# The Potential of Myelin-Sensitive Imaging: Redefining Spatiotemporal Patterns of Myeloarchitecture

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### **ABSTRACT**

Recent advances in magnetic resonance imaging (MRI) have paved the way for approximation of myelin content in vivo. In this review, our main goal was to determine how to best capitalize on myelin-sensitive imaging. First, we briefly overview the theoretical and empirical basis for the myelin sensitivity of different MRI markers and, in doing so, highlight how multimodal imaging approaches are important for enhancing specificity to myelin. Then, we discuss recent studies that have probed the nonuniform distribution of myelin across cortical layers and along white matter tracts. These approaches, collectively known as myelin profiling, have provided detailed depictions of myeloarchitecture in both the postmortem and living human brain. Notably, MRI-based profiling studies have recently focused on investigating whether it can capture interindividual variability in myelin characteristics as well as trajectories across the lifespan. Finally, another line of recent evidence emphasizes the contribution of region-specific myelination to large-scale organization, demonstrating the impact of myelination on global brain networks. In conclusion, we suggest that combining well-validated MRI markers with profiling techniques holds strong potential to elucidate individual differences in myeloarchitecture, which has important implications for understanding brain function and disease.

https://doi.org/10.1016/j.biopsych.2022.08.031

Myelin ensheathes axons of the central and peripheral nervous systems, providing the structural basis for fast and stable impulse propagation (1). The process of myelination is highly dynamic, involving rapid changes after even a few hours of a task and continuous reorganization across the lifespan (2-6). Indeed, the dynamic nature of myelin is thought to be central to its role in enabling flexible responses to rapidly changing environments and to maximizing the efficiency of neural communication (1). Elucidating the temporal dynamics and spatial patterns of myelination has been a topic of interest for over a century (7,8). As myelination throughout life is activity dependent (9), it has been theorized that the distribution of myelin reveals privileged microcircuits in the cortex that are relevant to understanding the structural basis of specific functions (10). Defining region-specific trajectories of myelination is thought to inform upon the maturational sequence of functional specialization in the brain (7,11), while region-specific breakdown of myelin in older age may reveal the structural underpinnings of cognitive decline (12). Classically, the spatial distribution of myelin has been described from postmortem study (13-16), but such approaches are inherently limited in characterizing intraindividual changes across time. Uncovering the spatiotemporal patterns of myelin necessitates in vivo imaging of myelin.

Magnetic resonance imaging (MRI) holds promise for enabling in vivo histology, whereby the cellular composition of living tissue may be discerned noninvasively. Crucially, MRI allows large, longitudinal cohort studies to track individual trajectories of tissue changes (17–19). Nevertheless, challenges remain for adoption of MRI for myelin mapping, as many previous approaches have been limited to indirectly measuring myelin via magnetic fields and confronted by the apparent discrepancy between imaging resolution and the size of myelin. New imaging sequences, contrasts, and biophysical models are helping to address these limitations, though not all MRI-derived myelin markers are equally valid. As interest in myelin mapping increases across foundational and clinical neuroscience, it is high time to discuss the validity of emerging techniques and identify promising avenues for future work.

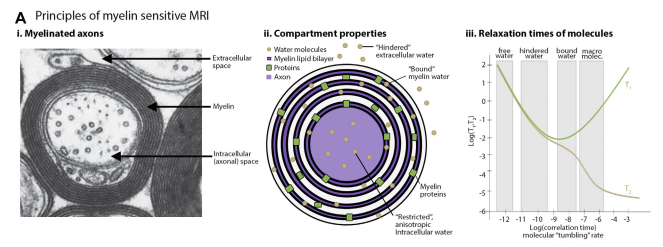
In this review, we aim to demonstrate the significant contribution that myelin-sensitive imaging can make to understand the spatial patterns and dynamic changes of myeloarchitecture in the human brain. First, we lay the groundwork for how myelin markers are derived from MRI. Considering theoretical and histological validations, several multimodal approaches are highlighted that benefit disambiguation of myelin from other neurobiological factors. Next, we discuss the emergence of myelin profiling techniques, which provide nuanced characterization of the myeloarchitecture of cortical regions and white matter bundles. Finally, we highlight how several recent studies have capitalized on methodological improvements and in doing so have

advanced our understanding of how myeloarchitecture evolves across the lifespan.

# DEVELOPING AND VALIDATING IN VIVO MYELIN MARKERS

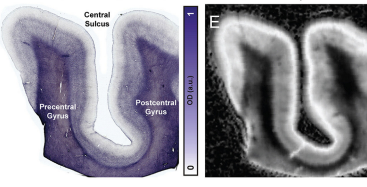
Myelin, the lipid-rich material that insulates nerve fibers, is amenable to MRI measurement because it contributes to key determinants of MR relaxation times: water mobility and the interaction between water and macromolecules (20,21) (Figure 1A). Bound water produces shorter longitudinal (T1) and transverse (T2) relaxation times than free water (e.g., in

cerebrospinal fluid) (Figure 1Aiii). As the primary location of bound water in the brain, myelin water has been shown to have distinctively short T1 and T2 relaxation times (30,31). However, looking across brain tissue types, the dominant source of T1 contrast are lipids (32). Cholesterol and cerebroside, in particular, both rich in myelin, are related to T1 shortening (20,33), which produces the distinctive appearance of gray and white matter in T1 images. In contrast, diffusion-weighted imaging targets anisotropy of hindered water (e.g., in axons or extracellular spaces), using multiple diffusion echo gradients to sensitize the MR signal to the random motion of water molecules. While myelin modulates anisotropy, for example by



## **B** Validation of an MRI marker of myelin

## i. Direct comparison of myelin basic protein stain and quantitative T



## ii. Cross-comparison of stains and contrasts

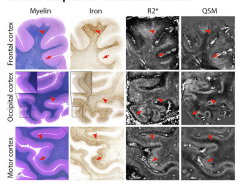


Figure 1. Sensitivity of magnetic resonance imaging (MRI) contrasts to myelin. (A) (i) Electron micrograph of a myelinated axon in the central nervous system (22). (ii) The schematic depicts distinctive features of myelinated axons that make them susceptible to MRI. [Adapted from Min et al. (23).] Myelin is composed of lipids (70%–85%) and proteins (15%–30%) (22). Water molecules are bound in the myelin sheath, which contrasts with the more motile water molecules in intra- and extracellular spaces [described as restricted and hindered water, respectively (24)]. (iii) T1 and T2 vary as a function of molecular tumbling rate (defined by correlation time) (25). Thereby, water in different compartments as well as macromolecules may be distinguished by T1 and T2. Positions of molecules along the x-axis are approximate (26). While T2 decay of macromolecules is too quick to be captured by human MRI scanners (26), the difference in T2 decay of water in different compartments can enable identification of myelin water [see, for example, (27)]. (B) (i) An exemplar validation study, Stüber et al. (28) showed the similar pattern of myelin basic protein staining and quantitative T1 in the same tissue. (ii) Cross-correlation of multiple stains and multiple contrasts can help disambiguate the contribution of different neurobiological features to an MRI marker. Here, Hametner et al. (29) show that iron is linearly related to R2\*, but iron and myelin interact to determine quantitative susceptibility mapping (QSM). For instance, in the frontal cortex, high myelin and high iron produce moderate QSM (top arrow), but low myelin and low iron also produce moderate QSM (lower arrow). In other words, the myelin and iron appear to cancel each other out, related to the diamagnetism of myelin (lowers QSM) and paramagnetism of iron (increases QSM). Notably, the study showed that removing the contribution of iron from the QSM resulted in a strong correlation of QSM with myelin.

decreasing the permeability of axons, its effects on water diffusion are minimal (34). Therefore, classic diffusion-based measures are not posited as specific proxies for myelin. Turning to the macromolecular makeup of myelin, its protein and lipid composition render it diamagnetic. This distinctive property may be detected by combining the magnitude and phase maps from a gradient echo sequence, known as susceptibility mapping (35,36). This technique can identify susceptibility anisotropy created by ordered molecular structures such as myelin sheath, offering a new opportunity for clinical phenotyping of neurological disorders such as multiple sclerosis, Parkinson's disease, Alzheimer's disease, and Huntington's disease (35–38).

While the spatial resolution of myelin-sensitive MRI techniques has become increasingly higher with advanced hardware and sequence development [~500 μm, (39)], it is still at a relatively coarse level, such that each voxel likely includes multiple tissue types, contaminating the purity of its signal. Biophysical models can theoretically disentangle such heterogeneous signal sources and potentially enhance specificity to myelin. For example, myelin water fraction (MWF) is based on the fact that in the central nervous system, the T2 signal decay follows a multiexponential curve (30). By fitting the decay curve using least-square methods, these models can distinguish the myelin water pool, with ultrashort T2 decay, from the nonmyelin water pool (Figure 1Aiii). In contrast, magnetization transfer (MT) indirectly measures myelin based on the exchange and crossrelaxation between macromolecules (found predominantly in myelin) and water (40,41). Longitudinal relaxation rate (R1, also reported as 1/T1) is similarly driven by the cross-relaxation of lipids and water (33). Overall, this nonexhaustive summary of imaging contrasts and biophysical models serves to demonstrate that the influence of myelin on water mobility, as well as the lipid composition of myelin, allow sensitization of the MR signal to myelin in various ways with different assumptions, complexity, and presumed specificity.

Empirical validation has been essential to support the myelin sensitivity of the MR markers described above. The principal approach for validation is comparison with myelinspecific staining in the same tissue. Early studies described the correspondence of T1-weighted imaging contrasts with myelin distribution, such as distinctive myeloarchitectural features (e.g., stria of Gennari) or highly myelinated areas (e.g., primary sensory areas and middle temporal visual area) (42-46). The potential of myelin-sensitive imaging was further established by showing reduced values in individuals with multiple sclerosis (47-49), nonhuman animals without myelin basic protein (shiverer mouse) (50,51) or myelin proteolipid protein (shaking pup) (52), and rodents with cuprizole-induced demyelination (53,54). More recently, statistical approaches have been used to estimate the correlation between MRI markers and the degree of myelin staining, comparing many matched samples across regions and/or across individuals (Figure 1Bi). Collating such studies, several recent metaanalyses have found that many MRI markers are positively correlated with myelin content (55-57). MWF and MT approaches exhibit the highest sensitivity to myelin, but effect sizes are heterogeneous. The heterogeneity in estimates, which produces wide confidence intervals and overlapping prediction estimates across MRI markers, suggests that no

single MRI marker may currently be considered the ideal myelin measure (55-57). Further studies are necessary to determine the interaction of MRI markers with influential methodological parameters, such as histological processing, sampling in gray or white matter, and statistical design. Alternative approaches include comparing MRI markers to oligodendrocyte- or myelin-related genes derived from brain transcriptomics. Such comparisons performed using T1weighted/T2-weighted (T1w/T2w) scans suggest enrichment for myelin-related genes, but the association with myelin is not specific, nor is it the strongest enrichment present for T1w/T2w (58,59). This approach has also been extended to other MRI markers such as MT and MWF by comparing transcriptomic maps in one set of subjects [e.g., Allen Human Brain Atlas; (60)] to imaging maps in another set of subjects. Associations of these MRI markers with myelin-related genes and oligodendrocytes have been reported, though effect sizes tend to be small (r < 0.4) (61,62), as may be expected due to the confounding influence of interindividual differences, especially when the samples are not age matched.

The utility of myelin-sensitive imaging is also dependent on reliability. Test-retest reliability is high (intraclass correlation coefficient > 0.8) for many proposed myelin markers, including T2-derived MWF, quantitative MT, R1, and calibrated T1w/T2w ratio, but more moderate for other markers (intraclass correlation coefficient 0.5-0.8; e.g., MTR and raw T1w/T2w) (63-68). While supporting a certain level of consistency of these markers, the estimates are mostly derived from moderate sample sizes (average n = 24). More systematic efforts to optimize these metrics based on a larger population and different sites are therefore required to confirm the high reliability of their use in an actual clinical setting. Notably, recent work has demonstrated that T2-derived MWF, quantitative MT, and R1 have good intersite reliability (65,67,69). This evidence further bolsters their potential to become reproducible and clinically translatable biomarkers that are especially relevant to neurological disorders with a well-defined myelin pathology, such as multiple sclerosis, and psychiatric disorders with likely myelin alterations, such as schizophrenia (70).

Future work may benefit from multimodal imaging, whereby the combination of several MRI markers can enhance in vivo approximation of myelin distribution. Indeed, discrimination of normal myelin, demyelination, and remyelinated lesions in mice is improved by using the combination of 3 contrasts (T1w, T2w, and MTR; 95% accuracy) (71). The combination of susceptibility mapping with T2\* (or R2\*) can help to disambiguate myelin from iron (72) (Figure 1Bii). Myelin and iron often colocalize in the cortex and additively contribute to certain MR contrasts (e.g., increase in R2\*) (73). However, they have opposing effects on resonance frequency (diamagnetic and paramagnetic, respectively), which may be detected by susceptibility mapping, and differ across tissue types, such as deep gray matter versus white matter (72). Consequently, contrasting R2\* with susceptibility maps may provide a more specific MRI marker of myelin (29,74,75). Notably, other multimodal approaches, such as T1w/T2w (46) and T2\*/B0 (76), use the commonalities of myelin and iron to characterize cortical microstructure with indiscriminate applicability to either source.

Multimodal protocols can also enable quantification of myelin sheath thickness relative to the thickness of myelinated

axon, namely the g-ratio (77). Known to be influential on conduction speed (78–80), the g-ratio is assessed using axonal volume and myelin volume fractions, which may be proxied in vivo using diffusion models, such as neurite orientation dispersion and density imaging (24), with myelin-sensitive imaging, such as MT. Notably, g-ratio values derived from MRI closely approximate g-ratio values calculated with electron microscopy, outperforming estimates of axonal volume and myelin volume fractions that come from each imaging modality separately (81). Therefore, while multimodal imaging approaches require extra consideration of theoretical assumptions and modality-specific distortions (82), this work demonstrates that combining modalities can benefit quantitative evaluation of myelin in vivo.

Overall, sensitivity, specificity, and reliability are important considerations in selecting a myelin-sensitive imaging sequence. However, these must be balanced against more practical requirements that vary across studies, such as acquisition time, available field strength, and the desired spatial resolution. R1 mapping has emerged as a simple and efficient method for myelin-sensitive imaging, with whole-brain coverage in only 8 minutes (magnetization-prepared 2 rapid acquisition gradient-echo: 1 mm on 3T, 0.7 mm on 7T) (67,83). Higher signal-to-noise ratio, spatial resolution, and image sharpness of R1 can be achieved using a recently developed multishot, inversion-recovery, echo planar imaging sequence, though the acquisition time is longer (20 minutes at 0.5 mm on 7T) (84). Toward multimodal imaging, the multiparameter mapping sequence provides 4 whole-brain quantitative maps (proton density, MTsat, R1, and R2\*) in approximately 20 minutes (1 mm on 3T, 0.5 mm on 7T) (85,86). By reducing the resolution of the multiparameter mapping sequence to 1.6 mm on 3T, the acquisition time may be reduced to less than 10 minutes (87). Finally, recent developments with fast acquisition with spiral trajectory T2 imaging allow for whole-brain MWF, which exhibits high variability across white matter (88), to be captured in 10 minutes (1 mm on 3T) (89).

### **MYELIN PROFILING**

A major advantage of myelin-sensitive MRI is the ability to explore myeloarchitecture in 3 dimensions. Myelin profiling, the measurement of myelin along biologically meaningful anatomical axes, is inspired by classic histology but benefits significantly from the 3-dimensional nature of MRI. While profiling in histological sections must follow the cutting plane, myelin profiling with MRI enables characterization of myelin along the natural courses of the cortex (from pial to white matter boundary) and along white matter tracts.

The origins of myelin profiling can be traced to the start of the 20th century. Around that time, several researchers developed myeloarchitectonic atlases of the cortex (8,15,16,90). In each case, cortical areas were delineated with respect to the vertical arrangement of myelinated fibers (Figure 2Ai). Myelinated fibers produce distinctive striations in the cortex, allowing categorization of types and the approximation of areal borders [see (97) for an excellent review]. Beyond qualitative types, histological studies have also used a photometric slice-capturing technique by which myeloarchitecture may be more quantitatively compared across the

cortex based on myelin density across cortical depths (98). Hopf (91,92,99) showed that this quantitative approach allows comparison of distributed areas, elucidating large-scale patterns of myeloarchitectural change (Figure 2Aii). In particular, he demonstrated that myelin content decreases with distance from primary sensory and motor areas, with the lowest levels in the paralimbic cortex, such as the medial orbitofrontal cortex. Legacy data from these classic studies has the potential to serve as a histological gold standard to validate contemporary in vivo myelin-sensitive imaging. However, the accessibility of such datasets has been severely limited, in part due to their qualitative reporting and the 2-dimensional illustrations of the brain that are incompatible with modern neuroimaging. Recently, leveraging seminal meta-analyses of the Vogt-Vogt school (93), a myelin-based cortical parcellation was generated for neuroimaging analysis, whereby the parcel boundaries represent histology-derived estimates of myelin (Figure 2Aiii). Notably, the atlas also incorporates intracortical myelin profiles of multiple cortical areas based on Hopf's photometric studies, providing a unique resource to bridge ex vivo and in vivo imaging studies (94) (https://bic.mni.mcgill.ca/~noel/noelmyelin).

Depthwise profiling of cortical myelin, pioneered in histological studies, has been increasingly adopted by in vivo imaging (100-102). In a typical workflow, cortical surfaces are segmented using a standard T1w image, multiple intracortical surfaces are generated between the pial and white matter boundaries, and then the intensities of a coregistered myelinsensitive image are sampled along the intracortical surfaces at matched vertices [code for protocol may be found in (103)] (Figure 2Bi). While depthwise profiling is agnostic to cortical layers, it is important to note that the vertical arrangement of cells and myelinated fibers varies with cortical curvature (104). As such, depthwise profiling approaches can use equivolumetric surface generation to account for these effects and minimize the influence of curvature on profiles (105). Another key concern is resolution. Dinse et al. (39) showed that distinctive features within intracortical profiles, such as turning points, disappear at lower resolutions (toward 1 mm) (Figure 2Bii), but areal differences remain evident. Areal differences are more pronounced at 0.4- to 0.5-mm resolution, which is increasingly feasible for whole-brain neuroimaging studies [e.g., 7T magnetization-prepared 2 rapid acquisition gradient-echo (83)]. Furthermore, Dinse et al. (39) showed that areas are best discriminated by combining mean intensity and profile shape differences rather than using either feature alone, reinforcing the benefit of using profiles to describe myeloarchitecture. These features (mean and shape) represent unique axes of cortical differentiation (Figure 2Biii) (95). Mean decreases with distance from primary sensory and motor areas and is lowest in frontal and temporal poles, in line with histological evidence (91,92). Shape is often summarized by profile skewness, a parameter adopted from cytoarchitectural histology (106), which pertains to the balance of intensities in upper versus lower layers. For MT-derived intracortical profiles, primary sensory-motor areas exhibit negative skewness, which is related to the gradual increase in MT across cortical depths, whereas cingulate and inferior temporal areas exhibit high skewness, which is related to relatively flat profiles with a sudden uptick in MT in the deepest layers (Figure 2Biii).

#### A INTRACORTICAL MYELIN PROFILING WITH HISTOLOGY i. Myelin striation in cortex ii. Myelin profiles and their graded changes across the cortex iii. MRI-compatible atlas DORSAL (SUPRA) TEMPORAL CORTEX Areas: ■ tmag d ■ tparti st transverse temporal gyr 3 2nd transverse temporal gyr 3rd transverse temporal gyri cortical depth cortical depth 50 LATERAL TEMPORAL CORTEX 51 60 tparti 6b tmag d myelin staining **B** Intracortical Myelin profiling with MRI i. Sampling technique ii. Impact of resolution iii. Spatial variations in profiles - mean and skewness **DIFFERENCES IN PROFILE MEAN** mean MT cortical depth (%) relative cortical depth **DIFFERENCES IN PROFILE SKEWNESS** MT skewness resolution depth 0.8 cortical 1.0 1600 2200 1800 2000 C Myelin Profiling of White Matter Bundles with MRI MWF Corticospinal Cingulum Cingulate Callosum Forceps Minor IFOF Callosum Forceps Major 0.12 0.12 0.08 0.16 0.1 0. 0.12 0.2 0.08 0.10 0.07 0.08 0.1 0.06

Figure 2. Myelin profiling. (A) (i) Classic drawings of myelin-stained cortical sections highlight how cortical areas differ with regard to myelin striation, signifying the potential to characterize areas with respect to depthwise variations in myelin density. [Reproduced with permission from Hopf et al. (91).] (ii) Histology-derived myelin profiles from different areas (left). Comparing profiles to areal positions (right) reveals a large-scale gradient in the myeloarchitecture; in this case "confirming steplike decrease in myelin content with increasing distance from the auditory region" (92). Based on the area-naming convention of the original text: tmag d, subregio temporalis magna dorsalis; tmag m, subregio temporalis magna ventralis; tpartr, regio temporalis paratransversa; ttr, regio temporalis transversa (primary auditory area). Additional anatomical landmarks are as follows: ITG, inferior temporal gyrus; MTG, middle temporal gyrus; STG, superior temporal gyrus. [Reproduced with permission from Hopf et al. (92); colors added to aid comparison between line and surface plots.] (iii) Histologyderived atlas of myeloarchitecture (93) generated on a magnetic resonance imaging (MRI)-compatible cortical surface (94), including intracortical myelin profiles for many areas, based on Hopf's photomicrograph studies. (B) (i) MRI-derived myelin profiling involves generating pial and white matter surfaces, generating equivolumetric surfaces between these 2 boundaries, and then sampling imaging intensities along a vertex that crosses the surfaces. (ii) Imaging resolution influences the smoothness of the profiles, yet distinctive elements of the profile shape remain evident even at 1-mm resolution. [Reproduced with permission from Dinse et al. (39).] (iii) The mean and skewness of MRI-derived myelin profiles capture unique information about myeloarchitecture as shown by the different patterns of each feature across the cortical surface. Cortical maps were generated using quantitative magnetization transfer (MT) imaging of healthy adolescents (95). Profiles on the left show extreme cases of high and low features, exemplifying how the features capture different aspects of the profiles. (C) Average myelin water fraction (MWF) across segments of white matter bundles. x-axes represent the dominant spatial dimension of the specific tract. Error bars show standard deviation across subjects, highlighting the robustness of tract profiles. IFOF, inferior fronto-occipital fasciculus. [Reproduced with permission from Baumeister et al. (96).]

15

spatial axes

0.06

15

0.04

0.08

0.06

Together, these variations in mean and skewness illustrate the existence of distinct, overlapping organizational axes of myeloarchitecture in the human cortex (Figure 2Biii). This work emphasizes the importance of incorporating multiple features of intracortical profiles in in vivo imaging studies to better understand myeloarchitectural differences and map large-scale patterns of cortical differentiation.

Extending the profiling approach past the gray/white matter boundary, recent work has evaluated the density of superficial white matter (SWM) (107). Historically, SWM (also known as the U-fiber system) has been difficult to study in vivo. The typical approach for fiber tracking, namely diffusion-weighted imaging, must be used at ultra-high-resolution for SWM (108) to account for its thinness and preponderance of crossing fibers (109). Alternatively, elevated iron levels in SWM (61) may be leveraged to target it with R2\* (107). Immunohistochemical analysis of SWM has shown that its iron content colocalizes with oligodendrocytes, reinforcing the relevance of this approach to understanding myelin processes. Using an extended profiling approach, Kirilina et al. (107) identified higher density of SWM in frontotemporoparietal association areas than primary sensorimotor areas. This pattern notably differs from the regional distribution of intracortical myelin (Figure 2B), suggesting that together these profiling approaches can reveal the unique combinations of short- and long-range fibers in different cortical areas.

Myelin profiling can also shed light on ensheathment along white matter tracts. Electron microscopy studies of nonhuman animal brains have shown that myelin thickness can vary along axons (110,111). Relaxometry, diffusion-based, and g-ratio measures are known to vary along white matter tracts (81,112,113). Explicitly profiling MWF along white matter bundles, Baumeister et al. (96) recently identified characteristic patterns for each tract that could be replicated in all subjects (Figure 2C). This approach is more in its infancy than intracortical myelin profiling, and potential caveats, such as crossing fibers, need to be addressed (114). Even so, profiling approaches show promise for disambiguating subbundles within fasculi (115). All in all, the profiling approach benefits from eschewing the assumption of uniform myelin distribution along white matter tracts and holds promise for offering greater sensitivity and specificity to inter- and intraindividual differences.

## LINKING LOCAL MYELIN MARKERS TO CONNECTOME ORGANIZATION

Next, we ask how local myeloarchitecture, revealed by myelin profiling, contributes to the large-scale function of the human connectome. In particular, we highlight 2 avenues of recent work that have probed the interrelations between local myelin markers and structural connectome topology and network efficiency.

Contemporary perspectives emphasize that cortical gradients capture multifactorial changes in neurobiological features (58,116–119). The most prominent gradient, the sensory-fugal axis, runs from primary sensory areas toward limbic areas (103,116) and involves concomitant changes in myeloarchitecture, cytoarchitecture, connectivity, and function (118,120,121) (Figure 3Ai, ii). Many projections pass stepwise

along the sensory-fugal axis, producing a set of parallel processing hierarchies, emanating from each primary sensory area (125) (Figure 3Aiii). Sequential processing through these hierarchies, with graded changes in underlying myelo- and cytoarchitecture, is thought to allow integration of information from several sources and gradual abstraction of neural code (116,118). We recently found that individual-level sensory-fugal axes may be defined by applying nonlinear dimensionality reduction to myelin profiles [(103); Figure 3B]. Thus, this approach links local properties of intracortical myeloarchitecture to global axes of cortical differentiation and provides a new foundation to map the dominant streams of information processing in individual human brains.

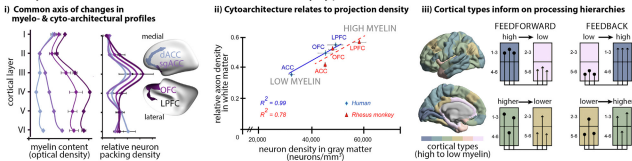
Turning toward white matter, the distribution of myelin influences the relationship between structural connectome topology and functional efficiency. The g-ratio, in particular, is a key contributor to conduction velocity (78). Thus, myelinsensitive imaging can be combined with diffusion imaging to approximate conduction velocity across the connectome (126). In doing so, recent work has shown that the rich-club has lower g-ratio than local edges, which may enable faster and more efficient propagation within a set of densely connected but widely distributed regions (127). Initial work in Parkinson's disease also suggests that mapping myelin across the connectome can help to identify aberrant tracts that are associated with scores on motor performance tasks, signifying the potential importance of local myelin measures on more distributed brain function (128).

## ADVANCING LIFESPAN RESEARCH WITH MYELIN-SENSITIVE IMAGING

The human lifespan is an ideal target for myelin research. Postmortem studies have long evidenced correlations between age and myelin, yet uncertainty remains regarding the principles of myeloarchitectural maturation. In vivo myelinsensitive imaging offers the opportunity to track spatiotemporal patterns of myelin across the entire lifespan in large samples, helping to show how local changes can shape trajectories of larger-scale brain organization and their interrelations with other neurobiological features.

A principal challenge for lifespan research is determining the maturational sequence of myelination. As early as the 1870s, Flechsig (129) sought to prove that certain laws dictate the developmental sequence of pathways in the brain and spinal cord. Through detailed examination of white matter tracts and intracortical myelin in postmortem tissue, he showed that developmental myelination is protracted and asynchronous (7). The onset and duration of myelination (the myelogenetic cycle) varies across fiber systems and regions, with cycles spanning from in utero to adulthood. Kinney et al. (130) set forth the following general rules that explain the temporal patterns of myelination: 1) proximal pathways myelinate earlier and have shorter duration of myelination than distal pathways; 2) sensory pathways myelinate before motor pathways; 3) projection pathways myelinate earlier and have shorter myelination intervals than associative pathways; 4) myelination progresses from the central sulcus toward the poles; 5) occipital, followed by frontal, and then temporal poles myelinate; and 6) posterior frontoparietaloccipital areas have faster myelination than

## A Myeloarchitecture as an index for connectivity types



## **B** From myelin profiles to the principal axis of myeloarchitectural differentiation

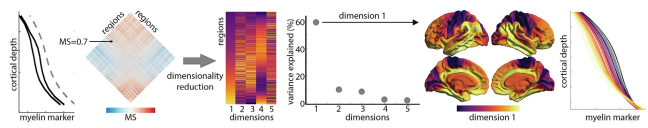


Figure 3. Relationship of local myelin to large-scale cortical organization. (A) (i) While myelin profiles (left) and cytoarchitectural profiles (right) differ in shape, they often capture a common spatial axis of changes across areas. For example, in the prefrontal cortex shown here, total myelin content and density of neurons in layer III increase along an axis that runs in a medial-lateral loop via the orbitofrontal cortex. dACC, dorsal anterior cingulate cortex; LPFC, lateral prefrontal cortex; OFC, orbitofrontal cortex; sgACC, subgenual anterior cingulate cortex. [Adapted with permission from Zikopoulos et al. (120). Line plot colors match the Brodmann areas delineated on the cortical surface (121).] (ii) Intracortical neuronal density is strongly correlated with the density of axons projecting from the cortical area, mirroring the spatial axis identified in Figure 3Ai and suggesting that intracortical myeloarchitecture is also associated with projection density [Reproduced with permission from Zikopoulos et al. (120). High and low myelin labels were added to aid interpretation.] (iii) Cortical types capture multifactorial changes in cyto- and myeloarchitecture that systematically vary along spatial axes (122). The relative type of 2 cortical areas indicates whether projections are in the feedforward (left) or feedback (right) direction as well as the laminar origin of projections. [Adapted with permission from Barbas and Rempel-Clower (123).] Thus, the relative myeloarchitecture of 2 cortical areas can index functional and structural characteristics of a neural network. (B) The principal axes of myeloarchitectural differentiation are resolved by first calculating myeloarchitectural similarity (MS) between regions based on the correlation of myelin profiles. Then, dimensionality reduction, typically diffusion map embedding (124), is applied to a matrix that contains MS between many cortical regions. The resultant dimensions reflect different spatial axes and are ranked according to the variance they explain in the MS matrix. In healthy adults, the first dimension runs from primary sensory and motor areas to limbic areas, closely approximating the sensory-fugal axis defined by postmortem histology (103). Coloring myelin profiles based on their position on dimension 1 illustrates how the axis reflects a decrease in myelin content as well as a shift from a concave to a convex curve. Notably, each dimension is sensitive to a different aspect of the myelin profile shape, related to myelin content at certain cortical depths. [Reproduced with permission from Paquola et al. (103).]

anterior frontotemporal regions. These large-scale rules provide a benchmark for imaging studies, which can in turn extend upon the postmortem research by providing nonbinary, quantitative assessment of myelin through investigation of healthy development and individual variability in large samples. For example, Kulikova *et al.* (131) demonstrated the potential of using multimodal parameters (R1, R2, and diffusion-based) to infer the maturational sequence of white matter bundles. Notably, the multivariate approach conformed to histological benchmarks with higher accuracy than did univariate approaches, supporting the utility of multimodal imaging for tracking myelin changes across the lifespan.

Recent studies have used MRI-derived myelin profiles, evaluated at multiple time points, to show how the patterns of myelin changes relate to brain organization at different life stages. Several theories have been proposed regarding the determinants of age-related myelin changes, such as the last-in/first-out hypothesis (132) and the spatial gradient hypothesis (130). Myelin-sensitive imaging is well positioned to test

these hypotheses, given the possibilities to examine myelin across the entire brain and to track individual developmental trajectories. In infants, Grotheer et al. (133) showed that myelin levels at birth and spatial position contributed to agerelated R1 changes in white matter bundles (Figure 4Ai). In certain tracts, speed of myelination is inversely correlated to the preexisting degree of myelin, whereas in other tracts, it is associated with spatial axes (Figure 4Aii). In adolescents, age-related changes in myelin also appear to reflect preexisting differences in intracortical myelin profiles (Figure 4Bi) (95). Specifically, areas that were less myeloarchitecturally distinct (relative to sensory and limbic areas) in early adolescence exhibit the strongest age-related changes throughout adolescence and young adulthood (Figure 4Bi). Notably, intracortical myelin profiles have linked local changes (i.e., myelination at a specific cortical depth) to largescale patterns of myeloarchitectural maturation (Figure 4Bii). Beyond mere correlative effects, myelin profiles may hold predictive power for later life. Imaging studies suggest that

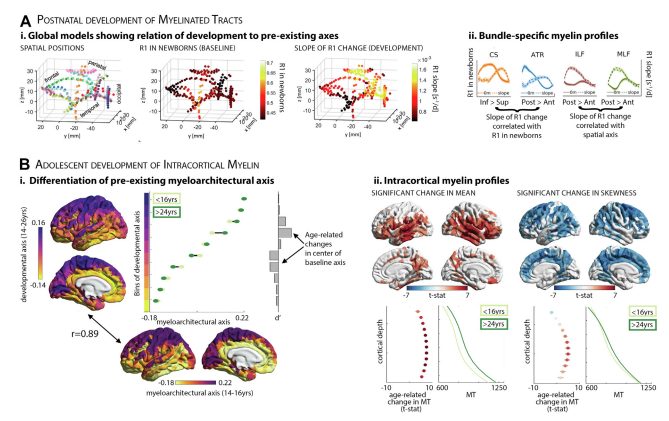


Figure 4. Myelin profiling approaches link local developmental changes to large-scale patterns of brain organization. (A) (i) Scatterplots represent sampling points along white matter bundles, colored by (left) bundle, (center) R1 in newborns and (right) development rate of R1 from 0 to 6 months. Across all points, development rate could be largely explained (67% of variance) by R1 in newborns and spatial position (anterior-posterior [ant-post] and inferior-superior [infsup] axes). [Reproduced with permission from Yeatman et al. (134).] (ii) Examining bundle-specific myelin, the developmental rate (slope, dotted line) is sometimes inversely correlated with R1 in newborns (0 m, full line) and at other times is correlated with a spatial axis (post-ant depicted as x-axis). [Reproduced with permission from Yeatman et al. (134).] ATR, anterior thalamic radiation; CS, cortico-spinal; ILF, inferior longitudinal fasciculus; MLF, middle longitudinal fasciculus. (B) (i) Regional variation in intracortical myelin profiles derived from magnetization transfer (MT) can be surmised by the myeloarchitectural axis. The more distant two regions are on the myeloarchitectural axis, the more dissimilar their intracortical myelin profiles are. In contrast, the more distant two regions are on the developmental axis, the more dissimilar their age-related changes in intracortical myelin profiles are. Notably, the baseline myeloarchitectural axis (derived from the earliest time points) and the developmental axis (calculated across the full age range) were strongly correlated (r = 0.89). Comparing myeloarchitectural axes at <16 and >24 years indicated that the most prominent age-related changes (Cohen's d effect size) were evident in the regions in the center of the developmental axis, suggesting differentiation of association cortex (orange) during this age range toward either the sensory (purple) or paralimbic (yellow) extremes. [Reproduced with permission from Paquola et al. (95).] (ii) Age-related changes in mean and skewness of myelin profiles shown on the cortical surfaces (threshold: q-value false discovery rate < 0.00625). Scatterplots show t statistic (mean  $\pm$  SD) for age-related changes in MT intensity at each sampled cortical depth within significant regions. Mean increases were balanced across surfaces, whereas decreases in skewness were driven by preferential MT increases at mid-to-deeper surfaces. Line plots exemplify how myelin profiles change from the lowest to oldest age groups in significant regions. [Reproduced with permission from Paquola et al. (95).]

earlier patterns of myeloarchitecture can predict myelin decline in older age. Specifically, the speed of myelin accumulation in adolescence can index the speed of myelin decline in older age (i.e., fastest in, fastest out hypothesis) (134). Fine-grained spatial variations in later life have not yet been explored with profiling approaches, which could help to discern bundle- and region-specific differences. Together, these studies demonstrate how myelin-sensitive imaging is helping in testing hypotheses of how development and degeneration progress across the brain. This research suggests that a basic set of organizational axes may govern bundle- and region-specific myelination across the lifespan.

In parallel, multimodal approaches are helping to elucidate the interrelation of age-dependent changes in myelin with other neurobiological features. Particular attention has been paid to disentangling the contributions of myelin and axonal properties from diffusion-based parameters (135). Furthermore, the balance between myelin and axonal measures can inform clinically relevant aspects of development and degeneration. For instance, throughout infancy, the g-ratios of white matter bundles decrease logarithmically toward adult levels (136), likely related to the increasing thickness of myelin sheaths (137) and the efficient conduction speed in brain networks (138). At the other end of the lifespan, the equilibrium between remyelination, myelin degradation, and axonal loss can be approximated by multiparameter decomposition of T2 and may help indicate healthy versus pathological aging (12,139).

# CONCLUDING REMARKS AND FUTURE PERSPECTIVES

The neuroimaging field provides strong evidence that myelin may be evaluated with in vivo MRI. Various MRI markers of myelin have been validated by comparing their values to well-established myelin markers acquired in the same tissue (i.e., histology, immunochemistry, or electron microscopy). Theoretically and empirically, T2-derived MWF, quantitative MT, and R1 provide good specificity to myelin. More caution is warranted in interpreting contrasts without such validation as specific myelin markers (e.g., T1w/T2w). Identifying the optimal multimodal combinations that can disambiguate myelin from artifactual and naturally occurring confounds, such as iron, is an important line of future research.

Moving forward, complementary "wide" and "deep" studies can help advance understanding of myeloarchitectural changes across the lifespan. On the one hand, large cohort studies that span a wide age range and demographic spectrum can leverage efficient whole-brain, quantitative, myelinsensitive sequences to confirm (or deny) whether laws of myelination (133) generalize across the population and to extend these laws to intracortical myelin and aging populations. On the other hand, ultra-high-field MRI studies can investigate attributes of myeloarchitecture that were previously only accessible through postmortem microscopy, such as laminar detail in the cortex (39) as well as myelin distribution along U-fibers (107) and deep white matter tracts (110). Ideally, such studies would focus on deeply phenotyping a few individuals with repeated scans, helping to reveal the dynamic intraindividual changes in myelin across short and long time frames.

Overall, a key advantage of progress in investigating myeloarchitecture in vivo is the ability to directly assess the relationship of myelin with function (10), cognition (140), behavior (141), and diseases (12,142). Previous MRI studies have shown that increases in myelin markers on specific tracts are associated with more mature activity patterns in certain brain regions, supporting the notion of concomitant maturation of myelin and function in the brain (143,144). Conversely, MRI markers of myelin degradation have been linked to cognitive decline in healthy older individuals and individuals with Alzheimer's disease (145). Further work is necessary to characterize the likely bidirectional relationship between myelin and brain function. "Wide" studies of myeloarchitecture across large cohorts will help establish the associations between regional myelin and cognitive skills, but "deep" approaches offer greater promise in disentangling their causal relationships. In particular, behavioral training and neurofeedback protocols may be able to reveal the dynamic intraindividual changes in myelin and brain activity that support cognitive development or decline (146).

Myelin-sensitive imaging has the potential to complement and advance upon current approaches for understanding psychiatric and neurological diseases, especially those with neurodevelopmental or neurodegenerative origins. Schizophrenia, for example, is associated with reduced oligodendrocytes and myelin-related gene expression [for review, see (147)]. However, the nature of dysmyelination in schizophrenia, especially its progression, remains a contentious issue.

Hypotheses variably focus on abnormal neurodevelopment (148) or accelerated neurodegeneration (149). The above-discussed advances in myelin imaging, analytics, and life-span research paved the way to accurately track and characterize aberrant age-related changes in myelin in individuals with schizophrenia, thereby disentangling competing theories, improving understanding of the etiology of the disease, and benefiting models of clinical course. Similar outcomes are now possible for a wide range of psychiatric and neurological diseases thanks to recent advances in myelin-sensitive imaging.

## **ACKNOWLEDGMENTS AND DISCLOSURES**

CP was funded by Helmholtz Association's Initiative and Networking Fund through the Helmholtz International BigBrain Analytics and Learning Laboratory under the Helmholtz International Lab grant agreement InterLabs-0015 and the Deutsche Forschungsgemeinschaft (German Research Foundation) (Grant No. 491111487).

The authors report no biomedical financial interests or potential conflicts of interest.

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Received May 18, 2022; revised Aug 12, 2022; accepted Aug 30, 2022.

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