White Matter Hyperintensity and Vascular Disease from Biological Basis to Clinical Significance

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Abstract

"White matter disease" identifies a series of different conditions and pathological mechanisms: autoimmune, infectious, toxic-metabolic and vascular. Each of these leads to a global impairment of the neural myelination process through the secondary destruction of previously myelinated structures. To date, the imaging spectrum represents an irreplaceable tool to detect these lesions, describe their distribution patterns and stage them over time. This study aims presents a pictorial review of white matter disease, from pathology to imaging spectrum, reporting current main staging systems with greater emphasis to relationship between cerebrovascular disease and white matter hyper-intensity appearance, and the newest advances in this field.

Keywords: CT = Computed Tomography; DWMH = Deep White Matter Hyperintensities; LA = Leukoaraiosis; MRI = Magnetic Resonance Imaging; PVWMH = Periventricular White Matter Hyperintensities; SVD = Small Vessels Disease; WMH = White Matter Hyperintensities

Introduction

“White Matter Disease”, formerly described as “leukoaraiosis” by Hatchinsky on Computed Tomography (CT) [1], can be depicted into two main types according to their causes: a) primary, derived from an unknown etiology and b) secondary, derived from a great variety of known etiologies [2]. White Matter Hyperintensity (WMH) is a purely descriptive term currently used on Magnetic Resonance Imaging (MRI), and it represents a very common finding in older patients affected by a wide range of diseases included autoimmune, infective, toxic-metabolic and cerebrovascular. WMH can be classified into Periventricular (PV-WMH) and Deep Subcortical (DS-WMH). PV-WMH is characterized by gliosis, loosening of the white matter fibers and myelin, loss around tortuous venules in perivascular spaces, whereas the main features of DS-WMH are demyelination, gliosis and increased tissue loss as the lesions become more severe. Although cerebral blood flow and the cerebrovascular endothelial surface area are both so large as to consent storage of fluid and proteins within interstitial spaces, these findings suggest that Blood Brain Barrier (BBB) impairment plays a key role on genesis of WM damage [3].

Several papers demonstrated that specific clusters of human perivascular tissue proteins in the WM would be involved in the process of arteriolar wall thickening. One of these clusters includes fibrinogen, immunoglobulins and thrombomodulin, which occur in subcortical grey and white matter [4]; another cluster includes albumin, PCR (not defined), homocysteine, interleukin-6 and the intracellular adhesion molecule-1, diffusively present in cerebral interstitial tissue in patients with increased risk of WHM development and lacunar stroke [5]. Cerebrovascular diseases are related to BBB damage and to WMH through a series of concatenated events. From a pathological point of view, acute subcortical infarct leads, in 90% of cases, to cavitation or narrowing of surrounding brain tissues and secondary gliotic reaction that could create a protective “white matter hyperintensities cap” [6]. The main vascular pathologies associated with WMH are shown in table below (Table 1) [7].

| Disease | Etiology | Damaged Area |
|---------|----------|--------------|
| Small Vessel Disease (SVD) | Hereditary/Sporadic | Basal ganglia; fronto-temporal and periventricular WM |
| Amyloid-related angiopathy | Hereditary | Cortical-subcortical temporal lobe |
| Primary Angitis of CNS | Idiopathic | Non-specific distribution |
| CADASIL | Hereditary | Bilateral subcortical white matter of temporal lobe; bilateral external capsule |
| CARASIL | Hereditary | Basal ganglia and thalamus; diffuse leukoencephalopathy; middle cerebellar peduncle across the pons (arch sign) |
| MELAS | Hereditary | Basal ganglia; middle cerebral artery; lacunes (arch sign) |
| Susac Syndrome | Idiopathic | Corpus Callosum |

Table 1: Main cerebrovascular disease causing WMH.
muscle loss from the media with a general thickening and stiffness of the arterial wall and genesis of microaneurysm and micro-atheroma associated with calcification [8]. SVD due to Cerebral Amyloid Angiopathy is characterized by a progressive storage of the beta-amyloid protein into small and medium size cells of the vessel muscle wall mainly located on the leptomeningeal space, cortex and also in capillaries [9]. When vascular injuries occur in vascular “end-zones”, WMH distribution pattern will be focal beginning or diffuse confluent, while this pattern will appear centripetal in the basal nuclei, centrum semiovale, corona radiata and brainstem (Figure 1 and 2) [10].

Further, the recent small subcortical infarcts (involving perforating arterioles) may evolves into rounded or ovoidal lacunes of 3-15 mm and represent the interrelate pathogenic substrate for WMH, which can be identified at the edge of ischemic territory [11]. CADA-SIL represents the most common hereditary cause of cerebral vascular pathology related to subcortical infarcts and leukoencephalopathy. WMH involves firstly, the temporal lobe; and then progress to the posterior temporal, frontal and parietal regions likely to the basal nuclei and thalamus [12]. PACNS is a very rare vasculitis syndrome, characterized by leptomeningeal and cortical vessels inflammation [13].

MRI shows different and nonspecific pattern: a) “hemorrhagic form” related to intra-parenchymal infarction; b) “pseudo-tumoral form” related to mass lesion with central necrosis, perilesional edema, infiltrating and surrounding cerebral structures. Hence angiography can demonstrate multifocal narrowing of large cerebral vessels [14].

Susac Syndrome is a rare and multifactorial disease that involves small vessels and leads to vascular occlusion with thrombus formation involving the thalamus, basal nuclei and subcortical white matter [15]. MELAS is hereditary syndrome due to mutations on the mitochondrial genome. MRI shows some signal alteration areas as hyperintense through T2 and FLAIR sequences frequently located on parieto-occipital cortex and less frequently in the temporo-parietal cortex. Characteristically, these lesions don’t respect the vascular distribution unlike ischemic lesions [16].

### Imaging Hallmarks

#### CT imaging

Hatchinsky in the 1980 was the first scientist who described on CT, some findings of irregular low attenuation signals in the periventricular and deep white matter [1]. The different location of damage probably depends on different pathological pathways which involve different levels of brain vascularization. In particular, SVD determines Deep subcortical White Matter Hyperintensity (DWMH) due to the higher grade of hypoxic sensitivity along the so called “border zones,” with subsequent appearance of ischemic lacunas that are visible as hypo-intense areas on CT (Figure 3 and 4). On the contrary, especially on chronic stage, the involvement of large vessels causes a variable and sporadic distribution pattern of WM damage [17]. Periventricular White Matter Hyperintensity (PWMH) relates to expansion of pre-existing periventricular damage due essentially to benign processes such as venous collag eosis involving an elderly cohort of patients [18]. Otherwise, brain CT scan showed a high capability to localize white matter changes as one or more hypo-intensity areas, based on European Task Force on age related white matter changes, allowing staging of them into a four points scale [19].

0 = no lesion
1 = focal lesion > 5mm
2 = early confluent lesion
3 = diffuse involvement of brain

![Figure 1: 77-yrs female with arteriosclerosis-related SVD sagittal view on FLAIR-T2 sequences showing diffuse sovra-tentorial white matter hyperintensity.](image1)

![Figure 2: Axial view on T2-FLAIR of watershed zones infarcts showing multiple white matter hyperintensity areas along ACA-MCA (anteriorly) and MCA-PCA (posteriorly) border zones.](image2)

![Figure 3: Multiple hypointensity(orange arrow) areas associated with enlarged cortical sulci (blue arrow) and ventricular spaces depicting hypoxic-ischemic encephalopathy.](image3)
BBB permeability measurement with two modalities: Perfusion MR can dynamically evaluate WM lesions through processes to orientation of WM tracts [20]. Therefore, DTI can detect (AD), can respectively quantify parallel or perpendicular diffusivity addiction with its two components Radial (RD) and Axial Diffusivity (FA=1) or not (FA=0) of unidirectional water diffusion. MD, in one dimensional value scaled from 0 to 1 which describes the presence of Fractional Anisotropy (FA) and Mean Diffusivity (MD). FA is a tify integrity of neuronal pathways of white matter with evaluation of WM tracts [20]. Therefore, DTI can detect any microstructural damage of WM due to the demyelination process predating axonal loss [21].

Perfusion MR can dynamically evaluate WM lesions through BBB permeability measurement with two modalities:

- Arterial Spin Labeling: it works applying the inversion of longitudinal magnetization of blood flow of afferent artery without the use of contrast media. This inversion pulse labels specific intracranial vessels sampling in the head [22]
- Paramagnetic Contrast Administration = it works by reducing relaxation time on vessels and surrounding tissues that show an increase of signal on T1-weighted and a loss of signal on T2-T2* sequences.

Recent advances in MRI allow the more careful description of the location but also the nature (ischemic or not) of white matter injuries. Hydrogene Magnetic Resonance Spectroscopy (H-MRS) shows a reduced level of N-Acetyl Aspartate (NAA) and Cr along ischemic WMH area, which is intended as axonal and neuronal injury markers, while they do not show any variation in normal elderly patient’s cohort [23]. Another interesting role of MRI analyses is the evaluation of BBB integrity. Dynamic Contrast-Enhanced MRI (DCEMRI) can evaluate and stage compromised permeability of BBB alongside the ischemic core and “penumbra region” in deep white matter. DCEMRI identify increased permeability signal surrounding WMH’s area and decreased permeability signal within WMH’s area [24]. More recently, the 7 Tesla coils MRI yields a better spatial resolution to well identify early WM injuries, detecting more detailed sub cortical infarcts, micro bleeds and therefore distinguishing ischemic lacunas from perivascular spaces [25].

| Imaging Technique                  | Common Features                                                                 |
|------------------------------------|---------------------------------------------------------------------------------|
| Computed tomography               | Multiple and/or single hypo-dense lacunes involving periventricular and deep white matter |
| Magnetic resonance                | Focal or diffuse areas of T2-FLAIR hyperintensity and T1 hypo-intensity; post-ischemic lacunes appear as a focal area of FLAIR hypo-intensity |
| Diffusion tensor imaging          | Visualize orientation to quantify integrity of neuronal pathways of white matter with evaluation of Fractional Anisotropy (FA) and Mean Diffusivity (MD) |
| Perfusion imaging                 | Evaluate and stage compromised permeability of BBB alongside the ischemic core and “penumbra region” in deep white matter |
| Spectroscopy                      | Shows a reduced level of N-acetyl aspartate (NAA) and Cr along ischemic WMH area |

Quantitative tractography of fibers crossing WMH with small and large volumes of WMH.

Low-volume WMH (a-c) and high-volume WMH (d-f). a, d T2 FLAIR MRI with WMH. b, e Three-dimensional region of interest of WMH. c, f Tracts crossing WMH. With kindly permission of Reginold, William & Luedeke, Angela & Tam, Angela & Iorralba, Justine & Fernandez-Ruiz, Juan &Reginold, Jennifer & Islam, Omar & Garcia, Angeles. (2015).

MRI grading scales

Some Authors tried to define the location and size of white matter damage trying to relate this one to progression of underlying pathology, as Fazekas did in 1987, when he proposed the “Fazekas Scale” to simplify the quantification of amount of WMH lesions [26]. This scale is still valid, and it remains a very useful and direct system to classify the severity of white matter diseases according to size, location and confluence of lesions (Table 2). Later, in order to classify white matter changes, Wahlund (Table 3) and Longstreth (Table 4) assessed other two minor methods that included basal ganglia and both DWMH and PWMH involvement. [27,28]. The width of the sub-arachnoid spaces and lateral ventricles are classified with the use of analog visual rating scale.

Figure 4: CT axial scans show large and confluent hypodensity «leukoaraiosis» areas related to moderate-severe SVD involving corona radiata and centrum semiovale with gliotic deterioration (white arrow).
Furthermore, Prins et al., [29], classified white matter lesions on the basis of their diffusion upon brain areas. This required defining periventricular a WMH area as directly extending along the ventricle’s borders and subcortical a WMH area as extending perpendicularly more or less 10 mm from the ventricle’s borders. This scale provides different scores depending on where the lesions are being located and their progression. According to this method, white matter lesions frame could include three main periventricular locations (frontal, parietal, temporal and occipital caps and lateral bands), and periventricular score fluctuates from -3 to +3; similarly, subcortical lesions includes four patterns of distribution (frontal, parietal, temporal and occipital) resulting in a subcortical score from -4 to +4.

**Table 2 (Fazekas Scale): Four-point scale to describe deep white matter and periventricular white matter damage.**

| Deep White Matter (DWMH) | P=0 | P=1 | P=2 | P=3 |
|--------------------------|-----|-----|-----|-----|
| Absent “caps” or pencil thin lining | smoothhalo | irregular periventricular signal extending into deep white matter |

| Periventricular White Matter (PVWMH) | P=0 | P=1 | P=2 | P=3 |
|-------------------------------------|-----|-----|-----|-----|
| Absent punctate foci | beginning confluenve | large confluentareas |

**Table 3 (Wahlund Scale): 4-point scale where focal lesions are not defined; they are interpreted as diffuse-focal lesion. Brain areas used for rating are: frontal, parieto-occipital, temporal, infratentorial and basal ganglia (globus pallidum, nucleus caudatus, putamen, thalamus, internal-external capsule and insula).**

| White Matter | Number of Lesions |
|-------------|-------------------|
| 0 | Absent |
| 1 | Focal |
| 2 | Beginning confluent |
| 3 | Diffuse of entire regions with or without subcortical U-fibers |

| Basal Ganglia | Number of Lesions |
|--------------|-------------------|
| 0 | Absent |
| 1 | 1 focal lesion (>5 mm) |
| 2 | More than 1 focal lesions |
| 3 | Confluent lesions |

**Table 4 (Longstrehth Scale): Eight-point scale for WM lesions graded at the level of the body of lateral ventricle.**

| P=0 | Absent |
| P=1 | Discontinuous periventricular rim with minimal dots of subcortical disease |
| P=2 | Thin continuous periventricular rim with a few patches of subcortical disease |
| P=3 | Thicker continuous periventricular rim with scattered patches of subcortical disease |
| P=4 | More irregular periventricular rim with mild subcortical disease and minimal confluent periventricular white matter hyperintensities |
| P=5 | Mild periventricular confluenve surrounding the frontal and occipital horns |
| P=6 | Moderate periventricular confluence surrounding the frontal and occipital horns |
| P=7 | Periventricular confluence with moderate involvement of centrum semiovale |
| P=8 | Periventricular confluence involving most of the centrum semiovale |

**Figure a Figure b Figure c Figure d**

Schematic “mild” WMH distribution along periventricular and deep subcortical brain areas using different staging scales. Red shadows highlight scattered WMH areas.

**Corresponding WMH Score:**

| Figure a | Figure b | Figure c | Figure d |
|----------|----------|----------|----------|
| Fazekas scale = 1 | Fazekas scale = 2 | Fazekas scale= 1 | Fazekas scale = 2 |
| Wahlund scale = 1.5 | Wahlund scale = 1.5 | Wahlund scale= 1.5 | Wahlund scale = 2.5 |
| Longstrehth scale = 2 | Longstrehth scale = 1 | Longstrehth scale= 3 | Longstrehth scale = 4 |

**Schematic severe WMH distribution along periventricular and subcortical brain areas using different staging scales. Red shadows highlight confluent WMH areas.**

**Corresponding WMH Score:**

| Figure a | Figure b | Figure c | Figure d |
|----------|----------|----------|----------|
| Fazekas scale = 3 | Fazekas scale = 3 | Fazekas scale= 3 | Fazekas scale = 3 |
| Wahlund scale = 2.5 | Wahlund scale = 2.5 | Wahlund scale= 2 | Wahlund scale = 2.5 |
| Longstrehth scale = 4 | Longstrehth scale = 4 | Longstrehth scale= 2 | Longstrehth scale = 4 |
Discussion and Conclusion

WMH lesions tend to appear heterogeneous, reflecting their degree of “whiteness”. This feature explains how cellular breakdown leads to tissue structural and vascular changes not only inside of “core” of WMH area, but also on surrounding white matter’s “penumbra” which evolve further away from the visible area of WMH. In particular, the extension of WMH edges relates to the improvement of functional outcomes in patients with a previous acute ischemic stroke or cerebral hemorrhage [30]. Saba et al., investigated a probable intrinsic relationship between LA and the distribution of cerebral micro bleeds, intended as an independent biomarker for small vessel stroke or cerebral hemorrhage [30]. Saba et al., investigated a probable intrinsic relationship between LA and the distribution of cerebral micro bleeds, intended as an independent biomarker for small vessel stroke or cerebral hemorrhage [30]. Saba et al., investigated a probable intrinsic relationship between LA and the distribution of cerebral micro bleeds, intended as an independent biomarker for small vessel stroke or cerebral hemorrhage [30].

Recently, many Authors suggested a determinant role of carotid disease [38,39] as specific predictor factor for increasing WMH areas. In particular, Lucatelli et al., carried on a study considering a leading role of carotid occlusion degree, evaluating in particular carotid Intima-Media Thickness (IMT) as the driving parameters for cerebrovascular risk such as atherosclerosis progression or regression. They enrolled a cohort of 61 patients to MRI examination and subsequently to carotid US that support the hypothesis according to which IMTV could represent a double increased hyper-intensity, suggesting a sensibility to post-recanalization blood flow necrosis in those areas in which there is an increased hyper-intensity signal. In particular, their results showed a double increased risk of intracerebral hemorrhage in acute ischemic stroke patients with detected LA. Therefore, the importance of detecting “leukoaraiosis” signs regards not only clinical outcome after stroke, but also the recovery after administration of rt-PA. Bahrami et al., proposed another interesting research perspective about WMH pathogenesis; they speculated about the specific relationship between WMH distribution (periventricular or deep) and decreased regional cortical blood flow. They found two important markers: the first was that deep WMH is associated with decreased regional cortical blood flow, and the second that blood flow in WM shows a greater reduction in the periventricular than in deep WMH areas. The strongest and most consistent risk factors that showed the clearest associations with WMH were age, hypertension, hypercholesterolemia, diabetes and smoke [33].

Despite the fact that many Authors considered hyperglycemia as having a primary role on genesis of SVD, there are many controversial associations between diabetes and WMH areas genesis, thus there is not sure concordances to consider diabetes as independent risk factor. This data discordance is likely due to a different and non-uniform sampling of case-control population [34]. DWMH or PVWMH can associate to different pathological correlates such as vascular dementia [35]. This particular form of dementia showed high relationships with subcortical impairment of the frontal white matter network and other single infarction areas with punctate or confluent “area foci”: thalamus, angular gyrus, deep frontal areas and the left hemisphere [36]. Even the small vessels circulation may be different between deep and periventricular areas, with higher periventricular white matter damage due to large vessels disease [37]. SVD leads to lacunar infarction that shows a tight connection to WHM. In particular, WMH proximal to normal course of perforating vessels, are highly distributed at the edge of incident lacunas and enlarge from periventricular to subcortical region [11]. These findings suggest that vascular risk factors with reduction of general cerebral blood flow are associated with a pervasive SVD involving both periventricular and deep white matter.

According to these results, Lin et al., conducted a recent meta-analysis by enrolling more than 6912 participants with a past ischemic stroke treated with an rt-PA thrombolytic therapy in acute stage, trying to identify eventual relationships between the rate of extension of WMH burdens and clinical outcome after administration of the therapy [32]. They confirmed that deep micro bleeds detected by T2*-weighted gradient echo sequences relate to the periventricular hyper-intensity, suggesting a sensibility to post-recanalization blood flow necrosis in those areas in which there is an increased hyper-intensity signal. In particular, their results showed a double increased risk of intracerebral hemorrhage in acute ischemic stroke patients with detected LA. Therefore, the importance of detecting “leukoaraiosis” signs regards not only clinical outcome after stroke, but also the recovery after administration of rt-PA. Bahrami et al., proposed another interesting research perspective about WMH pathogenesis; they speculated about the specific relationship between WMH distribution (periventricular or deep) and decreased regional cortical blood flow. They found two important markers: the first was that deep WMH is associated with decreased regional cortical blood flow, and the second that blood flow in WM shows a greater reduction in the periventricular than in deep WMH areas. The strongest and most consistent risk factors that showed the clearest associations with WMH were age, hypertension, hypercholesterolemia, diabetes and smoke [33].

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compartmentalization of vascular blood-flow along Circle of Willis (COW) demonstrating lower risk of stroke and transient ischemic attacks among people who have effective collateral circles, while higher risk in those who have poor collateral circles [43].

On this basis, Saba et al., investigated about any existing relationships between COW variants and LA volume analyzing COW variations and WMH’s area by MRI on FLAIR and MRA on TOF sequences. The results confirmed not only a different location of LA distribution volume with predilection for MCA and PCA territories, but also a significant association between more COW existing variants and more WMH’s extension areas detected [44]. Bilateral occlusion of carotid artery, detected with CTA, would be determinant to emphasize a perpetual state of inflammation of blood brain barrier, through the activation in loco of MMPs that contribute to brain injury. In particular, the majority of the emboli, originated from principal atheroma, addressed through "perforating arteries" to most hypoxic sensitive territory such as hippocampus, white matter and some areas of cortex with a subsequent ischemia and genesis of WMH’s areas [45].

Further, Saba et al., analyzed, in a multicentric study, 98 patients who underwent CTA examination of brain and supra aortic vessels in parallel both brain white matter damage either grade of stenosis and nature of plaque. This study confirmed that CTA scan detect the major risk patient’s cohort developing a future LA and lead to consider white matter change as intermediate of stroke’s brain damage evolution in elderly vasculopathic patients [46,47]. All of these studies consider the presence or not of carotid artery stiffness [48], carotid artery wall thickness, composition and behavior of carotid plaque detected by echo-color Doppler, as independent risk factors for WMH genesis and distribution alongside the brain. In particular, patients with an unstable carotid plaque developed an extended distribution of WMH’s burdens especially on periventricular area, with associated cognitive decline [49].

Thanks to these findings, we can assume a new interpretative role of combined imaging (ECD, CT and MRI) as predicting the outcome in patient’s cohort undergoing to Carotid Endarterectomy (CEA). The global vascular impairment observed leads to negative influence and also to general outcome and successful recanalization of carotid vessel after CEA and Carotid Stenting Procedure (CAS) [50]. The most reasonable hypothesis is that basal hypo-perfusion leads to a reduced wash out of emboli not only on the treated artery side but also on the contralateral, leading frequently to ischemic brain injuries [51]. This feature explains how much the extension of WM lesions, detected by MRI with DWI and FLAIR sequences, could influence a clinical outcome and predict a higher probability of a new ischemic event after CEA among patients with asymptomatic ischemic brain lesions [52]. In the last decade, many studies focused on clinical impact of WMH; in particular, the European Leukoaraisis and Disability (LADIS) study contributed to this porpouse substantially. This multicenter study provided harmonization between different diagnostic instruments (clinical, neuropsychological and radiological) to predict progression on global disability in subjects with a WMH load using Fazekas Scale and Scheltens Visual Rating Scale for global and temporal atrophy. After three years follow-up they found a significant correlation between moderate-severe WMH load and progression of clinical disability, such as speed and motor control, with annual transition of 29.5% [38]. They also suggested strong relationship between WMH load and cognitive decline as well as attention, visual-constructural and naming praxis depending from number and distribution of lacunar infarcts and depression [38]. Since depression can be described as a result of altered patterns of regulation between ventral structures (affective sphere) and dorsal structures (cognitive sphere), especially involving the prefrontal cortex, any vascular insults of this neural pathway can cause a disbalance and lack of modulation of limbic structures activity. Based on this hypothesis Aizenstein et al., conducted a functional imaging study using f-MRI among 33 elderly depressed patients and 27 nondepressed comparison subjects which demonstrated, among cohort of patients with middle and late-life onset of depression, a strong relationship between global cerebral WMH burden and higher degree of BOLD response of ventral and anterior structures (amygdala, rostral cingulate, prefrontal cortex and anterior insula) using faces and shapes affective reactivity task [39]. In particular, they observed in late-life onset of depression rostral cingulated hyperactivitation associated with a greater WMH load as the strongest predictors of treatment despite middle-life onset of depression [39].

Corpus Callosum is another emerging important topic of study which can relate with cognitive and motor disabilities and many studies suggest that its involvement may be a stronger predictor of worst evolution on memory performance, executive dysfunction and speed [53]. As described above, WMH represents a very useful radiological “primer” to identify many pathological processes. Correct imaging assessment and scare present a crucial pathway to improve clinical approach predicting outcome and staging risk factors among their nature. Further perspectives on this field need many longitudinal studies are needed to evaluated effective long-term risk factors and predictors of clinical outcome.

Conflicts of Interest

The authors declare no conflict of interest.

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