After 140 years from the discovery of Golgi’s black reaction, the study of connectivity of the cerebellum remains a fascinating yet challenging task. Current histological techniques provide powerful methods for unravelling local axonal architecture, but the relatively low volume of data that can be acquired in a reasonable amount of time limits their application to small samples. State-of-the-art in vivo magnetic resonance imaging (MRI) methods, such as diffusion tractography techniques, can reveal trajectories of the major white matter pathways, but their correspondence with underlying anatomy is yet to be established. Hence, a significant gap exists between these two approaches as neither of them can adequately describe the three-dimensional complexity of fibre architecture at the level of the mesoscale (from a few millimetres to micrometres). In this study, we report the application of MR diffusion histology and micro-tractography methods to reveal the combined cytoarchitectural organisation and connectivity of the human cerebellum at a resolution of 100-μm (2 nl/voxel volume). Results show that the diffusion characteristics for each layer of the cerebellar cortex correctly reflect the known cellular composition and its architectural pattern. Micro-tractography also reveals details of the axonal connectivity of individual cerebellar folia and the intracortical organisation of the different cerebellar layers. The direct correspondence between MR diffusion histology and micro-tractography with immunohistochemistry indicates that these approaches have the potential to complement traditional histology techniques by providing a non-destructive, quantitative and three-dimensional description of the microstructural organisation of the healthy and pathological tissue.

Keywords Diffusion tensor imaging · Tractography · Cerebellum · Post-mortem · Human · MR diffusion histology · Connectome · Multi-scale · Micro-tractography

Introduction

The year 2013 marks the 140th anniversary of Camillo Golgi’s discovery of the black reaction or “reazione nera” [1]. By applying his technique to the cerebellum, Golgi was able to describe the morphology of individual cells and, for the first time, the complexity of their dendritic arborisation and axonal “reticular” architecture. This work paved the way for Santiago.
Ramon y Cajal’s refined studies of histology and cerebellar connectivity and the definitive validation of the neuron doctrine [2, 3]. As a result of these pioneering anatomical works and subsequent physiological studies [4, 5], neuroscientists were able to integrate theoretical work with clinical observations. This led to the development of first computational models of cerebellar networks [6, 7] that were fundamental for understanding the basis for cerebellar physiology and motor control [8] and marked the beginning of modern Computational Neuroscience [9]. Studies of cerebellar connectivity were further extended to extra-cerebellar circuits with the use of viral tracers [10, 11] in animal models, and the demonstration of parallel cerebellar–cortical loops consolidated the link between the cerebellum and frontal lobes. This experimental evidence supported the clinical observation of disorders of cognition and behaviour associated with cerebellar pathology [12, 13]. More recent in vivo neuroimaging studies have used high-resolution structural MR imaging and high-field MR scanners to depict very fine details of the human cerebellar anatomy [14, 15], whereas diffusion-based tractography has allowed the study of major cerebellar pathways [16] in autism, schizophrenia and other disorders [17–21].

However, a fundamental gap between histological and current in vivo neuroimaging studies remains. While a detailed cytoarchitectural description of human cerebellar histology has long been established, a seamless integration with larger scale networks is incomplete. The study of connectivity poses a multi-scale problem ranging from major white matter tracts at the macroscale (centimetres to millimetres) to individual axons or even single synaptic connections at the micro/nanoscale (micrometres to nanometres). In between these two extremes is the mesoscale (millimetres to micrometres) comprised of cell assemblies and small groups of axons [22–24]. Each level of organisation requires appropriate tools to decipher connections, and current methods are not suitable to visualise connections at all levels.

Traditional fibre-tracing studies provide a reliable identification of neuronal connections at the microscale (microscopy) and in some cases at the nanoscale (electron microscopy), but are limited to these scales due to the thin sectioning in two dimensions [25]. Therefore, tracing over long distances is a major challenge, and three-dimensional reconstructions are problematic due to tissue distortions, small sample sizes and cutting artefacts [26, 27]. MRI-based tracers, such as manganese, can overcome some of these limitations [28] and provide an additional macroscopic assessment of fibre tracts, but are limited in providing a system view and cannot be applied to humans due to its neurotoxicity [29]. Although histological tracing studies in primates have been instrumental in establishing basic principles and evolutionary aspects of brain development, it remains difficult to draw reliable inferences about fibre tracts in humans based purely on those from other species [30–33]. This is particularly relevant in light of evidence suggesting a role of the cerebellum in higher cognition and behaviour that is unique to humans (e.g. language and executive functions) [11, 34–36].

Non-invasive white matter tracking in humans is routinely performed using diffusion-based MRI [37]. Here, the anisotropic movement of water molecules along fibre tracts allows a tractographical reconstruction of major white matter trajectories and offers the possibility of mapping large-scale networks in the living human brain [38–42]. However, two major limitations affect in vivo diffusion imaging today: the low spatial resolution of diffusion data achievable with current clinical hardware and the lack of histological validation for tractography and diffusion indices. Low spatial resolution limits current investigations to main brain features of white matter pathways and sometimes this results in partial or incorrect descriptions of the underlying tissue microstructure due to the amalgamation of properties from different tissues within the same voxel [43]. Also, little is known about the true correspondence between tractography reconstructions, diffusion indices and the underlying tissue organisation that can be determined, for example, from histology studies in healthy and pathological tissue [44]. In vivo methods are therefore currently unable to provide an adequate description of the small and local cortical networks between close regions leaving the mesoscale a largely unexplored domain.

A way to overcome these limitations is to perform diffusion imaging on the post-mortem human brain using preclinical scanners. By acquiring post-mortem data, imaging results can be promptly validated with histological analyses, and it is possible to investigate the direct correspondence between the diffusion signal and the real underlying microstructure [44–46]. Post-mortem imaging also removes some of the intrinsic time constraints of in vivo imaging, allowing significantly longer scan times that, combined with the more powerful hardware of preclinical scanners, give access to much higher spatial resolutions and signal to noise ratios [47–49]. Previous studies using ex vivo human tissue have shown that “grey matter” characteristics, such as cell layers, can be revealed using diffusion-based MRI [49]. In this study, we extend these results and show that diffusion MRI can be further applied to describe the combined cytoarchitectural organisation and connectivity characteristics at the mesoscale level. By combining complementary information about the microstructural organisation of the tissue, including fractional anisotropy (FA), apparent diffusion coefficient (ADC) maps and diffusion-weighted images (DWI)/T2-weighted images, we have created novel MR diffusion histology maps. When applied to a fixed post-mortem human cerebellum, these were able to show different mesoscale laminar features and their structural properties. To validate these results, we verified our findings with immunohistochemistry images of the corresponding region and with the known cytoarchitectural organisation of
the cerebellum. Additionally, using tractography reconstructions based on diffusion tensor imaging (DTI) data, we have reconstructed, for the first time, the main trajectories of axonal projections within individual cerebellar folia and exposed the fine details of intra-cortical structural organisation within the cerebellar cortex. Advantages and limitations of these approaches are finally presented in the discussion.

Methods

Post-mortem Tissue

The cerebellum of an 11-year-old female was made available from the Clinical Neuropathology Department at King’s College Hospital under the hospital post-mortem consent for medical and genetic research. The brain was extracted 45 h after death from acute haemorrhage due to a pilocytic astrocytoma of the spinal cord (grade I) and fixed by immersion in 10% neutral buffered formalin for 4 weeks. After fixation, the cerebellum was separated from the brainstem by severing the cerebellar peduncles. MR images were initially acquired from the entire cerebellum and then from smaller samples. The time from death to scan, or scan interval [48], was 3 months. To minimise long-term tissue changes. After imaging, the samples were returned to the Clinical Neuropathology Department for immunohistochemistry analysis.

MRI Acquisition

MRI data were acquired using a 7T Varian/Agilent (Palo Alto, USA) system using a 105/60 HD gradient system and a 39-mm quadrature volume RF coil. A Fast Spin Echo (FSE) T2-weighted sequence was acquired to provide a high-resolution structural reference images for planning subsequent diffusion imaging scans (TR=4,000 ms, TE=10 ms, averages=16, FOV=30×30 mm, matrix=128×128, 45 slices at 0.6 mm thickness and no gap). Diffusion MRI data were acquired using a Pulsed Gradient Spin Echo sequence (TR=3,600 ms, TE=36 ms, G=500 mT/m, δ=4 ms, Δ=15 ms, b value≈4,200 s/mm², averages=170, 3 non-diffusion-weighted volumes (b0) and 30 diffusion-weighted volumes/directions, FOV=51.2×6.4 mm, matrix 512×32, 65 slices at 0.1 mm thickness and no gap). Volume planning was set up to obtain the highest spatial resolution (100 μm in plane) orthogonal to most of the cerebellar folia as visible from the FSE images. Saturation bands were placed at the limit of the sample FOV along the phase direction. SNRb0 in the tissue was above 12. Total scan time was approximately 140 h.

MRI Data Processing

Diffusion data were processed using ExploreDTI (www.exploredti.com). Data were first pre-processed correcting for eddy current distortions and for long-term movements of the sample. Diffusion tensors were estimated using the full b-matrix, including the effect of the imaging gradient and applying a non-linear least square approach [50]. Mean ADC, FA, non-diffusion/T2-weighted (T2w) image and average DWI maps were generated. Tractography was performed using an Euler-like streamline algorithm with a step-size of 50 μm and the entire imaged sample as seed region. FA thresholds were set to 0.07 to initiate and continue tracking. The lower FA threshold relative to the usual in vivo threshold (e.g. FA=0.15/0.20) was required due to the intrinsically low diffusivity and anisotropy of the formalin-fixed sample. Angular threshold was set equal to 45°.

Immunohistochemistry

After cutting histological sections (50 μm) on a freezing microtome, sections were incubated overnight (21 °C) to detect white matter based on rat anti-myelin basic protein (MBP, 1:100, Abcam), neurons based on rabbit anti-neurofilament 160 kD (1:1,000, Abcam), Purkinje cells using a rabbit anti-calbindin (1:200, Abcam), and Bergmann glia using mouse anti-glial fibrillary acid protein (GFAP, 1:3,000, Sigma) primary antibodies. Appropriate secondary antibodies (Alexa350, 488 and 546, Molecular Probes) were applied for 1 h prior to three washes and coverslipping with Vectashield for fluorescence (Vector Labs). Where appropriate, Hoechst was used as a blue nuclear stain. Sections were visualised on a Zeiss Axioplan.

Results

Cytoarchitectural Basis of Diffusion Anisotropy

In the cerebellum, key neuroanatomical features of each folium, which represent the smallest structure on the surface of the cerebellum, consist of a white matter medullary layer (WM), an inner granule cell layer (cortical layer III), a middle Purkinje cell layer (II) and an outer molecular cell layer (I). These features vary in size from 25–40 μm for the Purkinje cell layer to 1 mm for the larger strand of white matter within a folium. With a spatial resolution of 100×100×200 μm, the data acquired in our experiment allow the visualisation of mesoscale histological features within individual cerebellar folia (Fig. 1). At this resolution, DWI, ADC and FA maps reveal different cytoarchitectural aspects of cerebellar
organisation and offer the opportunity to perform quantitative analysis of individual layers. For example, as the axonal white matter (WM) projections from the cerebellar folia disperse within the granule cell layer, the ADC increases while the FA decreases, suggesting an increase of water mobility due to changes in the structural organisation of fibres (e.g. loss of parallel pattern, branching, and reduced myelination). In the molecular layer (I), a uniform increase in FA and decrease in ADC is observed, suggesting a higher cellular density and possibly a more ordered axonal packing within this layer. By pseudo-colouring each MR image and merging these into a single image akin to fluorescence histology, an MR diffusion histology image was created to highlight anatomical features, such as the laminar organisation of the cerebellar cortex and tissue compartmentalisation between white and grey matter. These results appear consistent and uniform across the entire imaged sample. Direct comparison of MR diffusion histology images with immunohistochemical images of the same region corroborates these findings and confirms that both imaging techniques visualise similar cytoarchitectural features of the cerebellar folia, although at a different spatial resolution (Fig. 1).

The cellular layer specificity of these MR images is further shown by a quantitative analysis of changes in signal intensity (Fig. 2). The ADC of the granule cell layer (III) uniformly increases starting from low ADC values close to white matter and reaches a maximum in proximity or at the level of layer II. The high diffusivity (i.e. high water molecular mobility) in this region may suggest a decrease of tissue density or an increase of extracellular space. It is worth observing that this increase in ADC partially resembles the decrease of MBP contrast observed in the granular layer as show in Fig. 1. ADC sharply decreases within the molecular layer suggesting a sudden change in microstructural organisation that reduces the average mobility of water in this layer. The intermediate Purkinje cell layer (II) is consistently visible on T2-weighted images as a brighter separation between the larger layer III and layer I. In the MR diffusion histology image, however, this layer is not always visible due to partial volume effects and ADC values similar to layer III.

Tractography of Cerebellar Pathways from Macro- to Mesoscale

The ability to provide a three-dimensional description of the microstructural organisation of biological tissues is a unique feature of diffusion imaging. However, conventional in vivo

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**Fig. 1** MR diffusion histology and immunohistochemistry. The cytoarchitectonic features of the cerebellar cortex are shown by diffusion imaging maps at 100 μm resolution. White matter (WM), granular layer (III), Purkinje cell layer (II) and molecular layer (I) can be discriminated across the different T2-weighted images, diffusion-weighted images (DWI), apparent diffusion coefficient (ADC) and fractional anisotropy (FA) contrasts (scale bar=1 mm). A pseudo-colour visualisation of these maps is obtained by combining together FA (red), DWI (green) and ADC (blue) maps to mimic immunohistochemistry results and obtain a similar MR diffusion histology map. The comparison between MR diffusion histology and immunohistochemistry of the same region shows a good matching of features. Specifically, white matter tracts correspond to regions rich in myelin basic protein (MBP) and neurofilament, whereas a high cellular density, as indicated by the nuclear dye Hoechst, characterised the granular cell layer (III) (scale bar=250 μm). These cells were predominantly neurons (NeuN+ cells), whereas the Purkinje cell layer (II) was defined by calbindin+ cells. However, the homogeneous distribution of astrocytes (GFAP+ cells) was not associated with MRI-detectable features.
or post-mortem tractography performed with a relatively large voxel size can only describe macroscale features and does not provide a level of detail that can be integrated with microscale histology because multiple histological features are mixed within a single voxel (Fig. 3a). In contrast, microtractography performed at 100 μm allows the visualisation of mesoscale features that can be directly integrated with the histological organisation of the cerebellum. At this resolution, it is possible to visualise white matter tracts running into individual cerebellar folia and dispersing inside the granule cell layer. It is also possible to reconstruct cerebellar parallel fibres as a structural continuum of parallel orientations running inside the molecular layer (Fig. 3b). In this figure, we show all streamlines passing through either a region of interest (ROI) delineating the molecular layer or through a ROI defining the white matter medullary layer. Both ROIs were selected on two consecutive image slices at the centre of the sample and no exclusion ROIs were applied.

At a deeper level, micro-tractography provides a glimpse of the microstructural organisation within an individual layer. For example, the three-dimensional description offered by tractography consistently reveals the presence of repeated intra-cortical connections between the granular layer and the molecular layer as short strands that merge almost orthogonally with longer strands representing the parallel fibres (Fig. 4). At this resolution, these reconstructions are likely to be representative of the anatomy of a small group of hundreds of granular cell axons that coherently run within each voxel. Importantly, these tractography reconstructions follow the anatomical trajectories of real fibres (i.e. granular cell axons) that as short strands bend almost orthogonally at the level of the T-junctions to continue as parallel fibres inside the molecular layer. Although Purkinje cell bodies may also play a role in this reconstruction, their large cytoplasm (∼50 μm diameter) and lower density make them virtually undetectable inside the dominant diffusion component. In this figure, a first ROI was delineated around the white matter medullary layer to visualise only those streamlines passing through the white matter and the granular layer (green coloured). An exclusion ROI was used to exclude those streamlines reaching the molecular layer. Vice versa short strands and long strands (red coloured) were visualised selecting all streamlines passing through a ROI in the molecular layer. In this case, one exclusion ROI in the white matter medullary layer was used to exclude streamlines reaching the inner layer.

**Discussion**

Although a detailed cytoarchitectural description of the human brain by means of histology is long established, an integrated view of brain and cerebellar connections ranging from major white matter pathways to individual neurons or even small fascicles of a few hundred axons is still largely missing [22–24, 32]. In this preliminary study, we have shown that high-resolution MR diffusion imaging methods applied to fixed ex vivo human cerebellum have the potential to extend our ability to study brain connectivity, from a macroscopic description down to the mesoscale level, and at the same time provide results that can be directly compared with histological features observed at the microscale.

Conventional in vivo diffusion imaging applications acquire data with isotropic spatial resolutions of 2 mm (8 μl/voxel). Data acquired in our experiment had a spatial resolution of 100×100×200μm (2 nl/voxel), equivalent to a voxel volume
Fig. 3 Tractography from macro- to microscale. a A conventional clinical DTI scan at a resolution of $2 \times 2 \times 2$ mm (8 μl/voxel) reveals the directionality of major fibre tracts overlaid onto the main macroscopic anatomical features of the cerebellum. On a fixed cerebellum acquired at $500 \times 500 \times 500$ μm (0.25 μl/voxel, 32 times smaller voxel than clinical scan), it is possible to further focus on the cerebellum to reveal a far greater number of anatomical details and connections. b However, it is at a resolution of $100 \times 100 \times 200$ μm (2 nl/voxel, 4,000 times smaller voxel than on a clinical scan) that it is possible to clearly observe mesoscale connectivity features of white matter and histological details of the cerebellum grey matter. The arbor vitae (1), intra-folium axonal projections (2), granular cell layer (3), as well as molecular cell layer (4) are all identifiable within each cerebellar folium using MR diffusion histology and micro-tractography reconstructions.

Fig. 4 Cytoarchitectural tractography. Features of the cerebellar intracortical connectivity are revealed by micro-tractography in a portion of cerebellar folium. White matter (WM) trajectories propagate and disperse inside the granular layer (III) of the folium. At this interface, short strands, reflecting probably the average orientation of granular cell axons, project to the molecular layer (I) and merge with almost orthogonal and longer strands that consistently describe the dominant orientation of the parallel fibres (II) inside this layer.
4,000 times smaller than those acquired in clinical DTI scans. At this resolution, it was possible to start to investigate the cytoarchitectural basis of diffusion properties related to distinct histological components of cerebellar grey matter. Differences in ADC and FA between the white matter, granular layer and molecular layer were consistent with our knowledge of the microstructural and histological organisation of the cerebellum (for a review see [51]). Additionally, the comparison between MR diffusion histology maps and immunohistochemistry results showed a good agreement in identifying the same mesoscale features of the cerebellar cortex. We think that this approach, thanks to its ability to characterise cytoarchitectural organisation over relatively large brain samples, offers a new imaging tool for systematically investigating the microstructural and histological organisation of the cerebellum. While offering new insights at a range of scales not currently accessible with conventional clinical MRI scanners, these results could potentially integrate and complement other modalities to allow better interpretation of in vivo clinical and functional results. Moreover, due to the non-destructive imaging procedure, this technique can allow histological analyses of the same imaged sample and the study of the direct correspondence between the formalin-fixed cerebellum that required an uninterrupted scan time of approximately 6 days. This acquisition clearly represents a special case that cannot represent the standard acquisition protocols applicable to a large number of samples. Previous diffusion imaging studies on animals and also human samples provide important guidelines on how to sensibly reduce scan times and improve signal quality [59–62]. Our group is currently investigating these approaches, and future work will optimise novel and more specific sample preparation procedures together with new acquisition strategies.

Another important limitation is the long acquisition time of the formalin-fixed cerebellum that required an uninterrupted scan time of approximately 6 days. This acquisition clearly represents a special case that cannot represent the standard acquisition protocols applicable to a large number of samples. Previous diffusion imaging studies on animals and also human samples provide important guidelines on how to sensibly reduce scan times and improve signal quality [59–62]. Our group is currently investigating these approaches, and future work will optimise novel and more specific sample preparation procedures together with new acquisition strategies.

The ability to visualise the three-dimensional organisation of the mesoscale represents a unique feature of diffusion imaging and offers a new approach for investigating the anatomical basis of connectivity within the cerebellum and its complex functions [16]. The role of cortico-cerebellar connectivity in cognition, for instance, is still under investigation and represents a topic of intense debate and interest. Data on the precise anatomy of the intracerebellar connectivity in humans and the pathways that relay information from the cerebellum to associative cortex could help to confirm the involvement of the cerebellum in higher cognitive functions [31, 63, 64]. Mapping white matter pathways from the macro- to microscale could help create a more detailed atlas of the cytoarchitectural organisation and connectivity not only of the human cerebellum [65–67] but also of the cerebral cortex in general [68].
Conclusions

A detailed analysis of cerebellar connectivity could reveal unique features of human anatomy and address important questions related to the role of the cerebellum in higher cognitive functions. MR diffusion histology and micro-tractography represent two novel methods for obtaining a three-dimensional description of tissue cytoarchitecture at the mesoscale level. These methods produce quantitative information that, coupled with high-resolution visualisation of small fibres, can bridge the gap between large-scale network mapping and microscopic histology. We believe that this approach, applied to fixed post-mortem samples, represents an essential step forward in the understanding of the human cerebellum and its connectivity with brainstem nuclei and cerebral cortex. Future applications of these methods promise to have practical implications in the diagnosis and treatment of cerebellar disorders.

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Conflict of Interest The authors declare no conflict of interest.

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