White Matter Lesions in Adults – a Differential Diagnostic Approach
Läsionen der weißen Substanz im Erwachsenenalter –
ein differenzialdiagnostischer Ansatz

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ABSTRACT

Objective Cerebral white matter lesions on MRI in adults are a common finding. On the one hand, they may correspond to a clinically incidental feature, be physiologically or age-associated, or on the other hand they may be the overture to a severe neurological disease. With regard to pathophysiological aspects, practical hints for the differential diagnostic interpretation of lesions in daily clinical practice are presented.

Material and Methods With special regard to the vascular architecture and supply of the cerebral white matter, physiological structures are schematically represented and pathophysiological processes are highlighted by comparative image analysis of equally angulated MR sequences.

Results The most frequent vascular, inflammatory, metabolic, and neoplastic disease entities are presented on the basis of characteristic imaging findings and corresponding clinical-neurological constellations. The details of signal intensities and localization essential for differential diagnosis are highlighted.

Conclusion By means of comparative image analysis and the recognition of characteristic lesion patterns, taking into account anatomical principles and pathophysiological processes, the differential diagnostic classification of cerebral white matter lesions and associated diseases can be significantly facilitated. The additional consideration of clinical and laboratory findings is essential.

Key Points:
▪ Cerebral white matter lesions can be a harmless secondary finding or overture to a severe neurological disease.
▪ The comparative image analysis of different sequences with identical angulation is crucial.
▪ With special regard to the vascular anatomy, different lesion patterns can be identified.
▪ The consideration of neurological and laboratory chemical constellations is essential for the differential diagnosis.

Citation Format

ZUSAMMENFASSUNG

Ziel Zerebrale Marklagerläsionen im MRT beim Erwachsenen sind eine häufige Befundkonstellation. Sie können einerseits einem klinisch inapparenten Zufallsbefund entsprechen, physiologisch oder altersassoziert sein, oder andererseits die Ouverture einer schweren neurologischen Erkrankung darstellen. Mit Bezug auf pathophysiologische Aspekte werden praktische Hinweise für die differenzialdiagnostische Läsionsinterpretation im klinischen Alltag aufgezeigt.


Ergebnisse Anhand charakteristischer bildmorphologischer und klinisch-neurologischer Befundkonstellationen sind die wichtigsten und häufigsten vaskulären, entzündlichen, metabolischen und neoplastischen Krankheitssentitäten dargestellt und die für die differenzialdiagnostische Zuordnung essenziellen Details hinsichtlich Signalverhalten und Lokalisation hervorgehoben.
In adults, cerebral white matter lesions are a common MRI finding in the clinical routine [1, 2]. They can be either non-specific and age-related or indicative of the onset of severe neurological disease [1–4]. The image morphology is partially overlapping, with the hyperintense signal on the T2-weighted sequences (T2 WI) representing the common feature of these changes. Consequently, clinical-neurological data and laboratory chemical findings including CSF analysis are essential for differential diagnostic classification [1, 2, 5, 22]. The aim of this review is to explain diagnostic aspects of the classification of cerebral white matter lesions and to provide differential diagnostic tips with special regard to the vascular architecture of the white matter and pathological processes [6, 7, 10, 11].

### Technical Aspects

The MR sequences listed in ▶ Table 1 are necessary for the most reliable classification of white matter lesions. Diffusion tensor imaging (DTI), MR spectroscopy (MRS) and perfusion measurements are used as additional diagnostic tools, especially in cases of ambiguous diagnostic findings [1, 2, 5, 23]. Diffusion-weighted imaging (DWI) is acquired with diffusion gradients oriented in 3 orthogonal directions, which form the basis of directionally averaged DWI images (trace maps) [24]. The trace maps show the extent of diffusion of hydrogen protons, but not their directional dependence (anisotropy). The strength of diffusion weighting is described by the b-value [s/mm²], which is calculated from the properties of the diffusion gradients. Since the measured diffusion rates depend both on the chemical and physical tissue properties as well as the measurement conditions, the calculated diffusion values are referred to as apparent diffusion coefficient (ADC) [24]. Measurements with at least two different b-values are required to create ADC parameter images; b-values between 0 (pure T2-weighted image) and 1000 s/mm² are used for DWI measurements of the brain. Although two b-values are sufficient to create an ADC map, measurements with three b-values (b = 0, b = 500, b = 1000; each [s/mm²]) allow a more precise calculation of the ADC values. The lower signal-to-noise ratio (SNR) of the images with b = 1000 s/mm² leads to higher measurement inaccuracies, which are partly compensated by measurement with b = 500 s/mm². Compared to the ADC maps, the DWI images have the disadvantage that in the case of strongly T2-hyperintense changes on the DWI images, diffusion restriction is simulated (so-called “T2-shine through”), which is eliminated in the ADC maps [24].

In contrast to DWI, the DTI measures the anisotropy [25]. For this purpose, the diffusion images are measured with diffusion gradients oriented in at least 6 different spatial directions. Since the diffusion in the white matter through the fiber paths running in it is strongly directed along the fiber path, the parameters for the extent and direction of the anisotropy that can be calculated from the DTI are pathological early in most diseases of white matter [25].

In principle, 3D sequences offer the advantage of capturing small lesions more sensitively than 2D images with higher layer thickness because of the higher SNR, the high spatial resolution and the missing layer gaps [26]. In addition, 3D measurements in all planes can be reconstructed simply, curved multplanar or as maximum intensity projection (MIP). A disadvantage compared to the 2D sequences is the usually longer measurement times with resulting motion sensitivity, whereas flow artifacts are usually more pronounced in 2D FLAIR (fluid attenuated inversion recovery) images than in the 3D images [26].

### Vascular Anatomy of White Matter

1. **Terminal pial and medullary arteries (4–5 cm long)**

   They originate from the three large leptomeningeval arteries (anterior, middle and posterior cerebral arteries) and move perpendicularly through the cortex into the white matter. Due to only a few capillary anastomoses, they represent functional end arteries (see ▶ Fig. 1) [7, 10, 27].

2. **Subependymal arteries**

   These arise close to the ventricle from the choroidal arteries, which also extend perpendicularly into the deep white matter and are significantly shorter than the pial medullary vessels

3. **End-arteries of the medial and lateral lenticulostrate and thalamic perforators**

   Due to only a few capillary anastomoses, they represent functional terminal arteries (see ▶ Fig. 1) [7, 10, 28]. Thus, the deep white
matter is especially in the centrum semiovale and nearby the anterior horns border zone between superficial pial, deep subependymal and basal lenticulostriate and thalamo-perforating arteries [10]. This makes these deep white matter regions particularly vulnerable to vascular compromise (including “Steiner’s Wetterwinkel”). In contrast, the juxtacortical region with the U-fibers is better vascularized than the deep white matter due to the cortical network of arterioles and numerous anastomoses [7, 10].

<table>
<thead>
<tr>
<th>weighting</th>
<th>geometry</th>
<th>sequence</th>
</tr>
</thead>
<tbody>
<tr>
<td>T2</td>
<td>axial and sagittal</td>
<td>T2 spin echo (FSE, TSE)</td>
</tr>
<tr>
<td></td>
<td>axial 2D, possibly sagittal 3D with axial reconstructions</td>
<td>FLAIR</td>
</tr>
<tr>
<td>diffusion</td>
<td>axial</td>
<td>DWI with 3 spatial directions and at least 2 b-values (0, 1000 s/mm²)</td>
</tr>
<tr>
<td>T1</td>
<td>axial</td>
<td>spin or gradient echo native, possibly also after IV administration of contrast</td>
</tr>
<tr>
<td></td>
<td>3D</td>
<td>TOF²</td>
</tr>
<tr>
<td>T2⁺</td>
<td>axial</td>
<td>SWI or T2 gradient echo</td>
</tr>
</tbody>
</table>


1 Especially helpful for the detection of corpus callosum lesions and pattern recognition.
2 Intravenous administration of contrast medium is not routinely required, but depends on the pattern of lesions in the native T1 WI, taking into account the clinical symptoms and the specific issue.
3 TOF-MRA sequences are T1-weighted. Thus, substances with short T1 times (hematoma, gadolinium, fat) can cause a “shine through” effect and simulate a flow signal, which can then be misinterpreted in MIP reconstructions as a flow signal or vessel malformation.
4. Medullary white matter veins

Arterial and venous vascular systems run parallel in the white matter. Contrary to the arterial vascular supply, the veins penetrating the cortex are shorter and consequently the deep medullary veins draining to the medial center are longer, so that the venous watershed is closer to the brain surface [29].

5. Histological structure

a) Medullary white matter arteries

In the section near the origin, the terminal pial and medullary arteries are surrounded by pia mater and the subpial space to the brain parenchyma is delimited by the glia limitans [30]. Due to high cell density, this is very narrow at the level of the cortex and becomes subcortically widened [30, 31]. The lenticulostriate and thalamic perforators are surrounded by two leptomeningeal layers [28, 30].

b) Medullary white matter veins

The medullary veins are not enclosed by a pial layer; thus, the perivenous space communicates with the superficial subpial compartment [29].

c) Anterior outer edge of lateral ventricle (“Steiner’s Wetterswinkel”)

The white matter adjacent to the anterior horn and the cella media of the lateral ventricles is separated from the cerebrospinal fluid space by an only incompletely formed ependymal layer. This structure facilitates CSF diapedesis and causes age-related physiological appearance of hyperintense caps and strips on the T2 WI [6].

Para- and Perivascular Spaces (PVS)

Although the arterial subpial perivascular space is separated from the superficial subarachnoid space by the pia mater and filled with interstitial fluid [30], it appears CSF-isointense on the T2 WI, on FLAIR and on T1-weighted sequences (Virchow-Robin spaces, see ▶ Fig. 2, 3), and punctiform or elongated depending upon the section of the course of the penetrating arteries [1, 7–9]. These perivascular spaces can be cystically dilated (see ▶ Fig. 4) and in individual cases in mesencephalic position they can cause a disturbance of CSF circulation by constriction of the aqueduct [1, 9].

In addition to non-pathological congenital PVS size variations, increased dilation of these spaces in old age sometimes appears to be the result of impaired drainage of the interstitial fluid (lymphatic system) due to microangiopathy and represents possibly associated vascular induced impairment of cognition in the elderly [4, 23, 31–38]. The deposit of amyloid in the vessel walls near the cortex also have an amplifying effect. Expanded PVS can also occur in the context of metabolic diseases (see ▶ Fig. 5) [1, 15] and pathogen-induced inflammatory CNS diseases (see ▶ Fig. 6) [39]. The most common causes of cerebral white matter lesions are discussed below.
Microangiopathy (small vessel disease)

1. Vascularly-induced white matter signal changes (white matter changes; WMC)

PVS must be distinguished from vascularly-induced gliosis (see Fig. 3) which appears hyperintense on FLAIR sequences [1, 5, 7, 8]. Defective residuals after lacunar infarctions are CSF isointense and often have a narrow T2 hyperintense rim due to gliosis. They are oval or round-shaped with a longitudinal diameter ≤ 15 mm [3, 5, 7].

Vascular WMC are said to be caused by chronic hypoperfusion [40]. They typically occur bilaterally and symmetrically; 3 preferred regions are defined, whereby the perfusion areas in the terminal sections of the perforators play an important role: a) periven-

tricularly, b) in the deep white matter (centrum semiovale) and c) juxta- (sub-) cortically [10, 36]. The semiquantitative evaluation scale according to Fazekas et al. is often used [41, 42], whereby Grade 1 describes several punctiform lesions, Grade 2 partially confluent and Grade 3 extensive flat lesions (see Fig. 7).

As the volume of WMC increases, the risk of neurological functional deficits, infarcts, dementia and death increases [3, 4, 23, 32, 36, 43, 44]. While in age groups over 60 years WMC are typically found without a clinical correlate [35, 42], and some authors define age-associated WMC especially from the age of 75 years onward, there is no consistent information in the literature about the onset of these changes [43]. A higher incidence of WMC and possibly additional ovoid lesions in the border regions has also been described in patients with migraine with aura [45].

Fig. 3 a–c Comparative signalling of anatomical and pathological structures. Dilated perivascular spaces (PVS) (a: T2 WI ax.; b: FLAIR ax., arrows) with sharply delineated cerebrospinal fluid (CSF) isointense signal; vascular gliosis with hyperintense signal changes (a, b: arrowhead). In addition, numerous subcortical hypointense lesions (microbleeds; c: arrowhead) in a patient with a brain-organic psychosyndrome due to cerebral amyloid angiopathy (CAA).

Fig. 4 a–c Cystic dilated perivascular spaces (PVS) in a neurologically unremarkable patient (a: T2 WI ax.; b, c: T1 WI ax. and sag., arrow).
According to the etiopathogenetic classification of microangiopathies defined by Pantoni [3], age-associated arteriosclerosis with typical vascular risk factors (hypertension, diabetes mellitus) dominates as Type 1 and sporadic and hereditary cerebral amyloid angiopathy (CAA) as Type 2 (see ▶ Fig. 3); together they comprise more than 90% of microangiopathies. A common imaging characteristic of these etiologies is that white matter lesions usually spare the corpus callosum [46–48]. Type 3 describes other genetic microangiopathies (without CAA) such as CADASIL (Cerebral Autosomal Dominant Arteriopathy with Sub-

▶ Fig. 5 a–c Mucopolysaccharidosis. Enlarged perivascular spaces (PVS) (a: T2 WI ax.; b: FLAIR ax.; c: T1 WI ax., arrow) due to metabolite deposition (“Hurler holes”) with accompanying glial reaction (a, b: arrowhead).

▶ Fig. 6 a, e Microangiopathic hyperintense lesions (a: T2 WI ax., arrowhead) with microbleeds (e: T2* WI ax., arrow); b, f: enlarged PVS in mucopolysaccharidosis (“Hurler holes”; b: T2 WI ax.; f: FLAIR ax.; arrowhead, arrow); c, g: Cryptococcosis; hyperintense lesions (c: T2 WI ax., arrowhead) in the course of lenticulostriate perforators with punctiform diffusion restrictions (gelatinous pseudocysts; g: DWI ax., b = 1000 s/mm², arrows); d, h: lymphoma with perivascular spread and enhancement in the left hemisphere (h: pc T1 WI ax., arrowheads), consecutive asymmetrical perivascular spaces (PVS) on T2 WI (d: arrowhead) in left-right comparison.

Weidauer S et al. White Matter Lesions... Fortschr Röntgenstr
cortical Infarcts and Leukoencephalopathy) (see Fig. 8) [40, 49], MELAS (Mitochondrial Encephalomyopathy with Lactic Acidosis and Stroke-like episodes) or Fabry disease [50, 51]. Type 4 includes inflammatory and immunologically mediated angiopathies; Type 5, venous collagenosis; and Type 6, other microangiopathies [52–54].

2. Lacunar infarcts

Follow-up examinations of cerebral microangiopathies have shown that a) incidental lacunes typically develop close to the edge of WMC, taking into account the course of the perforators [40]; b) the increase in WMC develops from periventricular to central subcortically, and c) WMC increases around an incidental lacuna [40]. However, lacunes can also be caused by a macroangiopathy with atheromatous compromise of the perforator orifices, e.g. at the M1 segment of the middle cerebral artery, or by an embolism [3, 4, 33, 35, 55]. In the acute and subacute phase, DWI with evidence of a diffusion disorder due to circumscribed cytotoxic edema allows a reliable differentiation from pre-existing chronic WMC (see Fig. 9) [1, 5]. Macroangiopathy with corresponding upstream hemodynamically effective stenosis and possibly additional lesions in the border zone or terminal vascular bed is often present in the case of asymmetric WMC (see Fig. 10).

However, it is important to remember here that both autoimmune-associated inflammations such as multiple sclerosis (MS)
Fig. 8  a–c Axial T2 WI in a 45 year old woman suffering from CADASIL showing partially confluent and nearly symmetrical hyperintense lesions in the deep white matter (a: arrow), in the external capsule (b: black arrowhead) and the anterior temporal pole (c: black arrow); involvement of the U-fibres (a, c: white arrowhead).

Fig. 9  a–d Extended white matter changes (WMC) (a, b: T2 WI and FLAIR ax.) and additional acute lacunar infarct on the edge of the WMC (c, d: DWI ax; b = 1000 s/mm²; ADC-map).
or NMOSD (Neuromyelitis Optica Spectrum Disorders) [59, 60] as well as pathogen-related infections can cause a diffusion restriction in the acute phase, although it is usually lesser compared to an acute cerebral infarction (see ▶ Fig. 11) [39]. In contrast to cytotoxic edema, vasogenic edema, particularly in blood-brain barrier disruption, typically shows an anatomically predetermined extension along fascicles and fiber structures (see ▶ Fig. 12) [2, 6].

3. Microbleeds (MB)

The detection of MB with blood-sensitive sequences such as T2*-weighted gradient echo (GRE) sequences or susceptibility-weighted imaging (SWI) with frequently punctiform signal cannellation as well as evaluation of their localization are essential for differential diagnostic assessment [46–48, 61]. MB in the basal ganglia, thalamus, pons or cerebellum are usually caused by hypertensive vasculopathy with lipohyalinosis of the perforators; signal reductions correspond to hemosiderin deposits in the degeneratively altered vessel walls [7, 48]. Intracerebral bleeding typically occurs here, especially in cases of arterial hypertension. Often there are also dilated PVS up to so-called status cribrosus ("état criblé") and with sufficient high microscopic resolution the previously described "Charcot-Bouchard aneurysms" correspond to elongated and torqued perforators [8, 9, 46]. Differential diagnostically, cavernomas and parasitic/infectious diseases must be distinguished from MB.

[12, 56–58]
Subcortical MB suggest CAA. In addition to microhemorrhages, the revised Boston criteria [47, 48] also include focal cortical superficial siderosis (see Fig. 13) [61]. In CAA, small cortical and subcortical arteries with a diameter < 500 μm, capillaries, and to a lesser extent veins, are affected by amyloid deposits in the vessel walls. In the course of the disease, the neocortex in front of the allocortical areas is affected, and there is a close correlation with Alzheimer’s disease (AD) in parieto-occipital and temporal position (see Fig. 13) [23, 31, 62]. One variant is CAA-associated inflammation (CAA-RI), which histologically is

![Fig. 11 a–c Cytotoxic edema due to autoimmun-associated inflammation in a 34-year-old woman suffering from neuromyelitis optica spectrum disorders (NMOSD) with progressive hemiparesis re. within 3 hours. Axial T2 WI a disclose a polycyclic hyperintense lesion with broad contact to the left lateral ventricle (arrow); diffusion restriction (b: DWI; b = 1 0 0 0 s/mm², arrow) and expansion along the corticospinal tract (c: T2 WI sag., arrow, arrowhead).]

![Fig. 12 a–d “Finger-shaped” vasogenic edema (a–d: T2 WI ax., black arrows) corresponding to the course of the longitudinal inferior fascicle due to malignant glioma temporal left (a: arrowheads) with blood-brain barrier disruption.]

![Fig. 12 a–d “Finger-shaped” vasogenic edema (a–d: T2 WI ax., black arrows) corresponding to the course of the longitudinal inferior fascicle due to malignant glioma temporal left (a: arrowheads) with blood-brain barrier disruption.}

Weidauer S et al. White Matter Lesions... Fortschr Röntgenstr
perivasculitis and associated with extensive, sometimes space-occupying lesions in the white matter (see ▶ Fig. 14) [63, 64]. The lesion type is similar to that described in therapy studies with monoclonal antibodies against amyloid beta 42 (Aβ-42) in patients with AD [65]. The authors distinguish a hemorrhagic and an encephalitic variant (Amyloid-Related Imaging Abnormalities; ARIA-H, ARIA-E). For the sake of clarity, amyloid-β related angiitis (ABRA) and its differentiation from primary CNS angiitis (PCNSA) will not be discussed in detail; the latter exhibits no amyloid deposits [54, 63, 64]. If MB are subcortical and/or in deep white matter adjacent to various old (lacunar) infarcts and additional WMC, differential diagnosis should consider vasculitis of possibly only the small vessels (Pantoni Type 4) [3, 52]. In this context, reference is also made to the revised Chapel Hill 2012 criteria on systemic vasculitis [53].
Angiocentric Propagation Pattern

Vascular-associated diseases as well as inflammatory and neoplastic processes may exhibit an angiocentric lesion pattern on T2 WI due to the anatomically predetermined propagation along the PVS, which is illustrated by taking into account the T1 WI after contrast application (pc T1 WI) [39, 66–69]. In addition to sarcoidosis (see Fig. 15) [67, 68], in particular fungal infections [39], primary CNS lymphoma (PCNSL) as well as related diseases (e.g. lymphomatoid granulomatosis) [70] show a streaky or punctiform enhancement, with possibly continuous Gadolinium enhancement of the perivascular leptomeningeal structures along the course of the perforators with differently pronounced perifocal edematous (T2-hyperintense) reaction and possible circumscript diffusion restriction due to inflammation, high cell density or a resulting infarction.

Perivascular Inflammatory Lesions in Multiple Sclerosis (MS)

Multiple sclerosis (MS) is the most common demyelinating disease. The 2017 revisions of the MRI criteria for diagnosis define four characteristic regions for dissemination in space (DIS): a) juxta-/cortical, b) periventricular, c) infratentorial and d) spinal [12, 57, 58, 71]. Spatial dissemination requires at least one T2 hyperintense lesion in at least two regions, whereby a longitudinal extension of ≥3 mm is required e.g. periventricularly for lesions that often run perpendicularly to the ventricular wall (Dawson’s finger; see Fig. 16, 17) [57, 58]. Correlating with the histological description, the central vein in the plaques can be delineated at higher field strengths on SWI [12]. The simultaneous presence of a contrast enhancing lesion and a non-enhancing lesion, or a new lesion that occurs in the course of the disease, are criteria for dissemination in time (DIT) [57]. In contrast to atherosclerotic microangiopathy, U-fibers are also affected in juxtacortical MS lesions [57, 72].

In the case of acute demyelinating encephalomyelitis (ADEM), numerous monomorphic lesions, often with blurred boundaries, are clearly visible subcortically and in the basal ganglia with marginal contrast enhancement open towards the cortex (“open ring”) [11, 74]. A marked perivenous orientation of white matter lesions is also described in Behçet’s disease and in ANCA (antineutrophil cytoplasmic antibodies) associated vasculitis of the small vessels (e.g. microscopic polyangiitis) corresponding to its manifestation in capillaries and veins [75–77]. Regarding the ex-
Fig. 15  a–c Angiocentric lesion pattern in sarcoidosis. Symmetrical blurred lesions in the pons (a: T2 WI ax., arrow) with enhancement (b, c: pc T1 WI ax. and sag.; arrows) in the course of the perforators; note also superficial pial enhancement (b, c: arrowheads).

Fig. 16  a–f Multiple sclerosis (MS): Juxtacortical, periventricular (a, b: FLAIR ax., arrows) and infratentorial lesions (c: T2 WI ax.: arrow: trigeminal nerve nucleus; arrowhead: trigeminal nerve); d (FLAIR ax.) and e (SWI ax.): periventricular plaque (d: arrow) with central vein (e: arrow); f (T1 WI ax.): “black holes” (arrows).
tensive differential diagnosis of MS and atypical course variants, reference is made to the further special literature [68, 75].

Extensively Spread Lesions

The differential diagnosis of larger extended, often homogeneous white matter lesions includes metabolic (e.g. leukodystrophies, vitamin B12 deficiency, others) [22, 23, 78], inflammatory (e.g. HIV encephalopathy, progressive multifocal leukoencephalopathy; PML) [79–82], toxic (e.g. heroin-induced spongiform leukoencephalopathy) as well as radiogenic causes [18, 19]. Assessment of U-fiber involvement in addition to symmetry is necessary for a closer differential-diagnostic classification. PML emanating from the oligodendrocytes can be the overture to previously unrecognized immunosuppression due to HIV disease [39, 82]. Typical asymmetrical white matter changes are not space-occupying; they expand extensively along fiber tracts and involve the U-fibers (see Fig. 19) [80, 83]. T1 WI show a signal reduction and, depending on the immune status or the presence of an immune reconstitution inflammatory syndrome (IRIS), contrast accumulation and diffusion restriction may be present at the edges. In the case of therapy-related compromised immune system (e.g. natalizumab), a punctiform perivascularly oriented enhancement can occur in the vicinity of hypointense lesions on pc T1 WI (“milky way”) [83, 84]. In contrast, HIV-induced progressive diffuse leukoencephalopathy (PDL) typically spares the U-fibers (see Fig. 19e), is rather symmetrical and non- to slightly-hypointense on T1 WI. HIV encephalitis as an acute variant also affects the gray matter [39, 79].

Of the large heterogeneous group of leukodystrophies, metachromatic leukodystrophy (MLD) and adrenoleukodystrophy (ALD) deserve special mention, since both can also manifest clinically in advanced age [13, 22, 68]. Symmetrical, homogeneous changes in the white matter that leave out the U-fibers are apparent in the adult variant of MLD, which occurs in approx. 15% of cases and exhibits clinically slowly progressive psychiatric symptoms (see Fig. 19). ALD preferentially affects the parietooccipital white matter, shows marginal contrast enhancement depending on the stage of the disease and spreads into the descending fiber tracts [22]. A disorder related to vanishing white matter diseases (VWMD) can occur at any age. Mitochondriopathies such as MELAS or Kearns-Sayre syndrome (KSS) often also exhibit cortical involvement [50].
It is important to note that advanced chronic vascular and inflammatory etiologies may also exhibit extensive and largely symmetrical lesion patterns in the final stage [2].

**Cell or System/Fiber Tract-associated Lesions**

As an example from the group of neurodegenerative diseases, Waller’s degeneration in amyotrophic lateral sclerosis (ALS) with variable involvement of the 1st and 2nd motor neuron is described here [85]. This results in a symmetrical signal increase on T2 WI in the course of the corticospinal tract starting directly below the primary motor cortex (see ▶ Fig. 20). The ALS dementia complex preferentially discloses frontotemporal atrophy corresponding to a Tau protein negative, TDP-43 positive (TAR-DNA binding protein) frontotemporal dementia [23, 86]. Waller’s degeneration of the corticospinal tracts can also occur after damage to the 1st motor neuron due to other causes, e.g. after (sub-)cortical infarcts [85].
Checklist for Image Analysis and Diagnostic Algorithms

Table 2 and Fig. 21 summarize various imaging findings and corresponding differential diagnostic aspects. The age of the patient, the progression of the disease and the clinical and neurological findings as well as previous illnesses and laboratory parameters must be taken into account.

Conclusion

Comparative image analysis of different identically angulated MRI sequences including DWI and T2* WI or SWI is a prerequisite for the reliable differential diagnostic assessment of white matter lesions. Characteristic lesion patterns can be identified, taking into account the architecture and vascular supply of the white matter. The etiology of white matter lesions includes vascular, inflammatory, metabolic and neoplastic processes. Vascular pathologies are in the foreground, especially in the elderly; age-associated arteriosclerosis with typical risk factors and CAA account for over 90% of microangiopathies. Physiological changes such as dilated...
Fig. 20 a–c T2WI showing symmetric system associated hyperintense signal changes in the course of the corticospinal tract: in the subcortical region of the precentral gyrus (a: arrow heads), in the internal capsule (b: arrows) and in the cerebral peduncle (c: arrows) due to amyotrophic lateral sclerosis (ALS).

Fig. 21 White matter lesions in adults: hints for the differential diagnosis. ADEM: Acute demyelinating encephalomyelitis; ALD: Adrenoleukodystrophy; CAA: Cerebral amyloid angiopathy; CADASIL: Cerebral autosomal dominant arteriopathy with subcortical infarcts and leukoencephalopathy; CC: Corpus callosum; DIS: Dissemination in space; DIT: Dissemination in time; DWI: Diffusion weighted imaging; IRIS: Immune reconstitution inflammatory syndrome; MELAS: Mitochondrial encephalomyopathy with lactic acidosis and stroke-like episodes; MLD: Metachromatic leukodystrophy; MS: Multiple sclerosis; NMOSD: Neuromyelitis optica spectrum disorders; PCNSA: Primary central nervous system angiitis; PML: Progressive multifocal leukoencephalopathy; WMC: White matter changes.
PVS and sometimes age-associated WMC are to be separately considered. Involvement of U-fibers or the cortex is more often found in inflammatory or metabolic causes, although the latter are rare in adulthood.

Conflict of Interest

The authors declare that they have no conflict of interest.

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