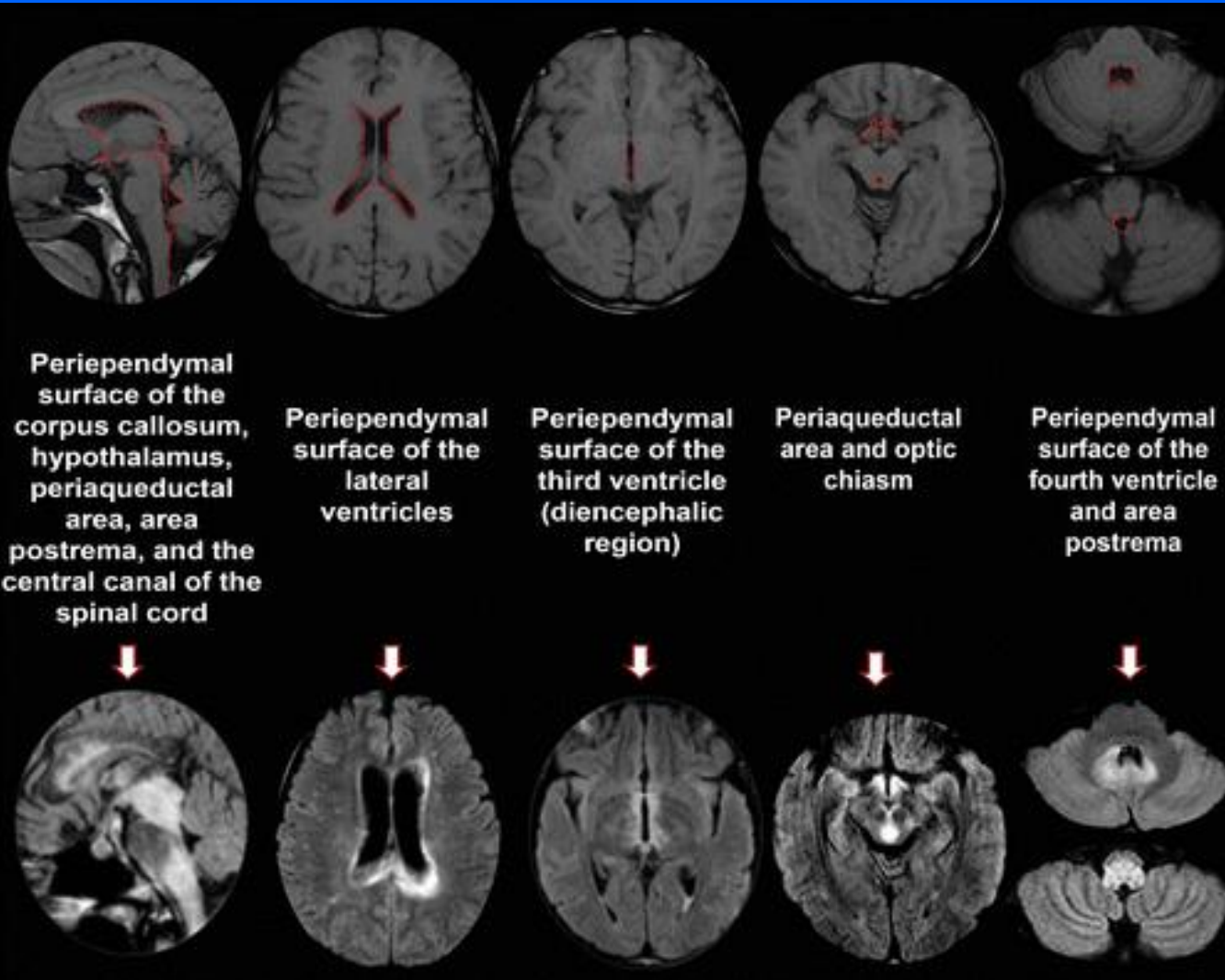
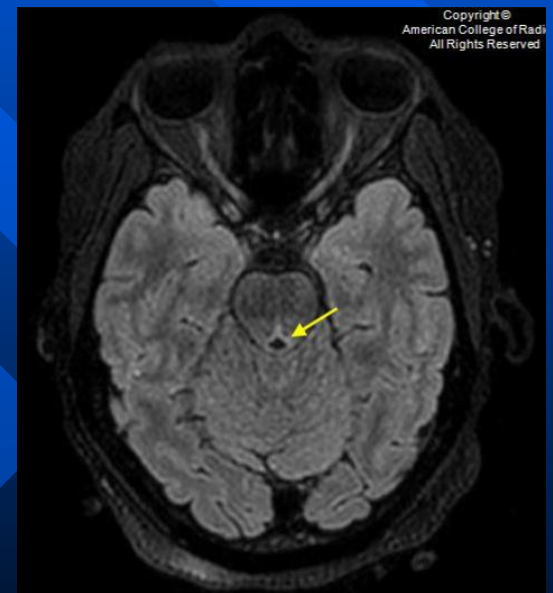
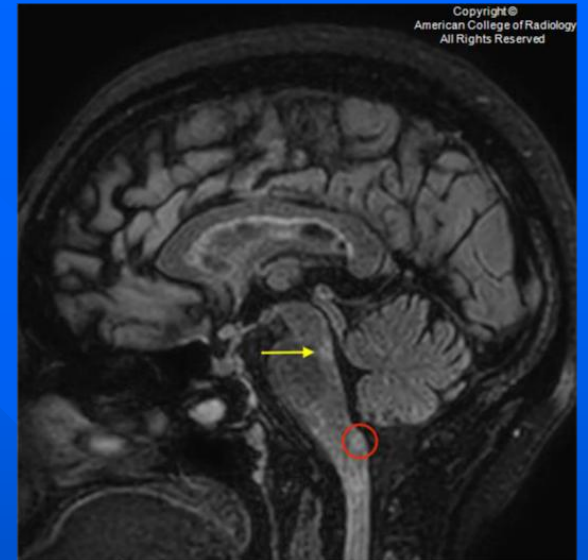
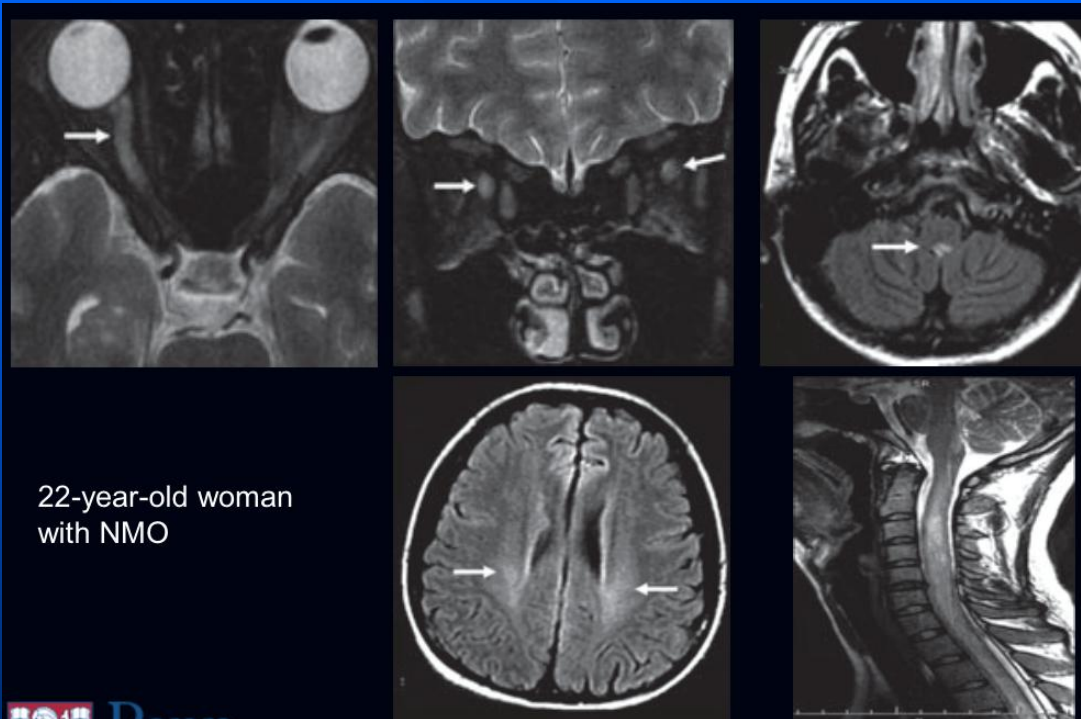


Neuromyelitis optica spectrum disorder

- Closely related severe demyelinating diseases caused by an autoantibody to the aquaporin-4 water channel.
- Classic presentation of NMO is with the triad of optic neuritis, longitudinally extensive myelitis, and positive **anti-AQP4 antibody**
 - although a far wider range of manifestations are now recognized as part of NMOSD
- Typically found in patients older than those with multiple sclerosis (MS), average age of presentation of 41.
- Even **stronger female** predilection (F:M 6.5:1).
- It is found more frequently in patients of **Asian, Indian, and African descent**





Brain

- Lesions which mirror the distribution of aquaporin 4 in the brain,
 - Particularly found in the periependymal regions abutting the ventricles
 - Next to the lateral ventricles, dorsal medulla, and periaqueductal gray
- Periventricular (hemispheric) confluent smooth sessile white matter involvement **usually no Dawson's fingers**)
- Hypothalamus/medial thalamus
- Dorsal pons/medulla
- Corpus callosum
 - multiple callosal lesions with heterogeneous signal leading to a marbled pattern
 - the splenium may be diffusely involved and expanded
- Deep (or less frequently subcortical) punctate white matter lesions (which may appear like those seen in MS)
- Larger >3 cm diameter hemispheric white matter lesions

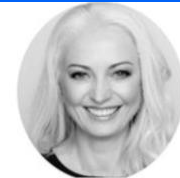


<i>Pathophysiology</i>	- Chronic CNS inflammation with antibodies against AQP4 channels
<i>Epidemiology</i>	- Prevalence: 0.5 – 4.4/100 000 - Ratio female/male: up to 9:1
<i>Clinical features</i>	- Optic neuritis, acute myelitis, area postrema syndrome, acute brain stem, or diencephalic syndrome
<i>Diagnosis</i>	- Combination of clinical and/or MRI in patients with and without AQP4 antibodies - Differential diagnoses: MOG-ab disease, other causes of LETM or optic neuritis (e.g., MS, ADEM, sarcoidosis)
<i>Treatment</i>	- Acute relapses: high-dose intravenous steroids and plasmapheresis - B-cell depleting (Rituximab) or other immunosuppressive medication
<i>Prognosis</i>	- Severely disabling CNS attacks may occur with incomplete recovery - The rise of modern treatment strategies may improve outcomes
<i>When to test for AQP4 antibodies?</i>	- In patients with LETM or extensive optic neuritis - In patients with atypical brain or brainstem lesions

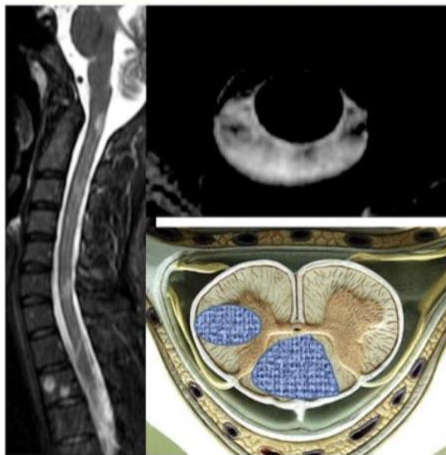
- Long lesions involving ≥ 3 vertebral segments are more characteristic of NMOSD,
- Short lesions are more characteristic of MS

Biomarkers and Imaging Findings in Myelitis

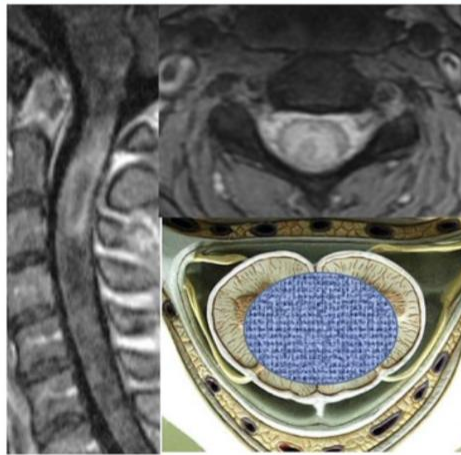
Majda M Thurnher, MD, EDiNR



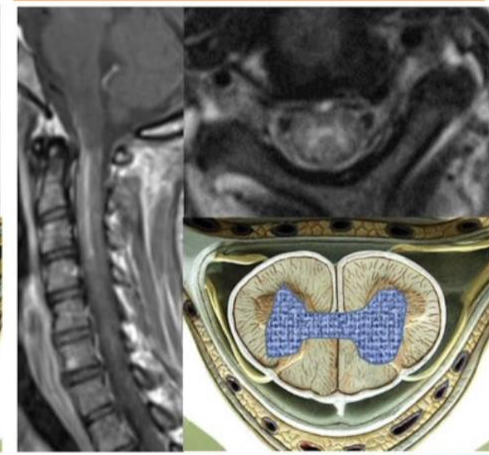
Multiple Sclerosis
(MS)



Neuromyelitis Optica
Spectrum Disorder
(NMOSD)



Myelin Oligodendrocyte
Glycoprotein-associated disease
(MOGAD)



#ASNR23

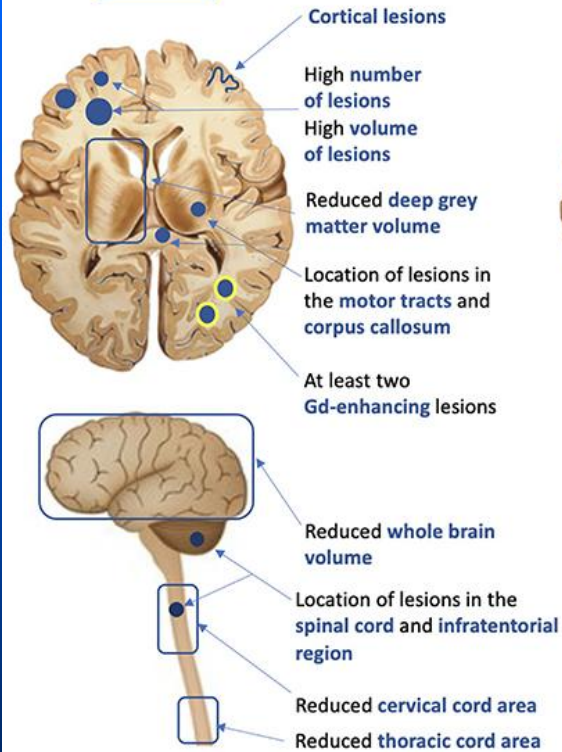
ASNR23

APRIL 29 - MAY 3
CHICAGO

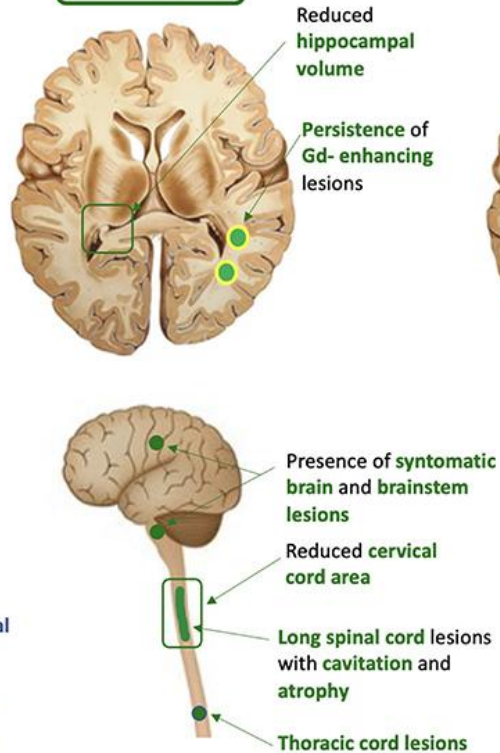
TRANSFORMING THE FUTURE
OF NEURORADIOLOGY



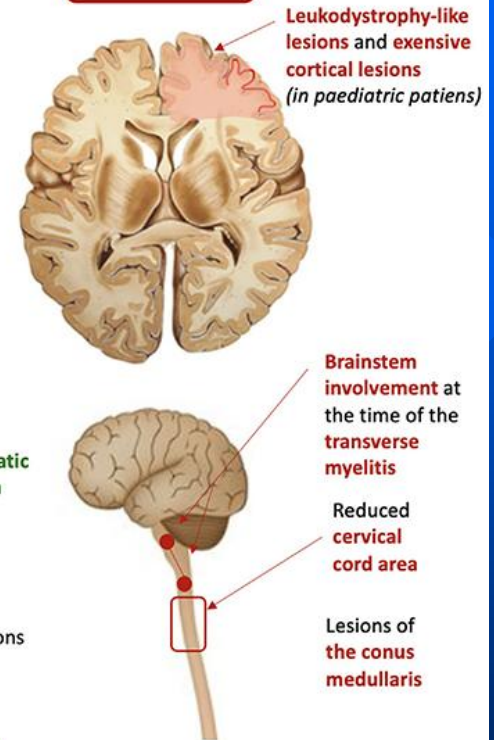
MS

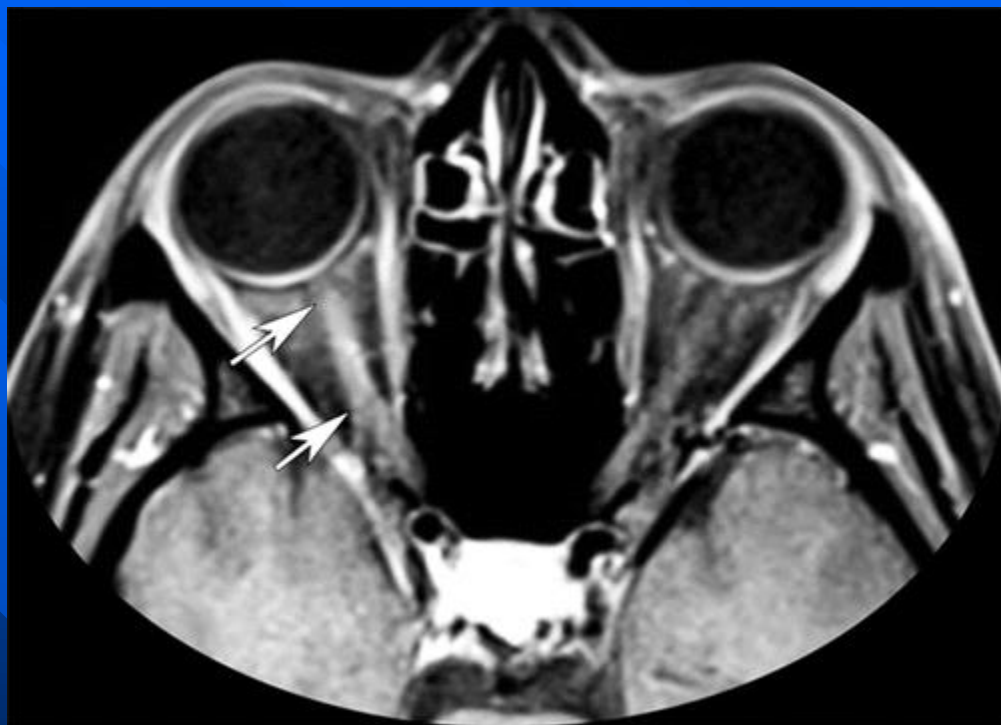


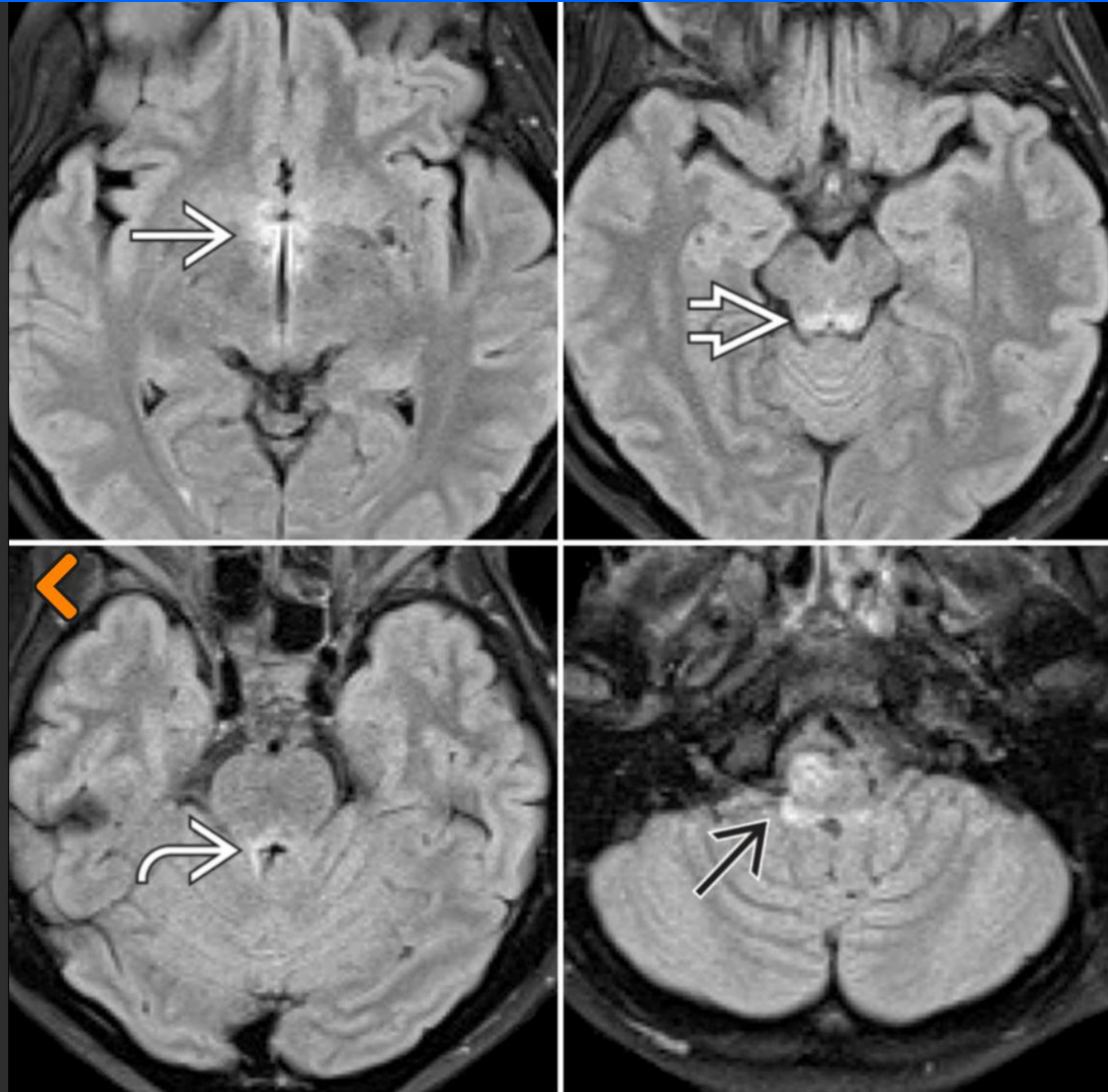
NMOSD



MOGAD







[View Full Screen Image](#)

Axial FLAIR MR images show the typical distribution pattern of brain lesions in NMOSD, including the peripendymal 3rd ventricle →, adjacent to the aqueduct ⇨, the peripendymal 4th ventricle, ↪ and the medulla ⇩.

