White Matter and Cognition: Making the Connection

Christopher M. Filley¹*

R. Douglas Fields²

1. Behavioral Neurology Section, Departments of Neurology and Psychiatry, University of Colorado School of Medicine, Aurora, Colorado, and the Denver Veterans Affairs Medical Center

2. Eunice Kennedy Shriver National Institutes of Child Health and Human Development, NIH

Contact information: Bldg. 35, Room 2A211, Bethesda, MD, USA 20892
(301) 480-3209 fieldsd@mail.nih.gov

*Corresponding author:
Behavioral Neurology Section
12631 East 17th Avenue, B-185
Aurora, CO  80045
Phone: 303 724-8225, Fax: 303 724-2202
christopher.filley@ucdenver.edu

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Abstract | Whereas the cerebral cortex has long been regarded by neuroscientists as the major locus of cognitive function, the white matter of the brain is increasingly recognized as equally critical for cognition. White matter comprises half of the brain, has expanded more than gray matter in evolution, and forms an indispensable component of distributed neural networks that subserve neurobehavioral operations. White matter tracts mediate the essential connectivity by which human behavior is organized, working in concert with gray matter to enable the extraordinary repertoire of human cognitive capacities. In this review, we present evidence from behavioral neurology that white matter lesions regularly disturb cognition, consider the role of white matter in the physiology of distributed neural networks, develop the hypothesis that white matter dysfunction is relevant to neurodegenerative disorders, including Alzheimer’s Disease and the newly-described entity chronic traumatic encephalopathy, and discuss emerging concepts regarding the prevention and treatment of cognitive dysfunction associated with white matter disorders. Investigation of the role of white matter in cognition has yielded many valuable insights, and promises to expand understanding of normal brain structure and function, improve the treatment of many neurobehavioral disorders, and disclose new opportunities for research on many challenging problems facing medicine and society.
Introduction

The central importance of the cerebral cortex in the elaboration of human behavior enjoys such wide popularity that this perspective has become axiomatic. The billions of cortical neurons in the human brain, and the trillions of synapses linking them, are regularly regarded as comprising the singular repository of cognitive function, so much so that neurologists routinely invoke the term “higher cortical function” to describe the study of brain-behavior relationships. Similarly, neuroscientists retain such firm allegiance to the hegemony of the cerebral cortex that the term “corticocentric myopia” has been used to describe the relative paucity of neuroscientific investigation devoted to other brain areas that may contribute to cognitive function (Parvizi, 2009).

Yet roughly half the brain is occupied by white matter, and it is reasonable to ask in what way this collection of myelinated tracts might inform the study of cognition (Fields 2008). The vast extent of white matter is an obvious datum meriting consideration: in one adult, the millions of myelinated fibers coursing throughout the brain have a combined length that would encircle the earth more than three times (Walhovd et al. 2014). Evidence from clinical neurology is now indicating that white matter serves a critical role in the organization of the distributed neural networks that are now conceptualized as the structural basis of evolved human behaviors (Mesulam 1990, Catani et al 2012). The advent of magnetic resonance imaging (MRI) in the last three decades has enabled the study of white matter in vivo (Catani et al. 2012), illuminating how lesions of white matter tracts disturb cognition. Many cellular changes in white matter can influence MRI, including differences in axon diameter, packing, or tortuosity, myelin, astrocytes, oligodendrocytes, and vasculature. MRI has also shown how differences in normal white matter structure correlate with level of performance in a wide variety of cognitive functions, such as
facility with arithmetic (Matejko and Ansari, 2015) and musical performance (Ullén, 2009). In parallel, experimental advances have occurred that elucidate basic aspects of white matter structure and function (Fields, 2008; Walhovd et al., 2014). Together, these developments have introduced a new multidisciplinary approach to the study of normal and abnormal cognition by focusing on a long overlooked region of the brain. In this review, we consider recent clinical and laboratory evidence informing the understanding of white matter-cognition relationships in health and disease.

The contribution of white matter to cognition

The neural basis of human cognition has been largely established by the clinical-pathological study of diseases that disrupt cognitive function. Behavioral neurology relies heavily on the “lesion method,” (Damasio 1984) which considers cognitive dysfunction in relation to brain lesions, and for most of the history of medicine, post-mortem analysis of the brain was the only means available to examine the lesions produced by neurologic disorders such as stroke, traumatic brain injury (TBI), and degenerative dementia. Together with gray matter, white matter was included in these studies, and classic observations in the 19th century identified neurobehavioral syndromes such as conduction aphasia, pure alexia, and ideomotor apraxia in which focal vascular lesions of white matter were found to play a pivotal role. These and other syndromes were famously summarized by the behavioral neurologist Norman Geschwind in his seminal 1965 paper on disconnection syndromes (Geschwind, 1965). According to Geschwind, disconnection of cerebral regions by white matter damage merited consideration equal to that given to focal lesions of the cerebral cortex.
Modern neuroimaging dramatically advanced the study of brain-behavior relationships, first in the 1970s with computed tomography (CT), when for the first time neurologists could see structural brain pathology \textit{in vivo} (Bradley, 1986). MRI, however, proved far more useful than CT in the visualization of white matter and its lesions (Bradley, 1986), allowing for correlation of these abnormalities with cognitive dysfunction during life (Tanridag and Kirshner, 1987). In the early years of the MRI era, white matter lesions were often called “unidentified bright objects” because of uncertainty about what specific neuropathology could be responsible (Kertesz et al., 1988), but with time it became clear that these white matter hyperintensities reflected the appearance of water accumulating in the presence of myelin disruption from many etiologies (Anderson et al., 2014). One of the most common MRI abnormalities was termed leukoaraiosis (LA), referring to scattered hyperintense lesions in the cerebral white matter of older people on T2 and fluid-attenuated inversion recovery (FLAIR) images (Hachinski et al., 1987). The origin of LA was determined to be most likely related to ischemia (Pantoni and Garcia, 1997), leading investigators to consider the possibility that LA is an early phase of the well-known vascular white matter dementia known as Binswanger’s Disease (BD; Babikian and Ropper, 1987). Indeed, evolving consensus opinion increasingly supports the view that LA is a clinically mild precursor state to the progressive dementia of BD (Rosenberg et al. 2016; Schmidt et al., 2016). Figure 1 shows axial MR brain images of older individuals to demonstrate the scattered ischemic white matter lesions of LA (Figure 1A) and confluent ischemic white matter hyperintensity consistent with BD (Figure 1B). The study of LA (a new observation) in the context of BD (an old disease) is but one example of how white matter-cognition relationships rapidly gained momentum with the advent of MRI (Catani et al., 2012).
The focal neurobehavioral syndromes related to white matter lesions (Geschwind, 1965) offered an important clue that disruption of distributed neural networks could be a general phenomenon underlying cognitive dysfunction, but the emergence of Alzheimer’s Disease (AD) as a major medical challenge (Katzman, 1976) focused much investigation on the cerebral cortex, where the signature microscopic lesions of this disease are found (Querfurth and LaFerla, 2010; Katzman 1976). As work on AD soon dominated the study of dementia, cognitive decline came to be widely conceptualized as a result of cortical disease (Geldmacher and Whitehouse, 1996), and pathology in subcortical regions received relatively little attention.

The cerebral cortex, however, occupies only the outermost 1-3 mm of the brain, and many subcortical structures play a crucial role in cognition. The idea of subcortical dementia, which gained prominence in the 1970s (Albert et al., 1974; McHugh and Folstein, 1975), proposed that diseases of the subcortical gray matter such as Parkinson’s Disease and Huntington’s Disease – with major pathology in the substantia nigra and caudate, respectively – feature cognitive loss because of disturbance in the timing and activation of cognitive functions. Whereas criticism was initially raised regarding the characterization of clinical and neuropathological features of subcortical dementia (Whitehouse, 1986; Brown and Marsden, 1988), the concept persists as clinically helpful in characterizing the phenomenology of dementing illness in relation to specific regions of pathologic involvement (Bonelli and Cummings, 2008; Ropper et al., 2014).

White matter is the other constituent of the subcortex. To illustrate the widespread distribution of myelinated systems, Figure 2 shows a diffusion tensor imaging (DTI) depiction of normal white matter tracts. Comprising roughly half the total brain volume, white matter tracts course throughout the brain to unite cortical and subcortical gray matter regions into functional
neuronal ensembles subserving cognition and emotion (Mesulam, 1990; Schmahmann and Pandya, 2006; Catani et al., 2012). White matter thus forms an essential component of the human connectome, or the structural description of the human brain (Sporns et al., 2005). An important observation is that white matter has actually expanded more in evolution than gray matter, reflecting the requirement for more myelination as cortical expansion leads to greater distance between neuronal cell bodies (Zhang and Sejnowski, 2000). The selective expansion of white matter, in turn, appears to contribute to the singular intelligence of *homo sapiens*; while other intelligent animals such as whales and elephants actually have larger brains, with nearly as many cortical neurons, humans have the most extensive cerebral myelination (Roth and Dicke, 2005). Thus human cognition depends as much on brain connectivity as on the activity of cortical neurons.

These considerations are consistent with prior observations on focal disconnection syndromes (Geschwind, 1965), and further predict that diffuse lesions of the white matter can reliably produce cognitive deficits that may even reach the severity of dementia. Evidence for this notion is steadily mounting, based primarily on the correlation of cognitive loss with MRI white matter lesions (Filley, 2012). A wide variety of white matter disorders can compromise cognition (Filley, 2012), and because these disorders typically produce widespread or diffuse neuropathology, the parallel involvement of multiple distributed neural networks is presumed responsible. MRI has been transformative in this context, and a newer and particularly appealing neuroimaging technology being investigated is DTI, which enables the detailed depiction of white matter microstructure by assessing the diffusion of water along myelinated tracts (Zhang et al., 2012). The most common DTI parameters, fractional anisotropy (FA) and mean diffusivity (MD), are used to quantitate the degree of abnormal (isotropic, or random) water diffusion within
white matter, but DTI is not yet capable of generating specific information as to whether axonal, myelin, or other pathology is involved (Zhang et al., 2012). This technique, however, holds much promise for the non-invasive identification and characterization of white matter pathology.

A convincing example of the potential for white matter neuropathology to produce dementia is toluene leukoencephalopathy (TL), a dementia syndrome resulting from intense and prolonged exposure to inhaled toluene as a result of solvent vapor abuse (Hormes et al., 1986; Rosenberg et al., 1988; Filley et al., 1990, Filley et al., 2004; Filley, 2013). This common but underappreciated form of substance abuse can produce severe neurotoxicity because of daily exposure to inhaled toluene that may be pursued for many years (Hormes et al., 1986). The high lipophilicity of toluene accounts for its predilection to damage myelin, and toxic exposure produces widespread cerebral myelin loss with concomitant dementia (Rosenberg et al. 1988). The degree of cerebral white matter injury correlates with the severity of cognitive impairment (Filley et al., 1990), supporting the notion that cerebral white matter injury in TL can be sufficient to cause dementia. The white matter regions specifically involved in the pathogenesis of dementia appear to be the large tracts of the cerebral hemispheres, as intracortical myelin has been noted to be unaffected (Rosenberg et al., 1988, Filley et al., 2004) or relatively spared (Kornfeld et al., 1994). Indeed, a systematic review of 30 empirical studies found that toluene can produce dementia by preferentially affecting white matter relative to gray matter, and that periventricular and subcortical white matter is most vulnerable (Yucel et al., 2008).

Based on the observations of TL, other neurologic disorders with prominent white matter involvement have been observed to manifest a similar clinical picture (Schmahmann et al., 2008). In addition to leukotoxic injury as exemplified by TL, vascular, traumatic, demyelative, inflammatory, infectious, metabolic, hydrocephalic, neoplastic, and genetic disorders can all
damage brain white matter and produce similar cognitive effects (Filley, 2012). Table 1 displays these ten categories and an example of a specific disorder within each. Many of these disorders feature co-existent gray matter involvement, and in many cases an admixture of gray and white matter pathology likely accounts for cognitive dysfunction and dementia. Yet all the entities in Table 1 share a substantial burden of white matter pathology, and evidence is accumulating that supports a selective role of white matter injury in cognitive dysfunction and dementia (Schmahmann et al., 2008). To highlight the potential for white matter disorders to produce this often devastating outcome, the term white matter dementia (WMD) was introduced in 1988 to help organize thinking about white matter dysfunction in relation to cognitive decline (Filley et al., 1988).

A unique profile of cognitive deficits has been seen to characterize WMD, and reflects the physiological role of white matter in normal cognition (Filley et al., 1988; Filley, 2012). The most important feature of WMD is cognitive slowing, an expected result of slowed impulse transmission in the brain resulting from damage to myelin, and in some cases axons as well. Other deficits include executive dysfunction, memory retrieval dysfunction, sustained attention deficit, visuospatial impairment, and various psychiatric disorders; conversely, language, extrapyramidal function, and procedural memory are relatively preserved (Filley, 2012). This profile differs from that of both the cortical dementia of AD – in which amnesia, aphasia, apraxia, and agnosia are typical (Geldmacher and Whitehouse, 1996) – and subcortical dementia, in which procedural memory is impaired (Saint-Cyr, Taylor, and Lang, 1998). Table 2 displays core cognitive differences between cortical, white matter, and subcortical dementia. These differences are most evident in the early stages of dementia, as with the progression of neuropathology all cognitive functions are eventually lost and the dementias become
indistinguishable. Yet the clinical distinctions in Table 2 are not merely academic; the profile of deficits and strengths revealed at an early stage can be very helpful in diagnosis and treatment. From the available evidence, therefore, it appears that diffuse damage to the white matter can produce a distinctive dementia syndrome, recognition of which can be clinically valuable (Filley, 2012; Filley et al., 1989, Lafosse et al., 2007; Schmahmann et al., 2008).

It should be noted, however, that dementia is generally encountered only with a heavy burden of white matter lesions. In contrast, many individuals harbor lesser degrees of white matter neuropathology, and have a less severe clinical syndrome. For example, the relatively mild white matter lesion burden of LA shown in Figure 1A produces cognitive slowing and executive dysfunction but not dementia, whereas the more advanced white matter disease shown in Figure 1B often culminates in the progressive dementia of BD (Filley, 2012). The early cognitive syndrome seen with mild white matter disease on conventional MRI may even result from involvement of what has been called the normal-appearing white matter (NAWM) (Filley, 2012), which can be found with the use of advanced neuroimaging technologies such as MR spectroscopy (MRS) and DTI. This more subtle syndrome, which has been termed mild cognitive dysfunction (MCD; Kozora and Filley, 2011), features a profile of cognitive slowing, inattention, and executive dysfunction, qualitatively similar to the pattern seen in WMD. MCD has clinical implications as a plausible precursor of WMD, which may appear later as the burden of white matter pathology worsens (Filley, 2012). The constructs of WMD and MCD need more study, but thus far these ideas are consistent with what is known of white matter structure and function in health and disease. That is, the structural alteration of multiple white matter tracts by various neuropathologies can disrupt the normal function of these tracts in a manner that begins with the subtle cognitive disturbance of MCD, and then, if unchecked, advances to the disabling
neurobehavioral syndrome of WMD. In clinical neurology, MCD could provide a much-needed
criterion for the early identification of cognitive loss from white matter involvement than can
potentially be treated before irreversible disability supervenes.

The role of white matter in distributed neural networks

In general, white matter can be seen as providing for the transfer of information within
distributed neural networks, while gray matter subserves information processing (Filley, 2012).
Accordingly, primary white matter damage results most prominently in cognitive slowing,
whereas primary gray matter disease leads to more specific cognitive deficits, most apparent in
the cortical dementia AD, which features amnesia, aphasia, apraxia, and amnesia related to
regional neuronal and synaptic loss (Geldmacher and Whitehouse, 1996; Table 2). However, as
discussed below, suboptimal conduction velocity can also impair information processing; a good
example of this phenomenon is optic nerve demyelination in acute optic neuritis that impairs visual
acuity in multiple sclerosis (MS; Thurtell et al., 2009). Neurons are the fundamental units of all
networks, and distinctions between the operations of white and gray matter are not absolute.
Disease may begin in the white matter or the gray, or progress from one tissue to the other, and
complex interactions between the two are typical. As myelocentrism can be as limiting as
corticocentrism, a balanced view of the representation of cognitive function is crucial. In this
section, we consider physiologic aspects of white matter underlying both information transfer
and processing as a basis for appreciating the effects of myelin and axonal dysfunction.

An understanding of white matter function begins with the details of its microscopic
anatomy. Figure 3 displays key aspects of white matter structure. Precision of spike time arrival
is critical for information processing and synaptic plasticity. Temporal summation of
postsynaptic membrane potentials from multiple synaptic inputs onto a dendrite requires
millisecond precision to depolarize the postsynaptic neuron to threshold for initiating a spike. In addition, the strength of synapses can be increased or decreased by the degree of coincidence of synaptic input relative to postsynaptic action potential firing (spike timing-dependent synaptic plasticity; Dann and Poo 2006). That is, synapses that fire coincidentally with or a few milliseconds before action potential firing in the postsynaptic neuron becomes strengthened, but synaptic strength is weakened in synapses firing a few milliseconds after the action potential. Despite the high temporal precision required for synaptic function in gray matter, conduction delays over long range axons in the mammalian brain (and especially large-brained primates), are on the order of tens of milliseconds or longer. Thus cognitive function can be impaired by suboptimal conduction times through white matter tracts connecting synaptic relay points, and complete impulse conduction failure following demyelination is not required for functional impairment.

Coordination of neural activity in large assemblies of neurons through phase and amplitude coupling is critical for cognition and consciousness (Singer, 2009; Buzsáki, 2006) and the conduction time between synaptic relay points is an important parameter affecting the coherence and frequency of brain wave activity. Brain waves, which represent the oscillations of neural activity at appropriate frequencies, and the coupling of oscillatory activity across long-distance cortical networks, are critical for cognitive function, gating of sensory information, and the binding of cognitive operations. White matter disease can disrupt normal electroencephalography (EEG) coherence patterns in association with impaired cognitive function (Nunez et al., 2015). Recent mathematical modeling predicts that conduction delays will have a profound effect on coupling oscillatory activity in the brain and that even small changes in myelination can produce substantial effects on coupling oscillatory activity in the brain.
Indeed, this modeling predicts that biological mechanisms for adaptively modulating conduction delays must exist to prevent hypo- and hyper- synchrony of coupled oscillators in the brain, and that changes in myelin would be one of the most effective means for such plasticity (Pajevic et al., 2014). A number of neuropsychiatric disorders are associated with disruption in brain rhythms that may derive from changes in myelin (Mathalon and Sohl, 2015; Uhlhaas and Singer 2015), including autism (Welsh and Ahn, 2005), schizophrenia, obsessive-compulsive disorder, depression (Schulman et al., 2011; Ferrarelli et al., 2012, Xu et al., 2013), and dyslexia (Ucles et al., 2009). These examples highlight the possibility that idiopathic disorders with cognitive dimensions may be understood by considering myelin disturbances that result in network dysfunction. The connectopathy so produced may include both psychiatric and neurologic dysfunction, as illustrated by the dysmyelinative disease metachromatic leukodystrophy (MLD), in which a typical progression from early psychosis to later dementia has been identified, presumably as a result of advancing white matter disease (Black, et al. 2003).

In addition to myelin defects degrading impulse conduction, the myelin sheath provides metabolic support for axons (Fünschilling et al., 2013; Lee et al., 2012). Demyelination and the resulting impaired trophic support for axons by oligodendrocytes can cause axonal and neuronal degeneration (Lee et al., 2012), further contributing to cognitive deficits in disorders such as MS (Koenig et al., 2014) and TBI (Armstrong et al., 2015). Axonal dysfunction in white matter disorders is well recognized to confer a worse prognosis than that implied by myelin damage alone (Trapp et al., 1998; Medana and Esiri, 2003). Thus whereas myelin damage slows network activity, the superimposed loss of axons may preclude any neural conduction and render the network inoperative.
White matter and neurodegenerative disease

An intriguing implication of the relationship between white matter and cognition is the etiopathogenesis of neurodegenerative dementia. The diseases within this group, the most common of which is AD, remain idiopathic, incurable, and a major threat to medicine and society. While the neurologic disorders capable of producing WMD do not include those in the category of neurodegeneration, white matter dysfunction has in fact been correlated with cognitive dysfunction in one such disease, fragile X tremor ataxia syndrome (FXTAS) (Filley et al., 2015), and no a priori reason exists as to why idiopathic neurodegenerative diseases should necessarily implicate selective gray matter involvement. The idea that white matter dysfunction may prove important in understanding the origin of neurodegenerative disease warrants consideration.

The problem of AD remains particularly disturbing because, despite much effort over the past three decades, no disease-modifying treatment is available. Reflecting the corticocentric bias that exerts a powerful influence on the study of dementia (Parvizi, 2009), AD is widely regarded as a cortical disease in which neuritic plaques and neurofibrillary tangles are entirely responsible for the dementia syndrome (Geldmacher and Whitehouse, 1996; Querfurth and LaFerla, 2010). Amyloid is thought to be the primary cause of cortical injury, followed in pathogenesis by tau neurotoxicity (Querfurth and LaFerla, 2010). The amyloid hypothesis currently dominates the AD field, postulating the centrality of amyloid β42 (Aβ) and its oligomers in producing cortical damage, and this notion has stimulated a great deal of work on identifying agents that can treat the disease by ridding the brain of this presumably toxic protein (Hardy and Selkoe, 2002). Although the pathogenetic role of Aβ in autosomal dominant early-onset AD offers support for
the amyloid hypothesis (Querfurth and LaFerla, 2010), it is sobering to consider that many
normal elders harbor sufficient amyloid to qualify for the diagnosis of AD, that the normal
function of amyloid and its relative amyloid precursor protein (APP) are unknown, and that all
efforts to treat AD by targeting brain amyloid have thus far failed (Castellani and Perry, 2014).
Nevertheless, amyloid and tau may plausibly disrupt cortical structure, and ongoing studies of
anti-Aß therapies may yet find that amyloid reduction can be effective at some point in the
course of AD. However, the idea has recently been offered that cortical pathology occurs as a
downstream event in AD pathogenesis, and that early white matter injury triggers an adaptive
response that produces the deposition of cortical amyloid and tau (Castellani and Perry, 2014;
Bartzokis, 2011). In this context, recent MRI evidence has supported the notion that white matter
hyperintensities are prominent early features of both late-onset (Brickman, 2013) and dominantly
inherited AD (Lee et al., 2016). Moreover, DTI has shown that AD is associated with disrupted
connectivity between various cortical and subcortical regions (Bozzali et al., 2010). Thus the
“myelin model of AD” proposes that upstream events within the cerebral white matter, such as
vascular disease and head trauma, initiate the deposition of amyloid and then tau as later end-
products (Bartzokis 2011). Indeed, evidence exists for both vascular (Marnane et al., 2016) and
traumatic white matter damage (Scott et al., 2016) promoting amyloid deposition in the cerebral
cortex. If the myelin model proves to be correct, preventive intervention long before the onset of
cognitive dysfunction may be profoundly important.

Cholinergic deficit is a prominent feature of AD, and acetylcholinesterase inhibitors
(AchEIs) such as donepezil and rivastigmine are used therapeutically in patients with AD. While
AchEIs are known to act at the cholinergic synapse, the benefits of AchEI treatment in AD
patients have also been linked to direct effects on oligodendrocytes and myelination (Bartzokis,
2007). A paradoxical increase in white matter connectivity in the internal capsule of AD patients is associated with AChEI treatment (Bozzali et al., 2012), and donepezil-induced oligodendrocyte differentiation is inhibited by the nicotinic acetylcholine receptor antagonist mecamylamine, but not by the muscarinic acetylcholine receptor antagonist scopolamine (Imamura et al., 2015). Interestingly, butyrylcholinesterase (BChE), an enzyme closely related to AChE, is found mainly in white matter and glia, and a BChE genotype influences white matter loss in AD (Lane and He, 2013). BChE becomes expressed in association with cerebral cortical Aβ plaques (Darvesh 2016), linking acetylcholine to both gray and white matter pathology in AD.

Another neurodegenerative disease attracting much attention is chronic traumatic encephalopathy (CTE), in which repetitive mild TBI in athletes and soldiers is proposed to lead to a dementia syndrome consequent to cortical tauopathy that appears many years later (McKee et al., 2013). The highly visible publicity surrounding CTE should not obscure significant controversy over its very existence, as critics have pointed out that tauopathy has not been definitively shown to cause dementia, and that the clinical features of CTE can be explained by alternative diagnoses such as depression and frontotemporal dementia (Iverson et al. 2015). However, because mild TBI regularly involves diffuse axonal injury (DAI) spread broadly throughout the white matter (Alexander, 1995), it is plausible that the insult of repetitive concussion – or even repeated subconcussive blows (Bailes et al., 2013) – may activate a neuropathological cascade that manifests later as hyperphosphorylated tau accumulation in the cortex that produces dementia. In health, tau is a normal protein primarily associated with microtubules within brain axons, and observations of boxers with dementia pugilistica, a neurodegenerative disease very similar to CTE, have documented tau hyperphosphorylation as a
result of repetitive mild TBI (Bartzokis, 2011). Although tauopathy is emphasized as the
centerpiece of CTE neuropathology, white matter injury is present in all stages of the disease
(McKee et al., 2013), implying that DAI may be the trigger for progressive tau deposition. The
progressive accumulation of abnormal tau that is thought to occur, possibly via a mechanism
similar to prion propagation (Prusiner, 2013) might therefore actually be a downstream
phenomenon in CTE, similar to the proposed role of amyloid in the myelin model of AD
(Bartzokis, 2011). While CTE remains highly controversial, the possible long-term effects of
traumatic white matter injury have far-reaching implications for both civilians and the military.

Prevention and treatment of white matter lesions

Many opportunities can be considered for preventing and treating white matter lesions
that can produce cognitive dysfunction and dementia. Medical intervention and public policy
strategies merit attention because many white matter disorders are related to vascular risk factors,
head injury, and intoxication with a variety of leukotoxins (Filley, 2012). Standard medical care
clearly plays a role, offering much benefit in terms of the control of hypertension, diabetes,
hyperlipidemia, metabolic syndrome, cigarette smoking, and obesity, and by preventing TBI and
substance abuse. Attention to vascular risk factors is crucial, as these problems are strongly
associated with white matter ischemia because of the selective atherosclerotic vulnerability of
small penetrating arterioles irrigating cerebral white matter. Hypertension, for example, is a
powerful risk factor for white matter ischemic lesions and stroke (Debette and Markus, 2010),
and a large prospective study found that antihypertensive therapy was not only effective for
stroke prevention but also reduced the incidence of dementia (Forette et al., 2002), raising the
possibility of an effect on ischemic white matter disease. Prospective studies evaluating the
efficacy of primary cerebrovascular care and physical exercise are underway to address the hypothesis that targeting white matter ischemia can prevent dementia (Prins and Scheltens, 2015). Disease-specific medical treatment can also contribute, as in the demyelinating disease MS, in which a variety of medications have been shown to reduce white matter disease burden (Wingerchuk and Carter, 2014). A novel line of inquiry now attracting considerable attention is the microbiome, and recent findings in mice that normal gut microbiota promote normal prefrontal cortical myelination (Hoban et al., 2016) may have clinical relevance. Public policy intervention is also important, improving the health of white matter by policies intended to enhance access to medical care, enact seat belt and helmet laws, support education to encourage lifelong cognitive engagement, and promote physical fitness.

Another approach to prevention and treatment involves the exploitation of intrinsic white matter repair mechanisms. Plasticity in the brain, meaning the capacity of neurons to be modified as a result of experience, has traditionally been considered a function of the gray matter, but recent findings have found that white matter also exhibits this phenomenon (Fields, 2008; Zatorre et al., 2012). Clinical and basic science investigations have been foundational in elaborating this novel idea (Fields, 2015).

White matter plasticity has been shown in both normal individuals, such as piano players whose pyramidal tract integrity correlated with number of hours practiced (Bengtsson et al., 2005), and in neurologic patients, such as those with Broca’s aphasia in whom right arcuate fasciculus volume increased as Melodic Intonation Therapy improved language performance (Schlaug et al., 2009). Human and experimental animal studies using DTI have found that structural changes indicative of plasticity may occur in the fornix, as well as the hippocampus, within two hours after engagement in a learning task (Hofstetter et al., 2013). Physical activity...
also appears to be salutary, as aerobic exercise has been found to improve white matter integrity in community-dwelling older adults (Voss et al., 2013). Better performance in inhibitory control is associated with white matter structure (increased fractional anisotropy) in prefrontal and frontostriatal tracts in healthy subjects (Forstmann et al., 2008), and in children with attention deficit hyperactivity disorder (Liu et al., 2012). In animal studies, social isolation in mice can impair myelination of the forebrain (Liu et al., 2012; Makinodan et al., 2012), demonstrating experience-dependent plasticity of myelination. Motor skill learning on a treadmill requires oligodendrogenesis in adult mice (McKenzie et al., 2014). The exploitation of white matter plasticity may thus find clinical utility as a method of both maintaining normal tract function, and repairing damaged tracts to restore normal cognition. New findings in white matter neurobiology may open new avenues for preventing or improving cognitive dysfunction, and for clinicians and policy makers, the prospect of making evidence-based recommendations such as playing a musical instrument, learning a new language, and staying fit could indeed be appealing.

In the last ten years, considerable advances have been made in identifying molecular mechanisms regulating development of oligodendrocytes and myelination, clarifying how cognitive impairments associated with white matter disorders may be prevented and treated. Brain myelination is an extremely complex process, regulated by multiple intrinsic and extrinsic signals during development and throughout life. Myelination encompasses a broad scope of biological processes, including appropriate cell specification from progenitors, precise control of cell proliferation, selective survival and apoptosis of cells in the oligodendrocyte lineage, cell migration, differentiation, identification of the proper cellular targets to be myelinated (only appropriate axons are myelinated, not other cellular structures, and not all axons are myelinated),
the formation of specialized axon-glial membrane complexes providing intercellular
communication to induce myelin formation, intricate cytokinesis and synthesis of enormous
amounts of lipid and membrane protein, and compaction of multilaminar axonal ensheathments
into a compact spiral sheath of membrane that is unlike any other membrane specialization or
cell-cell contact known.

In addition, because myelination requires the formation of nodes of Ranvier to enable
saltatory conduction, an intricate paranodal apparatus is needed to segregate appropriate ion
channels in the axonal membrane that are essential for impulse conduction. This segregation is
achieved by precise subcellular localization of unique cell adhesion molecules - a trimolecular
complex of Caspr, Contactin, and NF155, and others (Zonta et al., 2008) - that are organized into
septate junctions that can be visualized only by high resolution transmission electron
microscopy. Thus many cellular signals are involved in the process of myelin formation, myelin
maintenance, and remyelination, and this complexity increases vulnerability to various genetic
and environmental factors that can cause abnormal myelination and dysfunction. As these
signaling molecules become identified, however, new potential approaches to therapeutic
intervention become available.

A number of membrane receptors are implicated in oligodendrocyte differentiation and
myelination (Table 3). Many of these molecules are of potential relevance to neurologic
disorders affecting cognition. In hypothyroidism, for example, one of the most sought after and
commonly treated causes of dementia, cognitive deficits may in fact be related to cerebral white
matter involvement because thyroid hormones are important for myelination (Barres et al.,
1994). Many implications for the treatment of myelin damage are also apparent. Neuregulin is
centrally involved in myelination, and has been proposed as a possible myelin repair strategy in
older people with ischemic or traumatic white matter injury (Bartzokis, 2011). In younger people with MS, agents targeting LINGO-1 activity are being tested in clinical trials as a means of enhancing remyelination based on the role of LINGO-1 in inhibiting myelination by oligodendrocytes (Mi et al., 2005).

Cell adhesion molecules and extracellular matrix molecules also influence oligodendrocyte differentiation and myelination, notably integrins (Decker and ffrench-Constant, 2004), Tenascin-C (Frost et al., 1996; Garcion et al., 2001) and others. Interestingly, experiments in which dorsal root ganglion neurons are stimulated to fire action potentials in different frequencies show that mRNA abundance for different cell adhesion molecules on axons (NCAM, n-Cadherin, and L1-CAM) is differentially regulated by specific patterns of action potential firing (Itoh et al. 1995; Itoh et al., 1997). Moreover, specific frequencies of firing that lower expression of L1-CAM, an adhesion molecule that is necessary for myelin wrapping, reduces the amount of myelin that forms (Stevens et al., 1998). Mechanisms regulating myelination according to the pattern of action potential activity in neural circuits are especially important with respect to involvement of white matter plasticity in cognitive function.

Indeed, neurotransmitter signaling has been found to be particularly important in modifying development of myelinating glia in an activity-dependent manner. Among these are the neurotransmitters ATP (Stevens and Fields, 2000), adenosine (Ishibashi et al., 2006; Stevens et al., 2002) GABA (Zonouzi et al., 2015), glutamate (Wake et al., 2011, 2015; Lundgaard et al., 2013), and acetylcholine (Abiraman et al., 2015; Deshmukh 2013; Mei et al., 2014).

Neurotransmitters can be released along axons firing action potentials by both vesicular and non-vesicular release mechanisms (Fields and Ni, 2010; Fields, 2011). GABA (Zonouzi et al., 2015) and glutamate (Mangin et al., 2012) have been shown to influence oligodendrocyte
proliferation and differentiation. Inhibitors of muscarinic acetylcholine receptors are effective in improving myelination in animal models of MS (Abiraman et al., 2015; Deshmukh et al., 2013; Mei et al., 2014) and in Phase 2 MS clinical trials (Green, 2016). In addition to these developmental effects, activation of NMDA and mGluR glutamate receptors on oligodendrocyte cell processes stimulates the formation of an axo-glial signaling complex that generates intracellular calcium transients in the oligodendrocyte cell process adjacent to the axon in response to action potential firing. This in turn initiates myelination selectively on electrically active axons by stimulating the local synthesis of myelin basic protein from mRNA transported from the cell body into individual processes of oligodendrocytes (Wake et al., 2011). A consequence of this activity-dependent signaling would be that axons that are electrically active would become preferentially myelinated. This has been confirmed by recent research showing that when given a choice, oligodendrocytes will form functional contacts preferentially with electrically active axons. Oligodendrocytes then form myelin selectively on these electrically active axons while avoiding interaction with axons in the same vicinity in which glutamate release has been blocked by treatment with botulinum toxin (Wake et al., 2015). Studies in zebrafish show that although initiation of myelination is not affected, stabilization of the nascent myelin sheath is promoted by vesicle release from axons in zebrafish (Mensch et al., 2015; Hines et al., 2015). Activity-dependent specification of myelination would have significant effects on network function by the approximately 50 times increase in conduction velocity of myelinated axons compared with unmyelinated axons.

To summarize, activity-dependent plasticity of both synapses in gray matter and myelin in white matter is increasingly appreciated as being of fundamental importance for central nervous system function and cognition. Several medical implications of activity-dependent
myelination are apparent. First, the block of axonal conduction in demyelinated axons may impair remyelination, particularly in view of the many aspects of myelination that are stimulated by impulse activity, and axonal loss is one reason spontaneous remyelination is limited in patients with MS. Second, functional recovery after axotomy requires not only axon regeneration but also remyelination, suggesting that treatments to promote remyelination could promote recovery after axon injury. Finally, functional activity or electrical stimulation may be helpful therapeutically in promoting remyelination after demyelination and other white matter injury, as shown by the increased numbers of oligodendrocytes in rodents after spinal cord injury (Becker et al., 2010) and in cell culture (Gary et al., 2012) when electrical stimulation is delivered.

**Summary and Future Directions**

White matter provides the structural connectivity between gray matter regions throughout the brain, and the importance of myelinated systems in cognitive function derives support from both behavioral neurology and basic neuroscience. Complementing the information processing of the cerebral cortex, white matter tracts provide for the information transfer within the brain, enabling the rapid and efficient integrative capacity of neural systems necessary for the highly evolved cognitive operations of *homo sapiens*. Distributed neural networks are organized to mediate critical aspects of cognition such as attention, memory, language, visuospatial skills, and executive function, all of which depend critically on the structural connectivity provided by myelinated systems. Disrupted white matter has potentially profound effects, including dementia, and from this perspective a host of clinical and basic science implications become readily apparent.
Advances in this field can be facilitated by a focus on many unresolved questions. First, because white and gray matter lesions are often commingled in disease states, more specific study of the effects of white matter dysfunction in isolation would be useful, particularly as it evolves over time from subtle NAWM changes to grossly visible macrostructural lesions on conventional MRI. Second, research on early vascular and traumatic injuries of white matter, initially mild or even inapparent but much more significant in the years to follow, may lead to crucial insights into the etiopathogenesis and treatment of presently irreversible degenerative dementias. Finally, the prevention and treatment of white matter disease merits much more study, as there appear to be many opportunities to focus on this area using medical and public policy intervention, and by exploiting the emerging field of white matter plasticity.
GRANTS

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DISCLOSURES

No conflicts of interest, financial or otherwise, are declared by the authors.


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Fulmer CG, VonDran WM, Stillman AA, Huang Y, Hampstead BL, Dreyfus CF.


### Table 1: Disorders with cognitive impairment and prominent white matter pathology

<table>
<thead>
<tr>
<th>Category</th>
<th>Disorder</th>
</tr>
</thead>
<tbody>
<tr>
<td>Toxic</td>
<td>Toluene leukoencephalopathy</td>
</tr>
<tr>
<td>Vascular</td>
<td>Binswanger’s Disease</td>
</tr>
<tr>
<td>Traumatic</td>
<td>Traumatic brain injury</td>
</tr>
<tr>
<td>Demyelinating</td>
<td>Multiple sclerosis</td>
</tr>
<tr>
<td>Inflammatory</td>
<td>Systemic lupus erythematosus</td>
</tr>
<tr>
<td>Infectious</td>
<td>Human immunodeficiency virus infection</td>
</tr>
<tr>
<td>Metabolic</td>
<td>Vitamin B₁₂ deficiency</td>
</tr>
<tr>
<td>Hydrocephalic</td>
<td>Normal pressure hydrocephalus</td>
</tr>
<tr>
<td>Neoplastic</td>
<td>Gliomatosis cerebri</td>
</tr>
<tr>
<td>Genetic</td>
<td>Metachromatic leukodystrophy</td>
</tr>
</tbody>
</table>

### Table 2: Cognitive features of cortical, white matter, and subcortical dementia

<table>
<thead>
<tr>
<th>Cortical</th>
<th>White Matter</th>
<th>Subcortical</th>
</tr>
</thead>
<tbody>
<tr>
<td>Amnesia</td>
<td>Cognitive slowing</td>
<td>Executive dysfunction</td>
</tr>
<tr>
<td>Aphasia</td>
<td>Executive dysfunction</td>
<td>Cognitive slowing</td>
</tr>
<tr>
<td>Apraxia</td>
<td>Memory retrieval deficit</td>
<td>Memory retrieval deficit</td>
</tr>
<tr>
<td>Agnosia</td>
<td>Normal procedural learning</td>
<td>Impaired procedural learning</td>
</tr>
</tbody>
</table>
Table 3 | Molecules important for oligodendrocyte differentiation and myelination

<table>
<thead>
<tr>
<th>Cellular Function</th>
<th>Signaling Molecule</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cell adhesion</td>
<td>Contactin</td>
<td>Hu et al. 2003</td>
</tr>
<tr>
<td>Cell adhesion</td>
<td>Integrins</td>
<td>Decker and ffrench-Constant 2004</td>
</tr>
<tr>
<td>Cell adhesion</td>
<td>L1-CAM</td>
<td>Stevens et al. 1998</td>
</tr>
<tr>
<td>Cell adhesion (extracellular matrix)</td>
<td>Tenascin-C</td>
<td>Frost et al. 1996; Garcion et al. 2001</td>
</tr>
<tr>
<td>Growth factor</td>
<td>Insulin-like growth factor (IGF-1)</td>
<td>Carson et al. 1993;</td>
</tr>
<tr>
<td><strong>Growth factor</strong></td>
<td><strong>PDGF-alpha</strong></td>
<td>Zeger et al. 2007</td>
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<tr>
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<td>----------------</td>
<td>------------------</td>
</tr>
<tr>
<td><strong>Growth factor</strong></td>
<td><strong>Epidermal growth factor receptor (EGFR)</strong></td>
<td>Aguirre et al. 2007</td>
</tr>
<tr>
<td><strong>Growth factor</strong></td>
<td><strong>FGF</strong></td>
<td>Fumagalli et al. 2001</td>
</tr>
<tr>
<td><strong>Growth factor</strong></td>
<td><strong>BDNF</strong></td>
<td>Fulmer et al., 2014; Miyamoto et al., 2015</td>
</tr>
<tr>
<td><strong>Receptor tyrosine kinase</strong></td>
<td><strong>Neuregulin 1-ErbB</strong></td>
<td>Brinkmann et al. 2008</td>
</tr>
<tr>
<td><strong>Signal Transduction</strong></td>
<td><strong>Neurogenic locus notch homolog protein (Notch 1), membrane receptor for Jagged and Delta</strong></td>
<td>Wang et al. 1998; Genoud et al. 2002</td>
</tr>
<tr>
<td><strong>Signal transduction</strong></td>
<td><strong>Wnt (Frizzled family receptor, beta-catenin translocation to nucleus)</strong></td>
<td>Fancy et al. 2009</td>
</tr>
<tr>
<td><strong>G protein-coupled receptor</strong></td>
<td><strong>Gpr 17</strong></td>
<td>Chen et al. 2009</td>
</tr>
<tr>
<td><strong>Intercellular signaling</strong></td>
<td><strong>Leucine-rich repeat neuronal protein 1 (LINGO-1)</strong></td>
<td>Aguirre et al. 2007</td>
</tr>
<tr>
<td><strong>Thyroid hormone</strong></td>
<td><strong>Triiodothyronine/thyroid hormone 3 (T3)</strong></td>
<td>Barres et al. 1994</td>
</tr>
<tr>
<td><strong>Neurotransmitter</strong></td>
<td><strong>ATP/Adenosine</strong></td>
<td>Ishibashi et al. 2006; Stevens et al. 2002</td>
</tr>
<tr>
<td><strong>Neurotransmitter</strong></td>
<td><strong>GABA</strong></td>
<td>Zonouzi et al. 2015</td>
</tr>
<tr>
<td><strong>Neurotransmitter</strong></td>
<td><strong>Glutamate</strong></td>
<td>Wake et al. 2011, 2015, Lundgaard et al. 2013</td>
</tr>
<tr>
<td><strong>Neurotransmitter</strong></td>
<td><strong>Acetylcholine</strong></td>
<td>Abiraman et al., 2015; Deshmukh et al., 2013; Mei et al., 2014</td>
</tr>
</tbody>
</table>

**Figure 1** | Axial fluid-attenuated inversion recovery (FLAIR) MRI scans showing A) the scattered white matter lesions of leukoaraiosis, and B) confluent white matter hyperintensity consistent with Binswanger’s Disease.

**Figure 2** | DTI scan of a normal adult brain showing three white matter tracts. Color coding permits the demonstration of tracts oriented within the right-left, anterior-posterior, and superior-inferior planes. Thus red indicates the corpus callosum, green represents the arcuate fasciculus, and blue depicts the corticospinal tract. (Courtesy of Mark S. Brown, PhD)

**Figure 3** | White matter structure. (A) Approximately 40 μm by 60 μm by 60 μm volume of white matter (mouse optic nerve) reconstructed from serial block face electron microscopy, showing the composition of axons, myelin, astrocytes, oligodendrocytes, and vasculature. Changes in any of these components or in the tortuosity of the fibers will influence diffusional
MRI, where the voxel volume is 100 μm 100 μm 100 μm in high-resolution MRI of rodents, and typically 2 mm by 2 mm by 2 mm in human brain imaging. (B) Optic nerve axons in cross section showing multilaminar wrapping of compact myelin. (C) Three-dimensional reconstruction of a node of Ranvier in mouse optic nerve from serial block face electron microscopy. Myelin (purple), paranodal loops (salmon), perinodal astrocyte (green and blue). Inset shows a longitudinal slice through the node of Ranvier revealing the nodal gap (gray) between the paranodal loops of myelin containing high density voltage-sensitive sodium channels. Axon (white). (D) Illustration of myelinated axons, showing the multilaminar myelin sheath and electrogenic node of Ranvier. Scale bars = 10 μm in A and 1 μm in B-D.
Figure 1
Figure 2