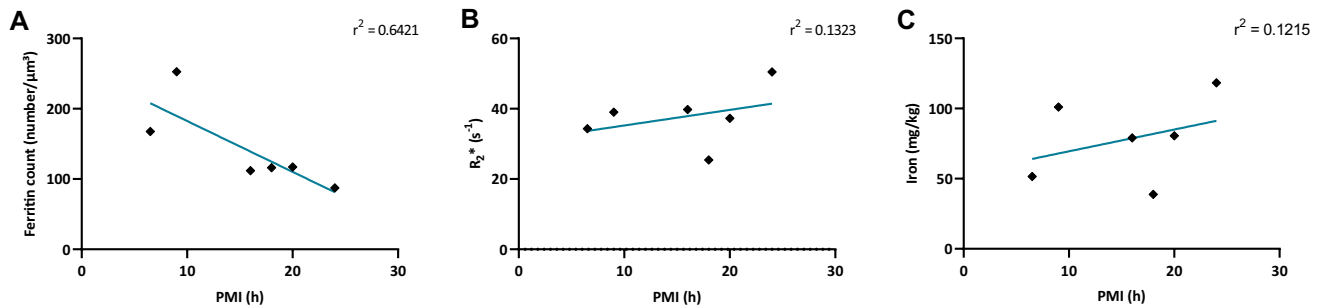
Our analysis showed that the mean ferritin count was inversely correlated to the PMI (Fig. 8a). In contrast, neither the  $R_2^*$  value (Fig. 8b) nor the ICP-MS determined iron concentration correlated with the PMI (Fig. 8c). In other words, ferritin loses iron from its core with progressing autolysis

while the absolute iron concentration remains stable in each patient.

### Cell Type Identification Reveals that Ferritins are Highly Concentrated in Glial Cells, Especially Oligodendrocytes

The counted ferritin was assigned to different cell types by four independent researchers. Representative examples of these cell types are shown in Fig. 9. Oligodendrocytes are recognized by their electron-dense cytoplasm and their heterogeneous nuclear chromatin pattern in their electron-dense





**Fig. 8** Ferritins disintegrate and their concentration is reduced over post-mortem time. **a** Mean values of the ferritin concentration (EFTEM) in all the brain regions plotted against PMI,  $r^2=0.6421$ ,  $p=0.0553$ . **b** Mean values of the proton transverse relaxation rate (qMRI) in all the brain regions plotted against PMI,  $r^2=0.1323$ ,  $p=0.4785$ . **c** Mean values of the iron concentration (ICP-MS) in all the brain regions plotted against PMI,  $r^2=0.1215$ ,  $p=0.4983$ . The

ferritin concentration decreases with PMI in regions of the brain (**a**) whereas the total iron load in the four regions of the brain does not seem to be influenced by the PMI (**b**, **c**). Each dot represents the mean ferritin concentration or the mean iron concentration measured in one patient, respectively ( $n=6$ ). All the plots are based on mean values of the FGM, FWM, the putamen, and the GP. \* $p < 0.05$ . Linear correlation analysis

nucleus (Fig. 9a). Astrocytes have a pale, sparse cytoplasm with few electron-dense inclusions in their perikaryon. They possess a thin rim of heterochromatin right beneath the nuclear envelope (Fig. 9b). A dark, electron-dense cytoplasm of microglia is similar to that of oligodendrocytes, but only “on the first sight”. Namely, the microglial cytoplasm often contains phagocytosed material and lipid inclusions in the form of lipid droplets (Fig. 9c). The glial cells that we could not assign to a specific cell type were labeled as undefined glial cells (Fig. 9d). Neurons have cytoplasm densely filled with Nissl bodies. The nuclei are dark with uniformly dispersed heterochromatin (Fig. 9e). The iron-filled ferritin cores identified in axon terminal and dendritic spines were also assigned to the category of neurons. Axon terminals in a synaptic connection are readily identified with synaptic vesicles in the pre-synaptic terminal and dendrites with a post-synaptic density (Fig. 9f). If myelinated, axons are wrapped by layers of myelin sheath by an oligodendrocytic process and have microtubules and mitochondria in their cytoplasm (Fig. 9g). The cellular ferritin distribution showed that most of the 2195 ferritin cores that could be assigned to a specific cell type were found within glial cells. Oligodendrocytes represented the major cell type within this fraction (59%). 17% were found in neurons, 8% in astrocytes, and 1.7% in microglial cells (Fig. 9h).

### Ferritin Retention in Neurons Varies with Both Autolysis and the Total Ferritin Count

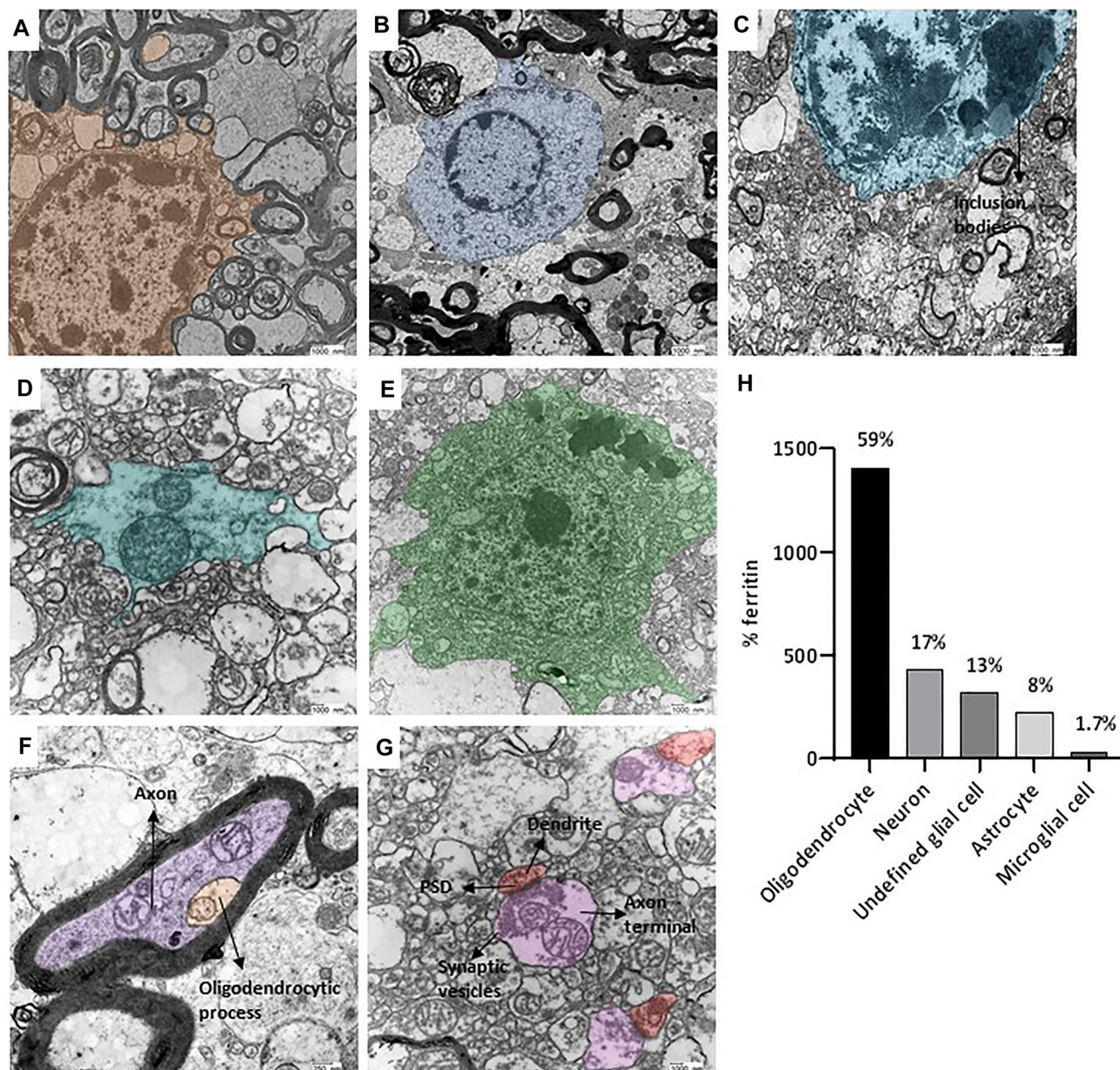
Next, we studied the influence of autolysis on the iron load within ferritins in neurons. We found that the percentage of iron-filled ferritin cores found using EFTEM in neurons was inversely proportional to the PMI (Fig. 10a), indicating that unloading of ferritins during autolysis is faster in neurons than in glial cells (Fig. 10b).

To evaluate whether the cell-specific accumulation of ferritin is dependent on the total ferritin amount, we plotted the percentage of ferritin found in neurons against the total ferritin count in each sample. We observed that the percentage of ferritins stored in neurons was proportional to the total ferritin count. In other words, when the ferritin concentration increased in a patient, the probability of ferritins accumulating in neurons increased. Thus, although the majority of ferritins were situated in glial cells, total ferritin load scaled with the proportion of ferritins found in neurons (Fig. 10c) and vice-versa for glial cells (Fig. 10d).

## Discussion

Our studies implemented analytical electron microscopy to determine the concentration of ferritin cores filled with iron and used ICP-MS and  $R_2^*$  relaxation rate from qMRI to determine the absolute iron concentration in the human brain. Samples from four brain regions of six deceased subjects were examined. The subjects had not suffered from a known neurological disorder and the samples were taken during routine autopsies.

Due to the electron density of the ferritins’ iron cores, they are directly visible in classical electron micrographs (reviewed in Iancu 2011), but they can become obscured by the contrasting agent in routine contrasted sections (Iancu 2011). Accordingly, we chose an energy filter to create iron maps for trustworthy ferritin core identification. The size (Iancu 2011) and detectability of these clusters indicate that they correspond to the iron-loaded cores of ferritins. After identifying the ferritin cores, we compared the elemental maps with a classical electron micrograph of the same area for cell-type identification and ferritin localization. We were able to demonstrate that



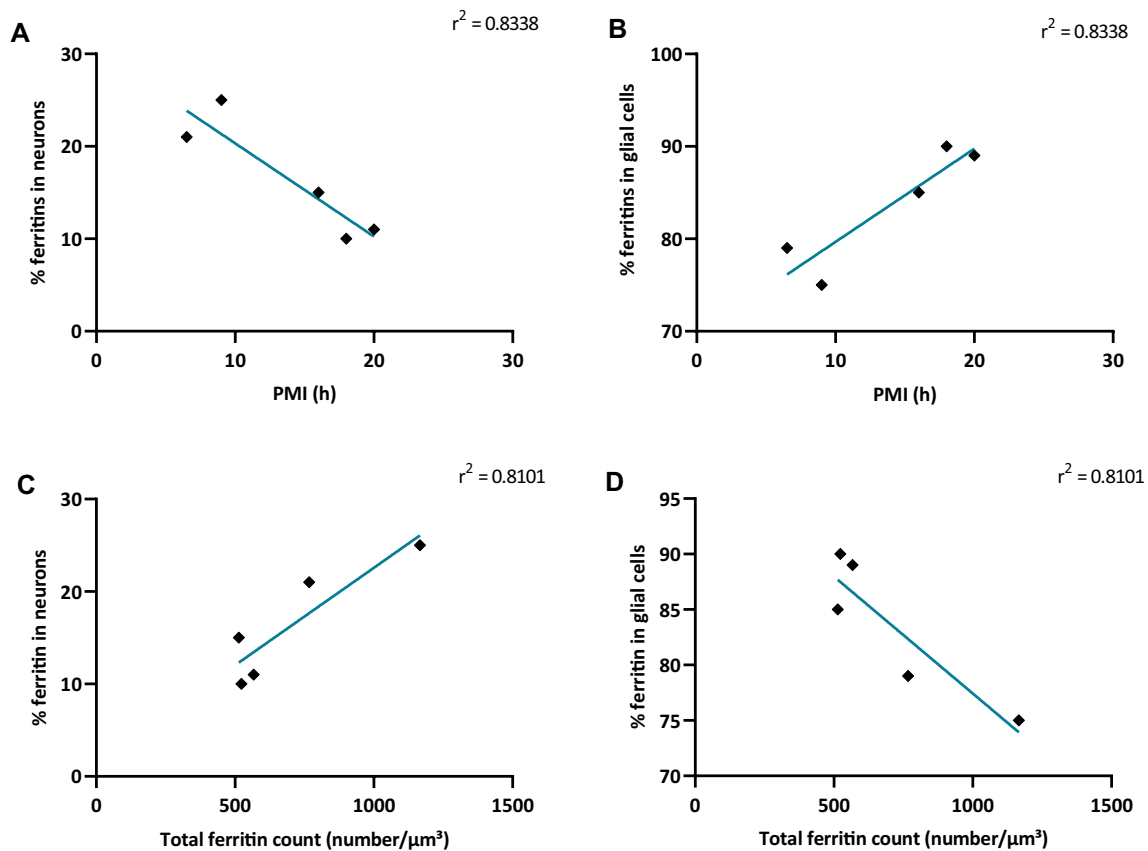
**Fig. 9** Identification of cell types and cellular distribution of ferritin. **a** Oligodendrocyte. **b** Astrocyte. **c** Microglial cell. **d** Unidentified glial cell. **e** Neuron. **f** Myelinated axon and oligodendrocytic process. **g** Dendritic spine and axon terminal in synaptic junction. **h** Ferritin is predominately stored in glial cells (82%) and especially in oligodendrocytes (59%); followed by 8% in astrocytes and 1.7% in micro-

glial cells of the identified glial cells. 17% is stored in neurons. The graph **h** shows those ferritins in the FGM, FWM, putamen, and GP of patients 1–5 whose cell type could be determined. One-way ANOVA with Tukey's multiple comparisons test was used for statistical analysis

the putamen and the GP store more ferritin iron clusters than the FGM and the FWM, confirming a trend also seen in data obtained with both qMRI and ICP-MS.

The EFTEM method is based on visualizing iron clusters inside ferritin using analytical electron microscopy. Ferritins are known to vary in their iron load (Harrison and Arosio 1996; Iancu 2011; Jian et al. 2016) and those ferritin proteins that contain a low amount of iron remain undetectable

using EFTEM. Instead of using a fixed detection threshold, this study relied on four independent researchers agreeing on the ferritin count from both iron L—elemental maps and iron L—jump-ratios. There is high variability in the data obtained using the EFTEM method due to the small sampling size. Hence, the mean concentrations of ferritin both from each patient and from each region were used for this study.



**Fig. 10** The percentage of ferritin in neurons varies with post-mortem time and with total ferritin concentration. **a** Percentage of ferritins in neurons from patients 1 to 5 plotted against PMI. Mean values of the percentage of ferritin concentration in neurons (EFTEM) reduce over time with increasing PMI,  $r^2=0.8338$ ,  $p=0.0303$ . **b** Percentage of ferritins in glial cells from patients 1 to 5 plotted against PMI. Mean values of the percentage of ferritin concentration in glial cells (EFTEM) reduce over time with increasing PMI,  $r^2=0.8338$ ,  $p=0.0303$ . **c** Mean values of the percentage of ferritins found in

neurons in all brain regions from patients 1 to 5 were plotted against total ferritin count. The probability of ferritins being found in neurons rises with the total ferritin concentration,  $r^2=0.8101$ ,  $p=0.0374$ . **d** The probability of ferritins being found in glial cells decreases with the total ferritin concentration,  $r^2=0.8101$ ,  $p=0.0374$ . Each dot represents the mean percentage of ferritin found in neurons in all brain regions measured in patients 1–5, respectively ( $n=5$ ). \* $p < 0.05$ , Linear correlation analysis

We were able to demonstrate that the putamen and GP had a higher mean concentration of detectable (iron-loaded) ferritins than the FGM and FWM. The basal ganglia as the storage site for iron have long been known (Hallgren and Sourander 1958) and have since been confirmed in several different studies that used a variety of different methods (Krebs et al. 2014; Maeda et al. 1997; McAllum et al. 2020; Pfefferbaum et al. 2009; Ramos et al. 2014). To validate our findings, we measured the iron concentration in adjacent pieces of tissue taken from the same patients with ICP-MS, and measured the proton relaxation rate  $R_2^*$ , using qMRI in the contralateral region of the same patients. Our study on iron concentration validates and confirms earlier studies in that the basal ganglia have both higher iron concentrations and a higher  $R_2^*$  relaxation rate than the frontal grey and white matter (Langkammer et al. 2010).

Our study found a high correlation between the  $R_2^*$  relaxation rate (qMRI) and the absolute iron concentration (ICP-MS), confirming an earlier, comparative study of both methods (Krebs et al. 2014; Langkammer et al. 2010). In contrast, the mean ferritin concentration of all the brain areas did not correlate with the mean iron concentration, indicating that the number of iron-loaded ferritins was not directly dependent on the total iron concentration in each patient. It has to be borne in mind that the EFTEM method measures only those ferritin cores that are filled with a minimum threshold amount of iron (cargo), whereas both qMRI and ICP-MS measure the total elemental iron load of the brain sample. We thus investigated the factors influencing the iron cargo load of ferritins and found that this load is highly dependent on the post-mortem interval: the longer the PMI, the fewer iron-loaded ferritin cores found in a sample.

Several ways of iron release from ferritin in the human brain have been described under physiological conditions. Examples are, autophagy-mediated degradation of ferritin to release iron, activated by low iron concentration (Panther et al. 2022), or proteosomal degradation of ferritin triggered by its under-saturation with iron (Panther et al. 2022). Another way of iron release from ferritin can be induced by iron-induced oxidative stress, in a positive feedback loop (MacKenzie et al. 2008; Mills et al. 2010).

While the above mechanisms would have been functioning in the living brain, they would lead to balanced, less drastic changes in iron release than described here during autolysis. In contrast, our findings show a considerable decrease in ferritin concentration with ascending PMI while maintaining an unhindered absolute iron concentration suggesting that the release of iron from ferritin took place post-mortem in our study case.

Autolysis progresses very rapidly during the post-mortem time and leads to the unloading of iron from the ferritins' core. Two factors may contribute to unloading: either the ferritin shell shielding the iron core is damaged during autolysis causing the leakage of its cargo, or the ferritin holoprotein is decayed dissipating the iron. Unloading of ferritin cores explains why the absolute iron concentrations remained independent of the post-mortem time, whereas the number of iron-loaded ferritin cores decreased with increasing post-mortem time. We thus can only speculate as to the cellular mechanisms that govern the dispersion or decaying of ferritin during the post-mortem interval. Further studies will be necessary to detect possible changes in the oxidation state during the unloading or decay of ferritins.

Though it is not possible to extrapolate from post-mortem tissue to living humans, it would be possible to determine their iron distributions using quantitative MRI (magnetic susceptibility mapping or  $T2^*$  mapping), as this is non-invasive, but this does not enable to assign the ferritin to specific cell types. Organotypic brain slice cultures could be the closest option to understanding cellular ferritin distribution and human iron regulation in-vivo, which however are accompanied by both ethical and practical limits of obtaining suitable samples.

Our second aim was to reveal the role of different cell types in preserving ferritin during autolysis. We were able to determine the cell type (neuron, oligodendrocyte, astrocyte, or other glial cells) that contained most (86%) of the ferritins we had detected in five out of the six patients. We found that most of the iron in the human brain is stored in oligodendrocytes, but iron is also found in astrocytes or unidentifiable glial cells and neurons. Their presence in astrocytes is in line with previous light microscopic results demonstrating an increasing number of ferritin-positive astrocytes in the ageing human basal ganglia (Connor et al. 1990). Owing to

the fact that oligodendrocytes are the myelinating cells, more iron maybe is stored in oligodendrocytes since it is used as a co-factor in lipid biosynthesis (Connor and Menzies 1996; Todorich et al. 2009). Earlier studies support our results using a variety of different techniques (Connor et al. 1990; Meguro et al. 2008; Quintana et al. 2004, 2006).

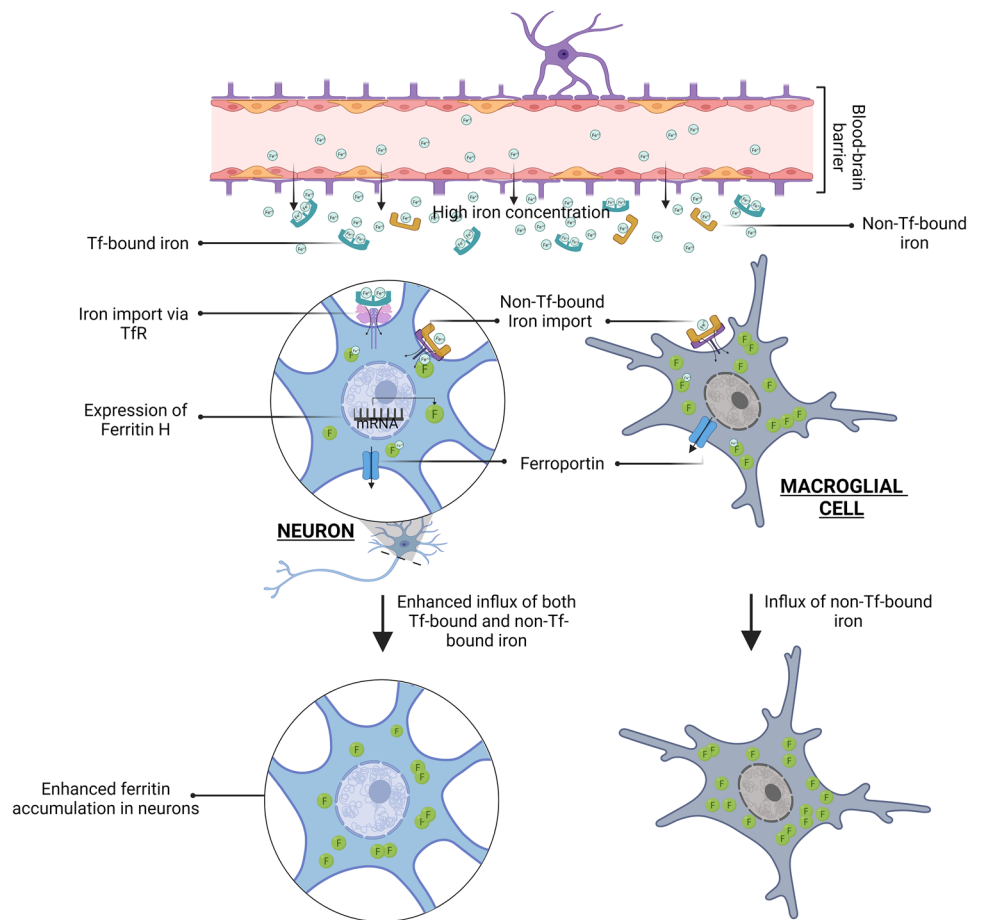
In examining the cell types that are involved in iron-filled ferritin loss during autolysis, we found that the loss occurs at a higher rate in neurons than in glial cells, as schematically depicted in the graphical abstract. Interestingly, we found that the probability of the ferritins being stored in neurons rather than glial cells scales positively with total ferritin concentration in each patient. To our knowledge, this is the first study that shows that the iron-filled ferritin concentration increases faster in neurons than in glial cells with increasing total ferritin concentration. In line with our data, in Parkinson's disease, iron overload has been shown to elicit an upregulation of ferritin synthesis in dopaminergic neurons (Aguirre et al. 2005; Griffiths et al. 1999; Oakley et al. 2007; Vela 2018; Wang et al. 2016).

Two possible mechanisms could explain this difference in loading rate between neurons and glial cells: First, entry of iron into the brain is in its transferrin-bound iron form and via the transferrin receptors (TfR) expressed on the brain capillary endothelial cells (Moos et al. 2007). Once entered, it is distributed between a variety of brain cells depending on the cellular requirement. In our studies, we discovered that the ferritin load increases in neurons when a higher concentration of iron is present in the brain region. Neurons were shown to have a higher ferritin H isoform distribution compared with glial cells (Han et al. 2002; Muñoz et al. 2011). Previous studies have shown that iron mediates the upregulation of ferritin H isoform in neurons (Nahirney and Tremblay 2021). Thus, iron overload might lead to increased production of ferritin H in neurons.

Second, neurons also express iron exporter ferroportin, which allows them to excrete any excess iron. Additionally, neurons express both TfR and divalent metal transporter 1, enabling the import of transferrin-bound iron (Burdo et al. 2001) (Moos and Morgan 2004). On the contrary, macro glial cells (oligodendrocytes and astrocytes) are devoid of TfR in-vivo (Li et al. 2013; Moos et al. 2007; Todorich et al. 2009). The source of iron in macro glial cells is thought to be a non-transferrin-bound form of iron taken up at the blood–brain barrier during the entry of iron into the brain (Moos and Morgan 2004; Todorich et al. 2009). From this, it is conceivable that the presence of TfR, and the ability of neurons to also take up non-transferrin-bound iron (Moos et al. 2007) might prompt neurons to take up more iron compared to glial cells when the total concentration of iron increases in the brain (Fig. 11).

In summary, we found high involvement of neurons in the storage of ferritins at high iron load.

**Fig. 11** Hypothetical schematic of the mechanism of ferritin accumulation in neuron with rise in total iron concentration



Moreover, iron-loaded ferritins are lost post-mortem, in a yet unknown process, and loss of iron from ferritin cores occurs at a higher rate in neurons than in glial cells.

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**Author Contributions** SS, SnR, and GL planned and performed the experiments at the electron microscope, and computed elemental maps and jump-ratios. SS, SnR, GL, and SL analyzed and quantified the raw data from elemental maps. JH, AMTB provided and assessed the brain tissue. StR, SE, and CB performed all qMRI experiments and analysis. WG performed ICP-MS experiments and analysis. PS proofread the manuscript. GL designed the project and study plan. All authors revised, read, and approved the manuscript.

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**Data Availability** The data presented in the manuscript is fully available upon request to the authors.

## Declarations

**Competing interest** The authors have no competing interests to declare that are relevant to the content of this article.

**Ethical Approval** This work was approved by the Ethics Committee of the Medical University of Graz, votum number 28-549 ex 15/16.

**Consent to participate** Not applicable.

**Consent to publish** Not applicable.

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