Differential diagnosis and evaluation in pediatric inflammatory demyelinating disorders

ABSTRACT
Major advances have been made in the clinical and radiologic characterization of children presenting with the different forms of an acquired inflammatory demyelinating syndrome (ADS) such as acute disseminating encephalomyelitis, neuromyelitis optica spectrum disorders, and clinically isolated syndromes. Nevertheless, a proportion of cases that present with similar symptoms are due to a broad spectrum of other inflammatory disorders affecting the white matter, primary CNS tumors, or neurometabolic diseases. The clinician therefore has to be aware of the different forms of ADS, the risk factors for a chronic-relapsing course, and features that indicate an alternative diagnosis. The goal of this article is therefore to provide an outline of a pathway for evaluating pediatric patients with a presumed inflammatory demyelinating disorder and discussing the spectrum of the more common differential diagnoses. Neurology® 2016;87 (Suppl 2):S28-S37

GLOSSARY

ADEM = acute disseminating encephalomyelitis; ADS = acquired demyelinating syndromes; ANE = acute necrotizing encephalopathy; AQP4 = aquaporin-4; BBE = Bickerstaff brainstem encephalitis; BBGD = biotin-responsive basal ganglia disease; CIS = clinically isolated syndromes; DIS = dissemination in space; DIT = dissemination in time; GBS = Guillain-Barré syndrome; HLE = hemophagocytic lymphohistiocytosis; IgG = immunoglobulin G; LCH = Langerhans cell histiocytosis; LETM = longitudinally extensive transverse myelitis; MOG = myelin oligodendrocyte glycoprotein; MS = multiple sclerosis; NMOSD = neuromyelitis optica spectrum disorders; ON = optic neuritis; PACNS = primary angiitis of the CNS; PCNSL = primary CNS lymphoma; PML = progressive multifocal leukoencephalopathy; SLE = systemic lupus erythematosus; TM = transverse myelitis.

Acquired demyelinating syndromes (ADS) represent a group of disorders such as acute disseminating encephalomyelitis (ADEM), neuromyelitis optica spectrum disorders (NMOSD), and clinically isolated syndromes (CIS). A timely and precise diagnosis is mandatory in order to ensure adequate patient management, but due to the extensive differential diagnosis, it is not uncommon that children are assigned an incorrect diagnosis.

The focus of this review is to describe and delineate the different subtypes of ADS from the broad spectrum of other inflammatory disorders of the white matter, genetic disorders affecting the innate and adaptive immune system, primary CNS tumors, and neurometabolic diseases.

ACQUIRED DEMYELINATING SYNDROMES
The clinical picture of ADS comprises ADEM, characterized by multifocal deficits and encephalopathy; CIS, characterized by multifocal or subacute deficits without encephalopathy; pediatric multiple sclerosis (MS); and NMOSD (table 1). ADS occur as a monophasic illness, but may also represent the onset of a chronic relapsing disorder.

Among the various forms of ADS, ADEM offers the widest spectrum of differential diagnosis. The clinical picture of ADEM is characterized by a multifocal onset, associated with encephalopathy.1 Encephalopathy in ADEM is defined as a qualitative or quantitative disturbance of consciousness, such as irritability, somnolence, or coma, and must occur in the absence of fever. MRI is an important tool in distinguishing ADEM from other diseases and in particular from the first attack of MS.2 Lesions found on MRI are typically diffuse, ill-defined, >1–2 cm in size, and in the cerebral white matter, but may also affect the thalamus, basal ganglia, and spinal cord. T1-hypointense lesions are rare.3 Among the patients presenting with ADS, ADEM is found more...
Table 1  Acute demyelinating syndromes and recommended workup

<table>
<thead>
<tr>
<th>Conditions</th>
<th>Consider following investigations (in addition to MRI brain/spine)</th>
</tr>
</thead>
<tbody>
<tr>
<td>ADEM</td>
<td>CBC, liver enzymes, renal function, ESR, complement (C3, C4), ANA, dsDNA, 25(OH) vitamin D, ferritin, triglycerides, ACE, AQP4, MOG antibodies, bacterial and viral studies as indicated (blood, throat swab, stool)</td>
</tr>
<tr>
<td>NMO/DS</td>
<td>CSF studies with cell count, protein, glucose, oligoclonal bands, cytology, MRZ reaction; bacterial and viral studies as indicated</td>
</tr>
<tr>
<td>CIS</td>
<td>Evoked potentials (visual, somatosensory, auditory), ophthalmology with OCT, neuropsychological testing, urology, rheumatology</td>
</tr>
</tbody>
</table>

Abbreviations: ACE = angiotensin-converting enzyme; ADEM = acute disseminating encephalomyelitis; ANA = antinuclear antibodies; AQP4 = aquaporin-4; CBC = complete blood count; CIS = clinically isolated syndromes; dsDNA = double-stranded DNA; ESR = erythrocyte sedimentation rate; MRZ = measles, rubella, varicella zoster virus; OCT = optical coherence tomography; MOG = myelin oligodendrocyte glycoprotein; NMO/DS = neuromyelitis optica spectrum disorders.

frequently in younger children. Typically ADEM is a monophasic disorder, but may also represent the first attack of pediatric MS.

CIS are characterized by monofocal or polyfocal events without encephalopathy, unless explained by fever. The most common clinical presentations of CIS in children are optic neuritis (ON), transverse myelitis (TM), or brainstem/cerebellar syndromes or symptoms attributed to the cerebral hemispheres. Unilateral ON is defined by acute/subacute and painful visual loss, impairment of visual acuity and color perception, MRI evidence of optic nerve swelling, or abnormal signal or enhancement. Bilateral ON is more common in ADEM or NMO/DS. TM is characterized by acute/subacute onset of bilateral symmetric or asymmetric sensory or motor deficits, sometimes associated with sphincter dysfunction, with a defined sensory level, and MRI evidence of spinal cord swelling or abnormal signal intensity or enhancement. If brain MRI is normal at the time of a CIS, the risk of MS is low.

In contrast, in case of abnormal MRI studies, the distribution and morphology of lesions predicts the risk of a second clinical event with high sensitivity and specificity. Furthermore, the presence of oligoclonal bands in pediatric ON increases the risk of MS.

Pediatric MS is defined by 2 or more non-ADEM episodes of presumed inflammatory demyelinating origin, separated by >30 days and involving more than one CNS site or a first nonencephalitogenic episode with an MRI fulfilling the McDonald criteria for dissemination in space (DIS) and with a follow-up MRI showing a new enhancing or nonenhancing lesion consistent with the 2010 dissemination in time (DIT) criteria. In children <12 years of age, a single, non-ADEM episode with MRI lesions fulfilling the 2010 McDonald criteria for DIS and DIT may also satisfy the criteria for pediatric MS. In a child with an ADEM attack that is followed by a second non-ADEM event, 3 or more months after the first episode, and new MRI lesions that fulfill the 2010 criteria for DIS and DIT, the diagnosis of pediatric MS can be assigned.

ON or TM are the main presenting features of NMO/DS. Other core clinical characteristics comprise an acute area postrema syndrome (intractable hiccups or vomiting), an acute brainstem syndrome, or symptomatic diencephalic (symptomatic narcolepsy) or cerebral syndrome with NMO/DS-typical MRI lesions in conjunction with aquaporin-4 (AQP4) antibodies. While many clinical, imaging, and CSF features are similar to those found in adult NMO/DS, some caveats remain. In particular, longitudinally extensive transverse myelitis (LETM) may be less specific for pediatric NMO, since LETM also may be found in pediatric MS, as well as in monophasic ADEM. Recently, pediatric cases with the clinical picture of ADEM, accompanied by the presence of AQP4 antibodies, also have been described.

Among children with ADS, a subgroup of patients shows serologic evidence for myelin oligodendrocyte glycoprotein (MOG) antibodies. Children with MOG-associated demyelinating CNS disease may present with various clinical features, including ADEM, ADEM followed by ON, or NMO/DS. MOG-associated ADEM in children shows a uniform radiologic pattern with large, hazy lesions, the absence of abnormal MRI features, and involvement of more CNS areas including spine LETM.

Recent studies showed that the presence of MOG antibodies argues against a further diagnosis of MS. However, the long-term evolution of children with MOG antibodies needs further prospective evaluation.

RED FLAGS SUGGESTING AN ALTERNATIVE DIAGNOSIS IN CHILDREN WITH ADS In children with an acute demyelinating event, the following red flags should challenge the diagnosis of ADS: onset before age 1 year; history of developmental delay; consanguineous family; family history of severe, acute neurologic symptoms; gradual progression of symptoms; and multisystemic involvement (table 2). Unusual MRI features that are infrequent in ADS are a single supratentorial white matter lesion, symmetric white matter involvement, or lesions restricted solely to the brainstem and basal ganglia (table 3).
even conventional angiography can be normal despite the involvement of different organ systems in combination with anemia, thrombocytopenia, and the presence of antibodies to double-stranded DNA, or high anti-phospholipid antibodies. Few studies exist describing the MRI findings in children with SLE, which can include nonspecific T2 signal alterations in particular at the beginning of the disease, larger areas of demyelination, and TM in addition to cortical and subcortical strokes.16,17

Behçet disease. Behçet disease is a systemic disorder with neurologic symptoms in 20% of pediatric cases.18 The neurologic presentations can be variable and the sole presenting sign of the disease. The neurologic manifestations are often linked to high intracranial pressure due to cerebral venous sinus thrombosis, inflammation of the white matter, or stroke due to vasculitis. The course of the disease is relapsing, sometimes despite treatment. Cerebral MRI can demonstrate venous sinus thrombosis and less often white matter lesions reminiscent of MS, which is more common in adults.18 CSF studies are normal in up to 50% of cases.

Primary angiitis of the CNS. The diagnosis of PACNS is based on 3 criteria: (1) an acquired neurologic deficit, (2) angiographic or histologic evidence of cerebral vasculitis, and (3) absence of systemic

### Table 2  Clinical red flags for conditions other than acquired demyelinating syndromes

<table>
<thead>
<tr>
<th>Condition</th>
<th>Clinical red flags</th>
</tr>
</thead>
<tbody>
<tr>
<td>Multisystemic involvement</td>
<td>HLH, Behçet disease, CNS vasculitis, SLE, mitochondrial diseases, sarcoidosis, Sjögren syndrome, LCH, infections</td>
</tr>
<tr>
<td>Spastic paraplegia</td>
<td>SLE, Sjögren syndrome, Lyme disease, West Nile virus, vitamin B12 deficiency, spinal cord tumor/ischemia/AVM/trauma, familial spastic paraplegia, AMN, Krabbe disease, Alexander disease</td>
</tr>
<tr>
<td>Protracted headache ≥ stroke-like episodes</td>
<td>SLE, MELAS, MERFF, HIV, malignancy, CADASIL, HLH, CNS vasculitis, Fabry disease, Susac syndrome</td>
</tr>
<tr>
<td>Epilepsy</td>
<td>Mitochondrial diseases, B12, metabolism disorders</td>
</tr>
<tr>
<td>Ataxia</td>
<td>NPC, SCA, PDH</td>
</tr>
<tr>
<td>Extrapyramidal symptoms</td>
<td>Anti-NMDAR encephalitis, Wilson disease, mitochondrial diseases, biotinidase deficiency, biotin-responsive basal ganglia disease, LBSL</td>
</tr>
<tr>
<td>Psychiatric symptoms</td>
<td>Primary CNS angiitis, anti-NMDAR encephalitis, B12 metabolism disorders, mitochondrial disorders, SLE, Susac syndrome</td>
</tr>
<tr>
<td>Progressive disease with dementia</td>
<td>Leukodystrophies, mitochondrial diseases</td>
</tr>
<tr>
<td>Hypothalamic dysfunction</td>
<td>Sarcoidosis, LCH</td>
</tr>
<tr>
<td>Cranial nerve neuropathies</td>
<td>Krabbe disease, MLD, Alexander disease, Lyme disease, sarcoidosis</td>
</tr>
<tr>
<td>Peripheral neuropathy</td>
<td>Krabbe disease, MLD, mitochondrial disorders, ALD/AMN</td>
</tr>
<tr>
<td>Sensorineural hearing loss</td>
<td>Susac syndrome, mitochondrial diseases</td>
</tr>
<tr>
<td>Optic neuropathy</td>
<td>LHON, MELAS, MERFF, OPA-1</td>
</tr>
</tbody>
</table>

Abbreviations: ALD = adrenoleukodystrophy; AMN = adrenomyeloneuropathy; AVM = arteriovenous malformation; CADASIL = cerebral autosomal dominant arteriopathy with subcortical infarcts and leukoencephalopathy; LBSL = leukoencephalopathy with brainstem and spinal cord involvement and elevated lactate; HLH = hemophagocytic lymphohistiocytosis; LCH = Langerhans cell histiocytosis; LHON = Leber hereditary optic neuropathy; MELAS = mitochondrial encephalomyelopathy with lactic acidosis and stroke; MERFF = myoclonic epilepsy with ragged red fibers; MLD = metachromatic leukodystrophy; NMDAR = NMDA receptor; NPC = Niemann-Pick C; OPA = optic atrophy; PDH = pyruvate dehydrogenase deficiency; SCA = spinocerebellar ataxia; SLE = systemic lupus erythematosus.
vasculitis. PACNS affects the medium-sized and small vessels and remains a rare and difficult diagnosis. Clinical presentations in small vessel disease include seizures, headache, and cognitive decline but also ON or TM. In children with small vessel disease, systemic signs of inflammation are often present, such as an elevated erythrocyte sedimentation rate, Chronic Reactive Protein, or von Willebrand factor, which has been suggested as a clinical marker of disease activity. CSF studies often reveal an elevated opening pressure, pleocytosis, and increased protein levels. MRI may demonstrate white matter lesions reminiscent of ADS and normal magnetic resonance angiography. A definitive diagnosis requires brain biopsy encompassing the meninges, gray matter, and white matter, ideally prior to initiation of glucocorticoid treatment.

Neurosarcoidosis. Granulomatous disorders such as sarcoidosis are often included in the differential diagnosis of ADS. Although selected cases have been described, children rarely present with sole neurologic symptoms such as aseptic meningitis,
Table 4  Selected diseases mimicking an acute demyelinating syndrome with clinical clues and recommended workup that should be tailored according to clinical symptoms and likelihood of an alternative diagnosis

<table>
<thead>
<tr>
<th>Conditions</th>
<th>Clinical clues</th>
<th>Consider the following investigations (in addition to MRI brain/spine)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Systemic lupus erythematosus</td>
<td>Rash, arthralgias, headache, stroke, neuropsychiatric symptoms, cognitive changes, movement disorder, spastic paraplegia</td>
<td>ESR, complement (C3, C4), ANA, dsDNA, rheumatology evaluation</td>
</tr>
<tr>
<td>Behçet disease</td>
<td>Optic neuritis, uveitis, rash, arthralgias, oral/genital ulcers, spastic paraplegia, stroke, cerebrovenous sinus thrombosis</td>
<td>Examination: Oral and genital ulcers; skin pathology test</td>
</tr>
<tr>
<td>Neurosarcoïdosis</td>
<td>Basilar meningitis, uveoparotid fever (uveitis, parotid swelling, facial nerve swelling), cranial neuropathy, raised ICP, seizures, cognitive changes, peripheral neuropathy, spastic paraplegia</td>
<td>Serum/CSF ACE, calcium, ESR, IgG levels, CSF studies (flow cytometry CD4:CD8), CXR + high-resolution CT, bronchoalveolar lavage</td>
</tr>
<tr>
<td>Isolated or primary angitis of the CNS</td>
<td>Headache, stroke, seizures, encephalopathy, visual abnormalities, cognitive changes</td>
<td>ESR, CRP, von Willebrand factor, MRA, VZV antibodies, CSF studies, conventional angiogram, brain biopsy</td>
</tr>
<tr>
<td>Hemophagocytic lymphohistiocytosis</td>
<td>Fever, seizures, meningismus, motor deficit, affected sibling/consanguinity</td>
<td>CBC, triglycerides, ferritin, bone marrow aspiration, CSF studies, genetic testing</td>
</tr>
<tr>
<td>Immunodeficiency syndromes (e.g., XLP, NK)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Neuroborreliosis</td>
<td>Early: Meningoradiculitis, cranial neuritis (e.g., facial palsy), meningoencephalitis, plexus neuritis multiplex, erythema migrans</td>
<td>Serum antibodies against Borrelia burgdorferi, CSF studies with cell count (lymphocytic), AI IgG and PCR B burgdorferi</td>
</tr>
<tr>
<td>Progressive multifocal leukoencephalopathy</td>
<td>Immunosuppressed individual, hemiparesis, dysphasia, ataxia, cortical visual deficits, cognitive changes, seizures, headaches</td>
<td>CSF PCR for JC virus</td>
</tr>
<tr>
<td>Acute encephalopathies with autoantibodies</td>
<td>Seizures, neuropsychiatric symptoms, orofacial dyssomnias and sleep disturbances, autonomic dysfunction</td>
<td>Serum/CSF NMDA, GABA-A, glycine receptor antibodies, EEG</td>
</tr>
<tr>
<td>Steroid-responsive encephalopathy associated with autoimmune thyroiditis</td>
<td>Encephalopathy, seizures, focal neurologic signs, neuropsychiatric features</td>
<td>Antithyroid peroxidase, antithyroglobulin antibodies</td>
</tr>
<tr>
<td>Acute cerebellitis</td>
<td>Ataxia, headache, brainstem syndromes</td>
<td>CSF studies</td>
</tr>
<tr>
<td>Guillain-Barré syndrome and Bickerstaff brainstem encephalitis</td>
<td>Ascending sensorimotor neuropathy, ataxia, areflexia, extracranial movement abnormalities</td>
<td>CSF studies, GQ1 antibodies, NCS</td>
</tr>
<tr>
<td>Solid tumors (astrocytoma, glioma, oligodendroglioma, ependymoma)</td>
<td>Typically monofocal neurologic deficits, persisting symptoms, pain</td>
<td>CSF analysis including cytology, lesional biopsy, tumor-specific testing</td>
</tr>
<tr>
<td>CNS lymphoma</td>
<td>Headache, ataxia, seizures, hemiparesis</td>
<td>CSF analysis including cytology, lesional biopsy, CSF immunotyping</td>
</tr>
<tr>
<td>Langerhans cell histiocytosis</td>
<td>Abnormalities of the hypothalamic-pituitary axis, behavior changes, seizures, visual deficits, headaches</td>
<td>CSF studies, lesional biopsy, CBC, liver enzymes, immunoglobulin levels, ESR, bone marrow aspiration/Biopsy, BRAF gene mutation</td>
</tr>
<tr>
<td>Neurometabolic diseases (general)</td>
<td>Variable degree of acute or episodic neurologic deficits, neurologic regression</td>
<td>Muscle/skin biopsy, MRS, CSF/serum lactate, aminoacids, NCS, ophthalmology evaluation</td>
</tr>
<tr>
<td>Leber hereditary optic neuropathy</td>
<td>Unilateral or bilateral severe vision loss, abnormal retinal</td>
<td>Mutation in mtDNA m.3460G&gt;A, m.11778G&gt;A, or m.14484T&gt;C in 90%</td>
</tr>
<tr>
<td>Kearns-Sayre syndrome, Leigh syndrome, POLG-related disorders</td>
<td>Kearns-Sayre syndrome: Extracranial movement abnormalities, pigmented retinopathy, cardiac conduction abnormalities, myopathy, ataxia</td>
<td>Mitochondrial genetic testing, POLG mutation</td>
</tr>
<tr>
<td>Leukoencephalopathy with brainstem and spinal cord involvement and lactate elevation</td>
<td>Leig syndrome: Psychomotor regression, failure to thrive, hypotonia, ataxia</td>
<td>Mitochondrial genetic testing, POLG mutation</td>
</tr>
<tr>
<td>Acute necrotizing encephalopathy</td>
<td>Progressive spasticity, ataxia and dorsal column dysfunction</td>
<td>DARS2 mutation</td>
</tr>
<tr>
<td>Acute necrotizing encephalopathy</td>
<td>Acute encephalopathy, seizures</td>
<td>Liver enzymes, CSF studies, RANBP2 mutation</td>
</tr>
<tr>
<td>Biotin responsive basal ganglia syndrome</td>
<td>Acute encephalopathy with dystonia, seizures</td>
<td>SLC19A3 mutation</td>
</tr>
<tr>
<td>Migraine</td>
<td>Recurrent headache meeting international headache classification</td>
<td>Clinical history, exclusion of other diseases</td>
</tr>
</tbody>
</table>

Abbreviations: ACE = angiotensin-converting enzyme; AI = antibody index; ANA = anti-nuclear antibody; CBC = complete blood count; CRP = C-reactive protein; CXR = chest X-ray; dsDNA = double-stranded DNA; ESR = erythrocyte sedimentation rate; GABA = γ-aminobutyric acid; ICP = intracranial pressure; IgG = immunoglobulin G; MRA = magnetic resonance angiography; MRS = magnetic resonance spectroscopy; NCS = nerve conduction study; NK = natural killer; POLG = polymerase γ; VZV = varicella-zoster virus; XLP = X-linked lymphoproliferative syndrome.
cranial neuropathy, or space-occupying lesions. Most children have systemic symptoms at initial presentation such as arthritis or swelling of the parotid gland. In CSF, elevated opening pressure, lymphocytosis, and elevated protein are common. Brain MRI can demonstrate granulomata in the parenchyma, frequently enhancing after gadolinium administration and periventricular white matter lesions. A lesional biopsy may be required to establish the diagnosis.
Genetic defects affecting the immune response. Hemophagocytic lymphohistiocytosis (HLH) represents a group of inherited immunodeficiencies, including familial HLH, Griscelli syndrome type 2, and X-linked lymphoproliferative syndrome, which are caused by defects in natural killer cells and CD8 T-cell cytotoxicity. Secondary forms are found in combination with infections, cancer, and other autoimmune disorders.23 Primary HLH remains asymptomatic until a viral infection, such as Epstein-Barr virus, triggers the activation of CD8+ T lymphocytes and macrophages. Initial symptoms can be restricted to the CNS and consist of seizures, impaired consciousness, meningoencephalitis, and focal motor deficits.24 Biological markers indicating HLH are pancytopenia, low fibrinogen levels, and elevated liver enzyme and ferritin levels. Signs of hemophagocytosis are detectable on CSF/blood/bone marrow smears in two-thirds of children. CSF results can be abnormal in half of all children demonstrating high protein and increased cell count.25 Cerebral MRI can demonstrate symmetric white matter lesions sometimes difficult to differentiate from ADEM in addition to signal hypointensity on T1 sequences (figure, C, table 4).

Infections. Neuroborreliosis. In general, children with Lyme neuroborreliosis present with unilateral or bilateral 7th nerve palsy or meningitis. Most cases occur in the late spring and summer and in the 5- to 14-year age group with neurologic involvement in 5%-15% of children.26 Diagnosis relies on the presence of a CSF pleocytosis and an intrathecal synthesis of immunoglobulin G (IgG) and immunoglobulin M antibodies against the pathogen. Infections with Borrelia burgdorferi affecting the neuroaxis can present as ON or TM with or without radiologic evidence of LETM (figure, H) or CNS vasculitis with white matter lesions and focal neurologic signs.27

Progressive multifocal leukoencephalopathy (PML). JC virus is a ubiquitous polyomavirus that causes PML, a frequently fatal demyelinating disease, in particular in immunocompromised patients. Presentation can be insidious with cognitive decline or more rapid with seizures and gait abnormalities. MRI changes show T2-hyperintense lesions in the frontoparietal white matter. Contrast enhancement is unusual. Diagnosis is confirmed on finding JC virus DNA in the CSF.28

Autoantibody-mediated diseases associated with white matter changes. Several antibodies against neuronal surface antigens have been detected in recent years. Although they commonly present with encephalitis, psychiatric symptoms, refractory seizures, or movement disorders, some of them have been shown to be associated with white matter lesions on MRI. Glycine receptor antibodies usually observed in stiff-person syndrome have been found in children and adults with ON.29 CSF antibodies to the inhibitory γ-aminobutyric acid A receptor have been discovered in children with encephalitis and white matter changes.30 Titulaer et al.31 recently reported adults with NMDA receptor encephalitis who subsequently developed demyelinating syndromes associated with MOG or AQP4 antibodies.

Steroid-responsive encephalopathy is characterized by the combination of anti-thyroidperoxidase antibodies with symptoms such as acute encephalitis with seizures, focal neurologic signs, and neuropsychiatric symptoms. Brain MRI lesions can involve the white and gray matter in particular of the subcortical regions and brainstem, often in a symmetrical fashion, which are distinct from MS or ADEM in most cases.32

Acute inflammatory episodes with demyelination other than ADS. Acute cerebellitis is defined by symptoms and MRI changes related primarily to the cerebellum and signs of inflammation in the CSF such as an elevated CSF cell count. Acute cerebellitis can lead to severe brainstem dysfunction. In less acute forms, unilateral or bilateral signal changes of the cerebellar cortex around the dentate nucleus or white matter are observed (figure, G). In children in whom the MRI additionally demonstrates supratentorial, brainstem, or spinal lesions, an inflammatory process such as ADEM should be considered.

Miller Fisher syndrome, Guillain-Barré syndrome (GBS), and Bickerstaff brainstem encephalitis (BBE) have been considered part of a clinical spectrum with a range of symptoms involving the CNS and peripheral nervous system. BBE reflecting the severe end of the spectrum is characterized by encephalopathy and signs of brainstem dysfunction in addition to hyporeflexia or areflexia.33 CSF shows a normal cell count but an elevated protein. Increased titers of anti-GQ1b-IgG antibodies can be used in support of diagnosis. White matter lesions affecting the brainstem or other regions have been reported in up to 1/3 of patients with GBS and BBE (figure, F). Timely diagnosis is important because of the severity and usually prompt response to immunotherapy.34

TUMORS Solid tumors of the CNS. Tumors such as oligodendroglioma, ependymoma, low-grade astrocytoma, and high-grade glioma may initially lead to the incorrect diagnosis of ADS. Important signs are a gradual development of clinical symptoms, the absence of multiple lesions on initial MRI, and lesions that also involve cortical areas extending to other anatomical areas. Disseminated astrocytoma can have spatial dissemination. CSF cell count can be elevated in addition to a high protein. It is recommended to always include a cytospin CSF analysis in the workup of a child with an unusual clinical presentation and inconclusive MRI findings.
Primary CNS lymphoma (PCNSL). PCNSL is often included in the differential diagnosis of an acute demyelinating event. PCNSL is an exceedingly rare but challenging diagnosis in children. Localization of PCNSL is often in the cerebral hemispheres but can be at any anatomical site including the leptomeninges. Therefore, symptoms at presentation are highly variable, including headache, visual problems, ataxia, seizures, and hemiparesis. MRI often reveals lesions in the cerebral hemispheres, basal ganglia, cerebellum, or brainstem, occasionally with meningeal enhancement. Diagnosis is confirmed by brain biopsy or immunophenotypic analyses of CSF. The majority of children reported in the literature have a CNS non-Hodgkin lymphoma.

Langerhans cell histiocytosis (LCH). LCH is a multisystem disease and neurologic symptoms are primarily related to solitary or multifocal lesions in meninges and other areas of the CNS including the brainstem and cerebellum. In the absence of a clinical history of LCH, an isolated CNS lesion presents a diagnostic challenge. MRI findings in these patients include symmetric hyperintense signals in the cerebellar gray and white matter, cerebellar atrophy, T2-weighted hyperintense changes in the pons, and T1-weighted hyperintensity of the globus pallidum.

**NEUROMETABOLIC DISEASES** A range of neurometabolic conditions are associated with white matter changes in the CNS, many of which present acutely in the context of a nonspecific illness with encephalopathy and neurologic symptoms, reminiscent of ADEM. MRI changes that are symmetrical and involve the striatum should alert the clinician to investigate in particular treatable diseases such as biotin-responsive basal ganglia disease (table 4).

Mitochondrial respiratory chain defects can present with symptoms and MRI findings indistinguishable from ADS. An evaluation of lactate content and of lactate/pyruvate ratio in blood and CSF, spectroscopy focusing on white matter lesions, and a direct study of respiratory chain activity in fibroblasts or muscle biopsy may be required in addition to genetic testing to establish the diagnosis. Leber hereditary optic neuropathy, caused by point mutations in mtDNA, deserves special consideration because of its associations with MS and ON. Other important mitochondrial disorders include Leigh syndrome, Kearns-Sayre syndrome, or leukoencephalopathy with brainstem and spinal cord involvement. The latter disease can present in a previously healthy child with pyramidal and cerebellar dysfunction. MRI abnormalities include spotty, periventricular white matter changes, as well as cerebellar white matter, brainstem, and spinal cord imaging similar to LETM. Further episodes can occur and respond to steroids.

Two other diseases affecting cellular energy metabolism are biotin-responsive basal ganglia disease (BBGD), which is due to a SLC19A3 mutation, and acute necrotizing encephalopathy (ANE). In the latter, a heterozygous mutation in RANBP2 is found in familial and recurrent ANE, a rapidly progressive encephalopathy that typically occurs in previously healthy children between 6 and 18 months of age. Typically, there is a prodrome of a febrile illness prior to the acute phase of encephalopathy with loss of consciousness and seizures. Mortality and neurologic sequelae are common in particular in cases not treated promptly with steroids (figure, 1). Liver transaminases are mildly elevated with increased CSF protein in the absence of pleocytosis. MRI demonstrates symmetrical involvement of the thalamus, with additional regions affected including the periventricular white matter, putamen, internal capsule, brainstem, and cerebellum.

BBGD should be considered in the context of an ADEM-like presentation and symmetrical basal ganglia involvement and treated immediately with biotin and thiamin.

Leukodystrophies such as Alexander disease, X-linked adrenoleukodystrophy, or metachromatic leukodystrophy can present at various ages with atypical presentations and MRIs, but due to their progressive nature and MRI patterns, should be differentiated easily from acute demyelinating syndromes.

**MIGRAINE** The detection of nonspecific incidental white matter changes in children with headache, or more specifically migraine, is now well-established. Different series demonstrate a similar rate of up to 6% of white matter hyperintensities with a trend for a higher prevalence in migraine with aura. Lesions are only supratentorial. The distinction from CIS is usually on clinical grounds and on differences in signal intensity, localization, and absence of gadolinium enhancement.

**DISCUSSION** Precise and detailed description of the clinical picture, along with MRI and CSF studies, allow the characterization of the different ADS in the majority of children. Nevertheless, a proportion of cases are due to a broad spectrum of other inflammatory disorders of the white matter, primary CNS tumors, and neurometabolic diseases. The clinician therefore has to be aware of the different forms of ADS, the risk factors for a chronic-relapsing course, and features that indicate an alternative diagnosis.

**AUTHOR CONTRIBUTIONS**

Kevin Rostasy: concept, outline, and writing of the manuscript. Barbara Bajer-Kornek: writing of the manuscript. Sunita Venkateswaran: writing of the manuscript and tables. Cheryl Hemingway: writing of the
REFERENCES


Differential diagnosis and evaluation in pediatric inflammatory demyelinating disorders
Kevin Rostasy, Barbara Bajer-Kornek, Sunita Venkateswaran, et al.
Neurology 2016;87;S28-S37
DOI 10.1212/WNL.0000000000002878

This information is current as of August 29, 2016

Updated Information & Services
including high resolution figures, can be found at:
http://www.neurology.org/content/87/9_Supplement_2/S28.full.html

References
This article cites 40 articles, 5 of which you can access for free at:
http://www.neurology.org/content/87/9_Supplement_2/S28.full.html#
ref-list-1

Subspecialty Collections
This article, along with others on similar topics, appears in the following collection(s):
Acute disseminated encephalomyelitis
http://www.neurology.org/cgi/collection/acute_disseminated_encephalomyelitis
Multiple sclerosis
http://www.neurology.org/cgi/collection/multiple_sclerosis

Permissions & Licensing
Information about reproducing this article in parts (figures, tables) or in its entirety can be found online at:
http://www.neurology.org/misc/about.xhtml#permissions

Reprints
Information about ordering reprints can be found online:
http://www.neurology.org/misc/addir.xhtml#reprintsus