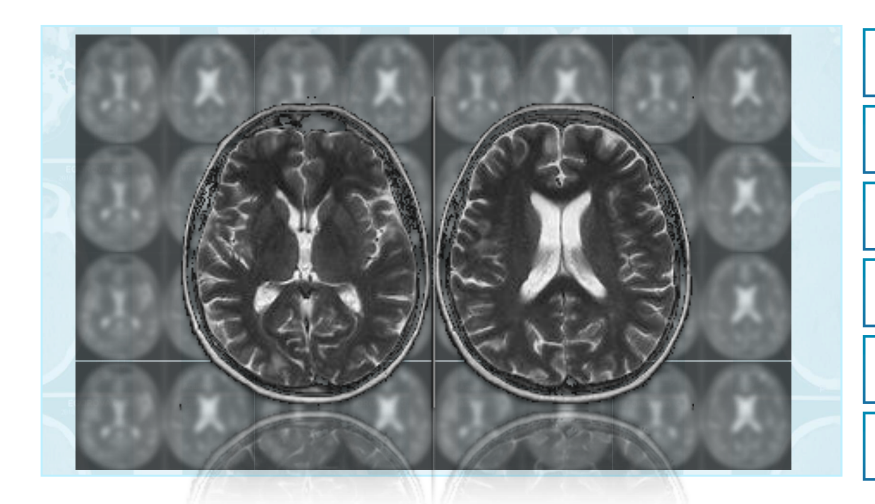




Outline slide



Introducing ARIA

Pathophysiology

Deeper focus on ARIA

Clinical manifestation of ARIA

Diagnosis of ARIA

Management of ARIA





What is ARIA?

- ARIA is a consequence of the presence of amyloid in cerebral blood vessel walls (cerebral amyloid angiopathy [CAA]).¹
 CAA can cause spontaneous ARIA in patients with AD and the risk of ARIA is increased with monoclonal antibodies that remove amyloid plaques¹
- Studies have suggested that ARIA-E and ARIA-H may be caused by disruption of vessels with CAA and the risk is increased by the clearance of Aβ from cerebral vessels, but other mechanisms are also hypothesized²
- An Alzheimer's Association workgroup defined the term "amyloid-related imaging abnormalities" or "ARIA," in AD based on MRI findings which are subdivided into ARIA-E or ARIA-H¹
 - ARIA-E: parenchymal vasogenic edema or sulcal effusions detected on FLAIR sequences³
 - ARIA-H: microhemorrhages, superficial hemosiderin deposition (superficial siderosis) detected on T2*GRE sequences³
- Most cases of ARIA in patients treated with monoclonal antibodies that remove amyloid plaque are asymptomatic; however, ARIA-E may have concurrent symptoms such as headache, confusion, dizziness, and nausea; less likely, gait disturbances, visual impairment, and rarely seizures. ARIA can be serious, and life-threatening and may require intervention beyond withholding treatment to address symptoms.

ARIA, amyloid-related imaging abnormalities; ARIA-E: ARIA-edema/effusion; ARIA-H: ARIA-hemosiderin/hemorrhage; FLAIR, fluid-attenuated inversion recovery; GRE, gradient-recalled echo

1. Sperling RA, et al. Alzheimers Dement. 2011;7:367–385; 2. Sperling RA, et al. Lancet Neurol. 2012;11:241–249; 3. Barakos J, et al. J Prev Alzheimers Dis. 2022;9(2):211–220; 4. Salloway S, et al. JAMA Neurol. 2022;79(1):13–21; 5. Cummings J, et al. J Prev Alzheimers Dis 2022:9:221–230

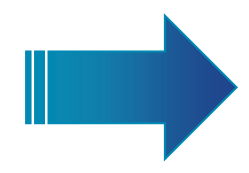


Emerging therapies aiming to remove amyloid beta (A β)

Monoclonal antibodies that remove amyloid



Strategies to target and remove amyloid are based on our understanding that interfering with the underlying pathophysiologic mechanisms of the disease process could slow disease progression, but need to be initiated early in the course of disease given these changes begin in the early stages of disease¹



Amyloid-related imaging abnormalities

Interfering/removing the amyloid deposition in the brain that has built up over years can impact the vessel vasculature in the brain which can result in signal changes identifiable on MRI: "amyloid-related imaging abnormalities or ARIA"²

ARIA is a known adverse reaction of monoclonal antibodies that remove amyloid plaque for AD

Aβ, amyloid beta; ARIA: Amyloid-related imaging abnormalities; AD: Alzheimer's Disease
1. Bateman RJ, et al. N Engl J Med 2012;367:795–804; 2. Sperling RA, et al. Alzheimers Dement. 2011;7:367–385



Neuroimaging: ARIA-E and ARIA-H



Interstitial vasogenic edema or sulcal effusion that manifests as parenchymal or sulcal hyperintensities



Microhemorrhages (mH) are observed as <1cm hypointense hemosiderin deposition in the parenchyma Superficial siderosis is observed as linear hypointense hemosiderin deposition in the leptomeningeal/subpial space

Primary MRI features

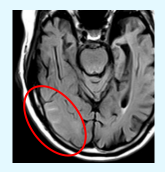
ARIA-E

Edema



FLAIR hyperintense; parenchymal edema in left occipital-parietal lobe^a

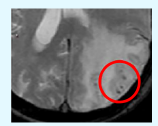
Effusion



FLAIR hyperintense; increased MRI signal in sulci within right temporal-occipital lobe^a

ARIA-H

Microhemorrhage



Punctate foci of signal void on T2*GRE in an area of parenchymal edema^a

Superficial siderosis



New right temporal superficial siderosis on axial T2*GRE imagingb

Intracerebral hemorrhage (also termed macrohemorrhage):

Rare lobar intracerebral hemorrhage occurs spontaneously in AD and with monoclonal antibodies that remove amyloid, related to underlying CAA²

Figures reproduced from ^aBarakos et al (2022); ^bCogswell et al (2022).

ARIA, amyloid-related imaging abnormalities; ARIA-E, ARIA-E, ARIA-H: ARIA-hemosiderin/hemorrhage; CAA, cerebral amyloid angiopathy; FLAIR, fluid-attenuated inversion recovery; GRE, gradient recalled echo; mH, microhemorrhage; MRI, magnetic resonance imaging; SWI, susceptibility weighted imaging.

1. Barakos J, et al. J Prev Alzheimers Dis. 2022;9(2):211–220; 2. Cogswell PM, et al. AJNR Am J Neuroradiol. 2022;43(9):E19–E35





ARIA-E and ARIA-H

ARIA is an umbrella term used to describe two types of imaging abnormalities¹

	ARIA-E ^{1,2}	ARIA-H ^{1,2}
PRIMARY DIAGNOSTIC IMAGING SEQUENCE	FLAIR	T2* GRE
NATURE OF LEAKAGE PRODUCTS	Proteinaceous fluids	Blood-degradation products
LOCATION OF INCREASED VASCULAR PERMEABILITY	Parenchyma: vasogenic edema Leptomeninges: sulcal effusions (i.e., exudates)	Parenchyma: microhemorrhages (typically defined as <10 mm) and intracerebral hemorrhage (≥10 mm) Leptomeninges: superficial hemosiderin deposits (superficial siderosis)
EVALUATION OF SEVERITY	MRI severity scales ³ and assessment of symptoms	The number of microhemorrhages and hemosiderin deposits on MRI and assessment of symptoms
IMAGE	ARIA-E seen on FLAIR images demonstrating increased signal in multiple regions of the right hemisphere, affecting both gray and white matter ⁴	ARIA-H seen on T2* GRE MRI. MRI reveals several microhemorrhages (<10 mm; red circle) ⁴

ARIA, amyloid-related imaging abnormalities; ARIA-E: ARIA-edema/effusion; ARIA-H: ARIA-hemosiderin/hemorrhage; FLAIR, fluid-attenuated inversion recovery; GRE, gradient-recalled echo; MRI, magnetic resonance imaging.

1. Sperling RA, et al. Alzheimers Dement. 2011;7:367–85; 2. Barakos J, et al. AJNR Am J Neuroradiol. 2013;34:1958–965; 3. Barkhof F, et al. AJNR Am J Neuroradiol. 2013;34:1550–1555; 4. Cogswell PM, et al. AJNR Am J Neuroradiol. 2022;43(9):E19–E35







Hypothesized pathophysiology of ARIA

ARIA is a consequence of the presence of amyloid in cerebral blood vessel walls (cerebral amyloid angiopathy [CAA]), which can cause spontaneous ARIA in patients with AD.1 The increased occurrence of ARIA-E seen with treatments that remove amyloid plaques is thought to be due to the removal of vascular amyloid and disruption of amyloid in blood vessel walls. Other mechanisms are also hypothesized.

Aggregation of toxic **Aβ species** in the brain (amyloid plaques) and blood vessels (CAA) contributes to Alzheimer's disease pathogenesis³

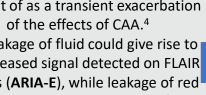


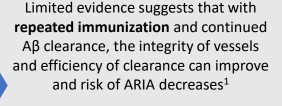
After the introduction of monoclonal antibodies that remove amyloid plaque, vascular amyloid deposits begin to clear leading to increased vascular permeability 1

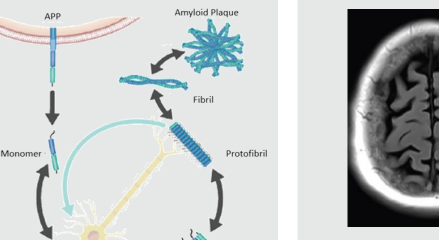


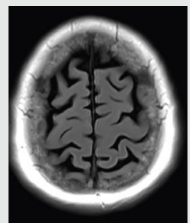
This loss of vascular integrity may be thought of as a transient exacerbation of the effects of CAA.4

The leakage of fluid could give rise to an increased signal detected on FLAIR images (ARIA-E), while leakage of red cells would result in ARIA-H1





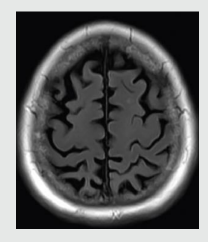




Example of Baseline MRI³



Example of ARIA-E post treatment³



Example of ARIA-E post treatment follow-up3

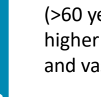
MRI images from Cogswell et al (2022);³ figure adapted from Hampel et al. (2021)⁴

AB, Amyloid beta; ARIA: Amyloid-related imaging abnormalities; ARIA-E, ARIA-edema/effusion; ARIA-H, ARIA-hemosiderin/hemorrhage 1. Sperling RA, et al. Alzheimers Dement. 2011;7:367–385; 2. Barakos, J et al. J Prev Alzheimer's Dis 2022; 9(2):211–220; 3. Cogswell, PM et al. AJNR Am J Neuroradiol. 2022;43(9):E19–E35; 4. Hampel H, et al. Nature. 2021;26:5481–5503





Increased risk of ARIA-E and ARIA-H in carriers of APOE E4



 APOF ε4 carriers (>60 years of age) have higher parenchymal and vascular Aß load^{1,2}

Therefore, when exposed to anti-Aß monoclonal antibodies, they would experience a larger antibodymediated shift in Aβ compared with non-carriers³



The presence of APOE ε4 alleles is one of the most robust known risk factors for ARIA-F³ and a proposed risk factor for ARIA-H⁴ occurrence in trials of monoclonal antibodies that remove amyloid plaque in patients with AD



• APOF ε4 carrier status is also a risk factor for spontaneously occurring ARIA-like events in microhemorrhage in the general population,⁵ microhemorrhage among patients in memory clinics, 6 and CAA-ri⁷

These findings support the hypothesis that vascular amyloid plays a key role in the induction of ARIA-E and ARIA-H^{1,2}

Aβ, amyloid beta; AD, Alzheimer's disease; APOE ε4, apolipoprotein Ε ε4; ARIA, amyloid-related imaging abnormalities; ARIA-E: ARIA-edema/effusion; ARIA-H: ARIA-hemosiderin/hemorrhage; CAA-ri, cerebral amyloid angiopathy-related imflammation. 1. Caselli RJ, et al. Neurosci Lett. 2010;473:168–171; 2. Cogswell, PM et al. AJNR Am J Neuroradiol. 2022;43(9):E19–E35; 3. Ketter N, et al. J Alzheimers Dis. 2017;57:557–573. 4. Arrighi HM, et al. J Neurol Neurosurg Psychiatry. 2016;87:106–112; 5. Poels MM, et al. al. Stroke. 2011;42:656–661; 6. Goos JD, et al. Neurology. 2010;74:1954–1960; 7. Kinnecom C, et al. Neurology. 2007;68:1411-1416;





Cerebral Amyloid Angiopathy (CAA) presentation and Cerebral Amyloid Angiopathy-related inflammation (CAA-ri)

What is CAA?



CAA is a type of cerebrovascular disorder characterized by the accumulation of Aβ peptide within the leptomeninges and small/medium-sized cerebral blood vessels in patients with or without AD symptoms¹

CAA presentation



Aβ deposition results in fragile vessels that may present with microhemorrhages, superficial hemosiderosis, or intracerebral hemorrhage (macrohemorrhage)¹

CAA-ri



CAA-ri is a rare and potentially life-threatening autoimmune response to vascular amyloid complication of CAA.² It can be a treatment-reversible disease, responsive to immunosuppressive therapies³

Aβ, amyloid-β; AD, Alzheimer's disease; CAA, cerebral amyloid angiopathy; CAA-ri, CAA-related inflammation.

1. Kuhn J, Sharman T. Cerebral Amyloid Angiopathy. 2022 Jun 6. In: StatPearls [Internet]. Treasure Island (FL): StatPearls Publishing; 2022 Jan; 2. Grasso, D et al. Radiol Case Rep. 2021 Sep:;16(9):2514-2521; 3. Antolini, L et al. Neurology 2021;97:e1809–e1822



Commonalities in pathophysiology between CAA-ri and ARIA

While ARIA and CAA-ri are separate entities, they share a number of similarities:

Risk factors



Increased number of microbleeds and APOE ε4 genotype

Location



CAA develops to a greater extent in cortical and leptomeningeal vessels (the locations where ARIA occurs)

Syndrome resemblance

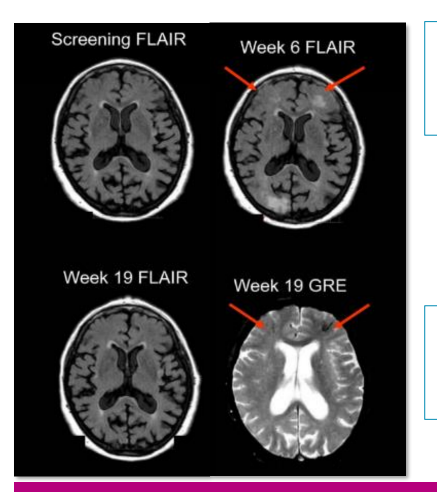


Infiltration of inflammatory cells (microglia, T cells, and Aβ-containing multinucleated cells) in CAA-ri suggests possible spontaneous anti-Aβ immunization

Aβ, amyloid-β; APOE ε4, apolipoprotein E ε4; ARIA, amyloid-related imaging abnormalities; CAA, cerebral amyloid angiopathy; CAA-ri, CAA-related inflammation. Greenberg SM, et al. Nat Rev Neurol. 2020;16(1):30–42

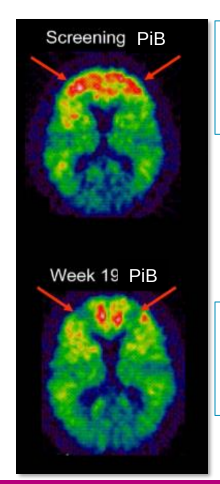


Relationship between amyloid removal with monoclonal antibodies and ARIA-E and ARIA-H



At Week 6, FLAIR MRI reveals bifrontal parenchymal hyperintensity (ARIA-E), which resolves by Week 19

At Week 19, T2*GRE sequence reveals the development of bifrontal microhemorrhages (ARIA-H)



Baseline PiB retention consistent with high fibrillar burden

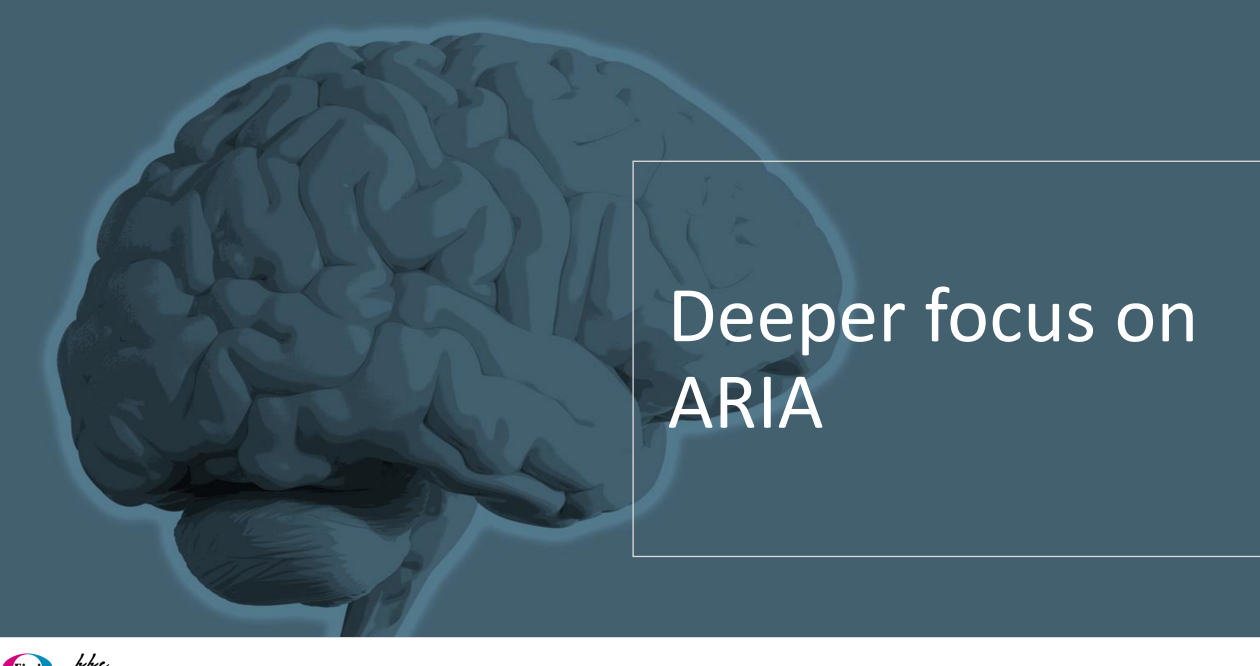
Week 19 PiB uptake is reduced representing clearance of fibrillar amyloid from plaque and cerebral vessels

Reduced PiB retention is temporally and regionally associated with ARIA-E and ARIA-H

ARIA-E: ARIA-edema/effusion; ARIA-H: ARIA-hemosiderin/hemorrhage; FLAIR, fluid-attenuated inversion recovery; GRE, gradient-recalled echo; MRI, magnetic resonance imaging; PiB-PET, Pittsburgh compound B-positron emission tomography Sperling RA, et al. Lancet Neurol 2012;11:241-249









ARIA-E

Parenchymal signal abnormalities (ARIA-E edema)

- Imaging features of **ARIA-E edema** are thought to reflect leakage of intravascular fluid and proteins into the parenchymal interstitial compartment1
- Parenchymal signal abnormalities can be quite subtle in a single region, multifocal, or nearly pan-hemispheric²



Figure from Barakos et al (2022)4

Sulcal FLAIR hyperintensities (ARIA-E effusion)

- The imaging features of ARIA-E effusion are thought to reflect leakage or effusion of proteinaceous fluid from meningeal vessels²
- Sulcal FLAIR hyperintensity in the leptomeningeal or sulcal space may be seen in isolation or near gray matter disturbances²



Figure from Barakos et al (2022)4

Additional analyses are required to confirm the prevalence of spontaneous ARIA-E³ In clinical trials, the rate of spontaneous ARIA-E in the placebo arm over 18 months has been found to range between 0.8% and 3.0%⁵⁻⁸

AD, Alzheimer's disease; ARIA, amyloid-related imaging abnormalities; ARIA-E: ARIA-edema/effusion; FLAIR, fluid-attenuated inversion recovery. 1. Barakos J, et al. AJNR Am J Neuroradiol. 2013;34:1958–1965; 2. Sperling RA, et al. Alzheimers Dement. 2011;7:367–385; 3. Carlson C, et al. Alzheimers Dement. 2011;396–401; 4. Barakos J, et al. J Prev Alzheimers Dis. 2022;9(2):211–220;

5. Budd-Haeberlein S, et al. J Prev Alzheimers Dis 2022;9:197-210; 6, van Dvck C, et al. N Eng J Med 2023;388;9-21; 7. Ostrowitzki S, et al. Alzheimers Res Ther, 2017;9(1);95; 8, Vandenberghe R, et al. Alzheimers Res Ther, 2016;8:18.





ARIA-H

Microhemorrhages

- Small deposits of iron in the brain parenchyma in the form of hemosiderin¹
- Typically manifest as new hypointense lesions on T2*GRE MRI sequences (typically defined by a cutoff of <10 mm)²
- Thought to represent residua of a small leakage of blood from a vessel into adjacent tissue¹
- The baseline prevalence of microhemorrhages is estimated to be 15.3%³
 - This prevalence increases with age: ~17% in people aged 60–69 years, ~29% in people aged 70–79 years, and ~36% in people aged 80–97 years³

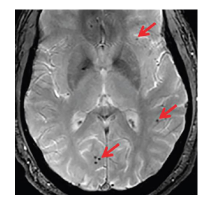


Figure from Cogswell et al (2022) 4

Less commonly, intracerebral hemorrhage (≥10 mm) can also occur²

Superficial Siderosis

- Curvilinear low intensities on T2*GRE MRI sequences that lie adjacent to the surface of the brain¹
- Attributed to the deposition of iron in the form of hemosiderin and is thought to represent residua of leakage of blood from a vessel into the adjacent subarachnoid space or the periadventitial compartment¹
- The baseline prevalence of superficial siderosis is estimated to be 0.21% in those aged 50–69 years and 1.43% in those >69 years old⁵

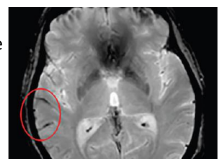


Figure from Cogswell et al (2022) 4

ARIA-H: ARIA-hemosiderin/hemorrhage; GRE, gradient-recalled echo; MRI, magnetic resonance imaging.

1. Sperling RA, et al. Alzheimers Dement. 2011;7:367-385. 2. Barakos J, et al. AJNR Am J Neuroradiol. 2013;34:1958-1965. 3. Poels MM, et al. Stroke. 2010;41:S103-S106 4. Cogswell PM, et al. AJNR Am J Neuroradiol. 2022;43(9):E19–E35; 5. Pichler M, et al. Stroke. 2017;48:3210-3214









Clinical manifestations of ARIA



In most cases, ARIA is asymptomatic. Moreover, most cases occur early in the treatment course and decrease with increased duration of exposure^{1,2}



The most commonly reported symptoms of ARIA-E are transient and nonspecific and include headache, confusion, dizziness, nausea and neuropsychiatric symptoms; less frequent symptoms include fatigue, visual impairment, blurred vision, and gait disturbance^{1,3}



Infrequently, severe symptoms occur (e.g., encephalopathy, focal neurologic symptoms, seizures), requiring hospitalization and specific treatments (e.g., intensive care unit admission, electroencephalography, corticosteroids, antiepileptics). ARIA can be serious and lifethreatening4

1. Filippi M, et al. JAMA Neurol. 2022;79(3):291–304; 2. Sperling RA, et al. Lancet Neurol 2012;11:241–249 3. Salloway S, et al. JAMA Neurol. 2022;79(1):13–21;; 4. Cummings J, et al. J Prev Alzheimers Dis 2022;9:221–230





ARIA experience from clinical trials



ARIA is more common in APOE £4 carriers^{1,2}



Most cases of ARIA-E and ARIA-H are asymptomatic and usually recognized as incidental ARIA during follow-up evaluation on MRI^{1,2}



Most cases of ARIA-E occur early in the treatment course and decrease with increased duration of exposure. 1–3 ARIA-E and ARIA-H may occur concurrently³



Most cases of ARIA-E resolve completely. Depending on severity, treatment may be continued, be interrupted, or discontinued.^{1,4,5,6} Some cases may require specific treatments or even hospitalization.⁶ ARIA-H stabilizes but can remain on subsequent imaging^{3,7}



Re-dosing following resolution is generally associated with a low rate of ARIA recurrence^{4,5}

APOE ε4, a polipoprotein E ε4; ARIA, amyloid-related imaging abnormalities; ARIA-E, ARIA-edema/effusion; ARIA-H: ARIA-hemosiderin/hemorrhage; MRI, magnetic resonance imaging 1. Sperling R, et al. Lancet Neurol 2012;11:241–249; 2. Filippi M, et al. JAMA Neurol. 2022;79(3):291–304; 3. Barakos J, et al. AJNR Am J Neuroradiol. 2013;34(10):1958–1965; 4. Ketter N, et al. J Alzheimers Dis 2017;57:557–573; 5. Ostrowitzki S, et al. Alzheimers Res Ther 2017;9:95; 6. Cummings J, et al. J Prev Alzheimers Dis 2022;9:221–230; 7. Salloway S, et al. JAMA Neurol. 2022;79(1):13–21;



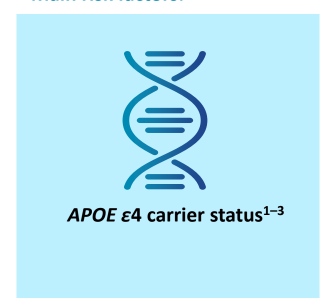






ARIA risk factors

Main risk factors:







APOE, apolipoprotein E; ARIA, amyloid-related imaging abnormalities (includes ARIA-E and ARIA-H); ARIA-e, ARIA-edema/effusion; ARIA-H, ARIA-hemosiderin/hemorrhage;
1. Filippi M, et al. JAMA Neurol. 2022;79(3):291–304; 2. Sperling RA, et al. Alzheimers Dement. 2011;7(4):367–385; 3. Cogswell PM, et al. AJNR Am J Neuroradiol. 2022;43(9):E19–E35



Grading scale for determining radiographic severity of ARIA

ARIA-E, ARIA-H microhemorrhage, and ARIA-H superficial siderosis are each categorized by radiographic severity (mild to severe) based on the following criteria

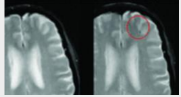
ARIA-E

Sulcal and/or cortical/ subcortical FLAIR hyperintensity

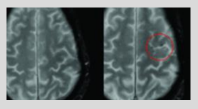
ARIA-H Superficial siderosis

ARIA-H Number of new Microhemorrhages

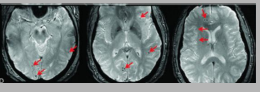
Mild Moderate Severe 1 location 5-10 cm OR 1 more location 1 location <5 cm >1 location each <10 cm > 10 cm Baseline **Posttreatment** Baseline **Posttreatment** Posttreatment Baseline 1 focal area 2 focal areas > 2 focal areas ≤4 5-9 ≥10



Baseline Posttreatment <5 treatment-emergent microhemorrhages



Baseline **Posttreatment** 5 treatment-emergent microhemorrhages



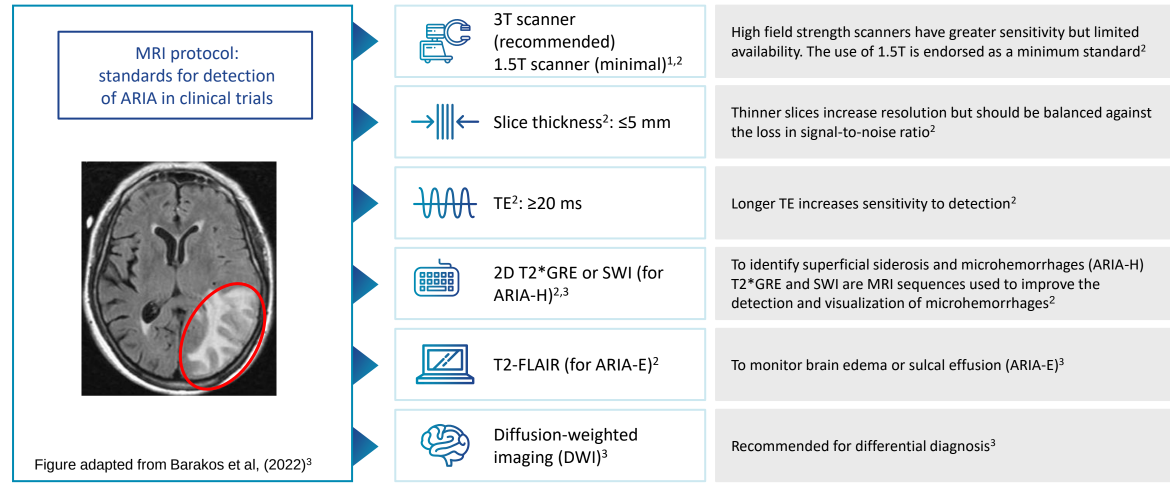
Posttreatment At least 12 treatment-emergent microhemorrhages (arrows)

ARIA, amyloid-related imaging abnormalities; ARIA-E, ARIA-vasogenic edema; ARIA-H, ARIA-microhemorrhages or hemosiderosis Cogswell PM, et al. AJNR Am J Neuroradiol. 2022;43(9):E19-E35

Figure adapted from Cogswell et al (2022)



Recommended MRI protocols for detection of ARIA



ARIA, amyloid-related imaging abnormalities; ARIA-E, ARIA-edema/effusion; ARIA-H, ARIA-hemosiderin/hemorrhage; GRE, gradient recalled echo; MRI, magnetic resonance imaging; T2-FLAIR, T2-weighted fluid attenuated inversion recovery; TE, echo time; SWI, susceptibility weighted imaging.

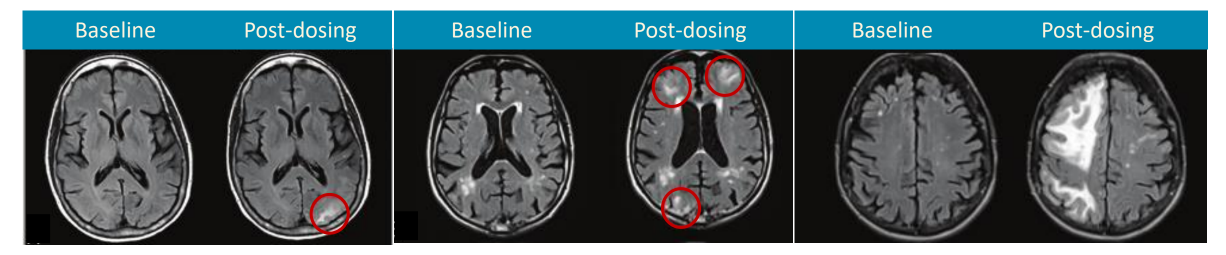
1. Cogswell PM, et al. AJNR Am J Neuroradiol. 2022;43(9):E19–E35; 2. Sperling RA, et al. Alzheimers Dement. 2011;7(4):367–385; 3. Barakos J, et al. J Prev Alzheimers Dis. 2022;9(2):211–220





Detection of ARIA-E, parenchymal edema, and corresponding grading

All figures adapted from Cogswell, PM et. al (2022)



Mild ARIA-E

T2-FLAIR hyperintense signal in the left parietooccipital subcortical white matter with mild local mass effect and sulcal effacement measuring <5 cm the transverse dimension

Moderate ARIA-E

New multifocal, patchy T2-FLAIR hyperintense signal in the bifrontal and right occipital subcortical white matter, each region measuring <5 cm. A single region measuring <5 cm would be classified as mild; >1 yields a moderate ARIA-E classification as long as each region is <10 cm in diameter

Severe ARIA-E

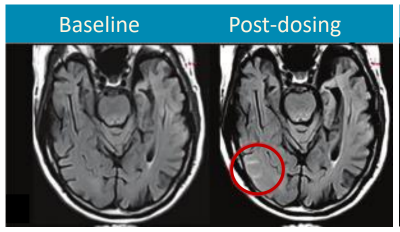
Development of extensive T2-FLAIR hyperintense signal throughout the right frontal and parietal lobes measuring >10 cm
Associated mass effect and sulcal effacement throughout much of the right cerebral hemisphere

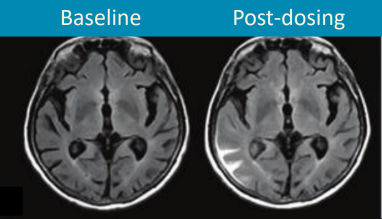
Data shown of 3 different patients
ARIA, amyloid-related imaging abnormalities; ARIA-E, ARIA-edema/effusion,T2-FLAIR, T2-weighted fluid attenuated inversion recovery;
Cogswell PM, et al. AJNR Am J Neuroradiol. 2022;43(9):E19–E35

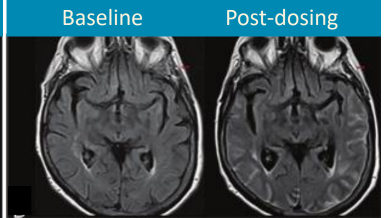


Detection of ARIA-E, sulcal effusion, and corresponding grading

All figures adapted from Cogswell, PM et. al (2022)







Mild ARIA-E

New sulcal T2-FLAIR hyperintense signal in the right temporal-occipital lobe measuring <5 cm in transverse dimensions

Moderate ARIA-E

New T2-FLAIR sulcal effusion involving the right posterior temporal and parietal lobes measuring 5–10 cm

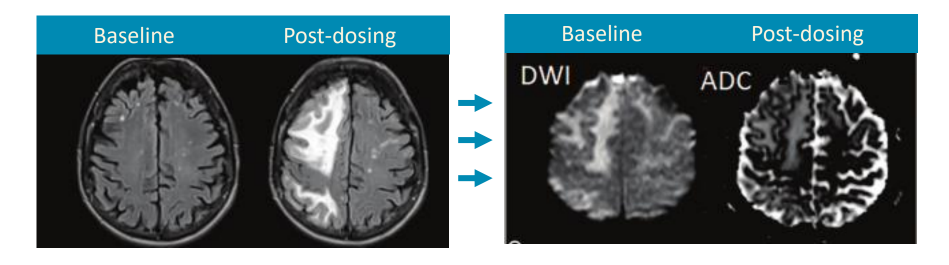
Severe ARIA-E

Extensive T2-FLAIR sulcal effusion involving the bilateral temporal and occipital lobes measuring ≥10 cm in extent

Data shown of 3 different patients
ARIA, amyloid-related imaging abnormalities; ARIA-E, ARIA-edema/effusion; MRI, magnetic resonance imaging; T2-FLAIR, T2-weighted fluid attenuated inversion recovery
Cogswell PM, et al. AJNR Am J Neuroradiol. 2022;43(9):E19–E35



Differentiating ARIA-E from Ischemic Stroke



Severe ARIA-E

Development of extensive T2-FLAIR hyperintense signal throughout the right frontal and parietal lobes measuring >10 cm (severe ARIA-E). Associated mass effect and sulcal effacement throughout much of the right cerebral hemisphere

Hyperintense signal on Diffusion Weighted Imaging (DWI) is confirmed to be T2 shine-through on the Apparent Diffusion Coefficient (ADC) map, differentiating ARIA-E from acute ischemia or other cause of cytotoxic edema

Data shown of 3 different patients

ADC, Apparent Diffusion Coefficient; ARIA, amyloid-related imaging abnormalities; ARIA-E, ARIA-edema/effusion; DWI, diffusion-weighted imaging; T2-FLAIR, T2-weighted fluid attenuated inversion recovery Cogswell PM, et al. AJNR Am J Neuroradiol. 2022;43(9):E19–E35



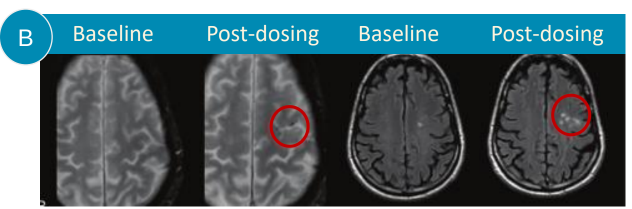
Detection of ARIA-H, microhemorrhage, co-occurring with ARIA-E

A leakage of heme products in the parenchyma, as a result of ARIA-E, can result in microhemorrhages*



T2*GRE T2-FLAIR

- Mild ARIA-H: few (<5) new peripheral left frontal microhemorrhages (red circle) that occur with new patchy T2-FLAIR hyperintense signal in that region
- Moderate ARIA-H: 5 treatment-emergent microhemorrhages (red circle) that occurred with regional mild ARIA- E
- Severe ARIA-H: ≥10 new microhemorrhages and associated extensive right cerebral hemisphere T2-FLAIR hyperintense signal (red circle)



T2*GRE T2-FLAIR Post-dosing Baseline Baseline Post-dosing

T2*GRE

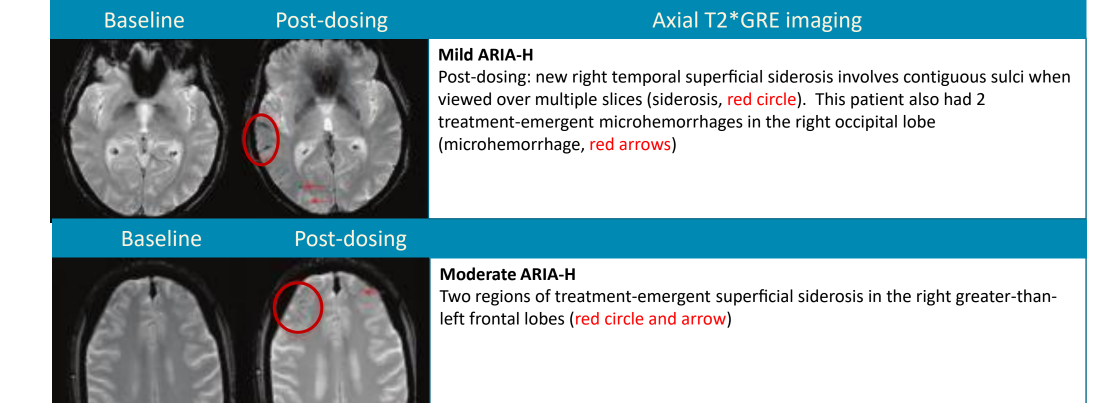
T2-FLAIR *Data shown of 3 different patients

ARIA, amyloid-related imaging abnormalities; ARIA-E, ARIA-edema/effusion; ARIA-H: ARIA-hemosiderin/hemorrhage; GRE, gradient recalled echo; T2-FLAIR, T2-weighted fluid attenuated inversion recovery Cogswell PM, et al. AJNR Am J Neuroradiol. 2022;43(9):E19-E35





Detection of ARIA-H, superficial siderosis, and corresponding grading



Figures adapted from Cogswell et al. (2022)

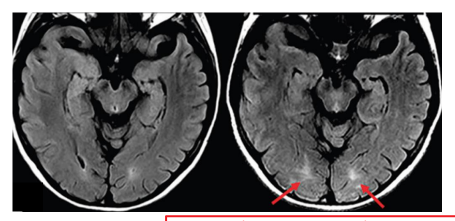
Data shown of 2 different patients
ARIA, amyloid-related imaging abnormalities; ARIA-H: ARIA-hemosiderin/hemorrhage; GRE, gradient recalled echo
Cogswell PM, et al. AJNR Am J Neuroradiol. 2022;43(9):E19–E35



Potential interpretation pitfalls of MRI when detecting ARIA-E

If a patient is imaged on different scanners, it may be difficult to distinguish true ARIA-E versus technical variation¹

Vendor 1: Time-point 1 Vendor 2: Time-point 2



T2-FLAIR hyperintense signal in the bilateral occipital white matter that may be mistaken for subtle ARIA-E, which appears to be new from the prior examination on vendor 1

Figure reproduced with permission from Cogswell et al (2022).

White matter signal may differ with scan technique and field strength, such as the use of 3D versus 2D FLAIR

Shading artifacts and scanner or sequence variability may make identification and interpretation of ARIA-E versus artifacts difficult

- Axial T2-FLAIR images from two time points with the two scans performed on different vendor scanners
- Repeat imaging of participant on vendor 1 showed that the apparent abnormality was resolved

ARIA-E can be identified using T2-weighted FLAIR sequences, but can be entirely obscured with T2-weighted imaging²

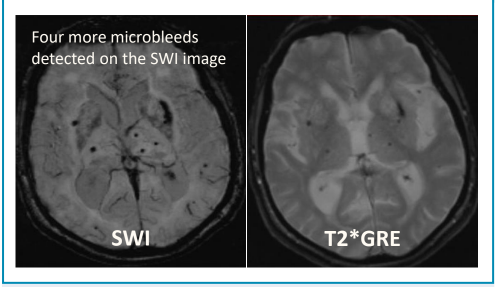
ARIA, amyloid-related imaging abnormalities; ARIA-E, ARIA-edema/effusion; MRI, magnetic resonance imaging; T2-FLAIR, T2-weighted fluid attenuated inversion recovery 1. Cogswell PM, et al. AJNR Am J Neuroradiol. 2022;43(9):E19-E35. 2. Barakos J, et al. AJNR Am J Neuroradiol 2013;34:1958-1965





Potential interpretation pitfalls of MRI when detecting ARIA-H

SWI is a more sensitive technique for detection of microhemorrhages than T2*GRE images¹



Images acquired from the same patient on the same day Figure reproduced with permission from Sperling et al (2011).

Enhanced sensitivity with SWI is accomplished by forming both a magnitude and a phase image and multiplying the magnitude image by the phase image³

The conspicuity of microhemorrhages can be increased based on sequence and magnetic field strength²

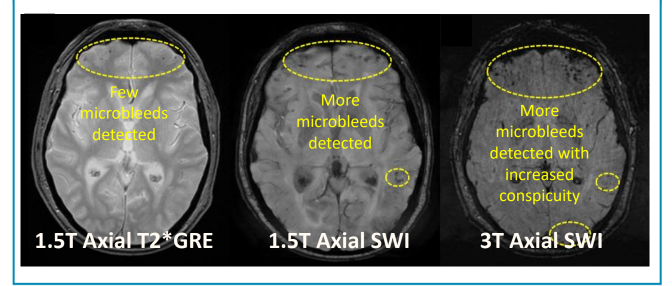


Image of a patient with spontaneous intracerebral hemorrhage Figure reproduced with permission from Puy et al (2021).

Thick-section acquisitions may make it difficult to distinguish a mH from a vessel flow void³

ARIA, amyloid-related imaging abnormalities; ARIA-H, ARIA-hemosiderin/hemorrhage; GRE, gradient-recalled echo; MRI, magnetic resonance imaging; SWI, susceptibility-weighted imaging.

1. Sperling RA, et al. Alzheimers Dement. 2011;7(4):367–385; 2. Puy L, et al. J Neurol Neurosurg Psychiatry. 2021;92(6):598–607; 3. Cogswell PM, et al. AJNR Am J Neuroradiol. 2022;43(9):E19–E35



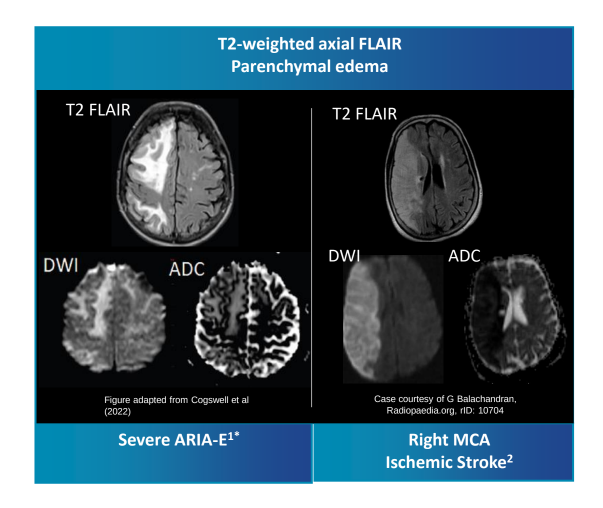
Differentiating ARIA from other pathologies

ARIA-E or ARIA-H should be considered as the presumptive diagnosis when signal abnormalities on MRI are identified in patients recently exposed to monoclonal antibodies that remove amyloid plaque and in whom no evidence of any other inciting cause or underlying lesion can be found¹

- In a suspected ARIA case, the full clinical picture must be taken into account before a diagnosis is confirmed¹
- MRI is key for the diagnosis and differential diagnosis of ARIA²
- CT would not be expected to detect milder forms of ARIA-edema/effusion (ARIA-E) and is insensitive to the detection of microhemorrhages and siderosis (ARIA-H)²
- Training should be provided to ensure reliable diagnosis of ARIA²



Differential diagnosis: acute ischemic stroke



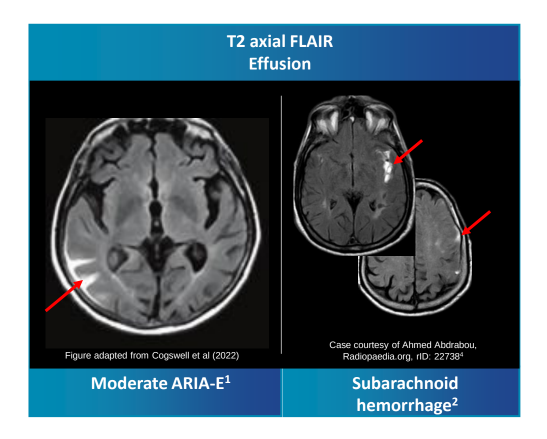
- Parenchymal FLAIR hyperintensity of ARIA-E edema may be mimicked by ischemic stroke³
- Diffusion weighted imaging (DWI) is needed to differentiate between ARIA-E and ischemic stroke³
- Signs and symptoms of ischemic stroke include: acute onset, hemiparesis, dysphasia or dysarthria, facial paresis, paresthesia, eye movement abnormalities, and visual field defects⁴
- Knowing if a patient is on monoclonal antibodies that remove amyloid helps with determining the diagnosis of ARIA³

^{*}Hyperintense signal on DWI is confirmed to be T2 shinethrough on the ADC map, differentiating ARIA-E from acute ischemia or other cause of cytotoxic edema ADC, Apparent Diffusion Coefficient; ARIA-E, ARIA-edema/effusion; DWI, diffusion-weighted imaging; T2-FLAIR: T2 Fluid attenuated inversion recovery; MCA, middle cerebral artery

1. Cogswell PM, et al. AJNR Am J Neuroradiol. 2022;43(9):E19–E35; 2. Bhuta S, et al. Radiopaedia.org https://radiopaedia.org/articles/13401; 3. Barakos, J et al. AJNR AM J Neuroradiol 2013;34:1958-1965; 4. Yew KS, et al. Am Fam Physician. 2015;91(8):528–36



Differential diagnosis: subarachnoid hemorrhage



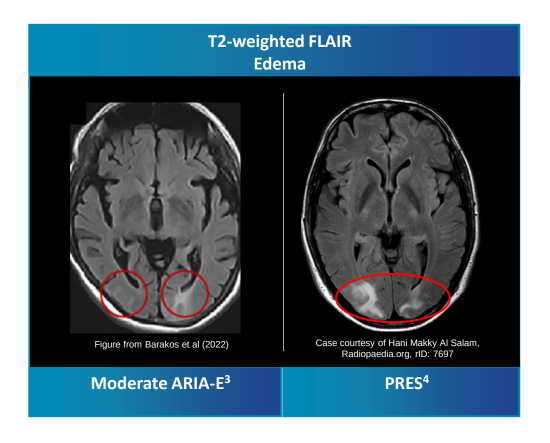
- Leptomeningeal FLAIR hyperintensity of ARIA-E effusion may be mimicked by SAH³
- Differentiating ARIA and SAH requires a systematic clinical and diagnostic approach³
- Subarachnoid hemorrhage typically presents with a number of signs and symptoms: severe headache accompanied by nausea or vomiting⁴
- Decreased level of consciousness and focal neurological signs can also be present⁴

ARIA-E, ARIA-edema/effusion; SAH, subarachnoid hemorrhage; T2-FLAIR, T2-Fluid-attenuated inversion recovery

1. Cogswell PM, et al. AJNR Am J Neuroradiol. 2022;43(9):E19–E35; 2. Abdrabou A. Radiopaedia.org https://doi.org/10.53347/rID-22738; 3. Barakos, J et al. AJNR AM J Neuroradiol 2013;34:1958-1965; 4. Tetsuka S, et al. BMC Neurol 2016;16:196



Differential diagnosis: Posterior Reversible Encephalopathy Syndrome (PRES)



- PRES could resemble ARIA-E on imaging¹
- PRES frequently develops from cytotoxic medication or disorders such as preeclampsia, sepsis, renal disease, or autoimmune disorders²
- Signs of PRES²:
 - Encephalopathy, epileptic seizures, visual disturbances, and focal neurological deficits
- Less specific signs include:²
 - Headache, nausea, vomiting
- In this case, clinical history is **important for** differentiation

ARIA, amyloid-related imaging abnormalities; ARIA-E, ARIA-edema/effusion; T2-FLAIR, T2-Fluid-attenuated inversion recovery 1. Barakos, J et al. AJNR AM J Neuroradiol 2013;34:1958-1965; 2. Fischer M, et al. J Neurol 2017;264:1608-1616 3. Barakos J, et al. J Prev Alzheimers Dis. 2022;9(2):211-220; 4. Gaillard F, et al. https://doi.org/10.53347/rID-1915









Management of ARIA



Refer to prescribing information of monoclonal antibodies that remove amyloid for monitoring and management guidelines of ARIA



Discuss ARIA and associated symptoms with patients and care partners before treatment initiation including the importance of MRI monitoring and seeking urgent evaluation in the case of ARIA clinical symptoms^{1,2}



MRI should be used to assess for ARIA symptoms where possible; CT scans can be deficient for detecting radiographic findings, particularly ARIA-H, owing to its relatively low spatial definition and resolution vs MRI³



ARIA is most frequently detected on routine surveillance MRIs in patients who are clinically asymptomatic, highlighting the need for monitoring early in the course of therapy⁴



In cases of severe or serious ARIA-E or ARIA-H, monitoring neurologic status closely and early empiric administration of high dose intravenous corticosteroids should be considered¹

ARIA, amyloid-related imaging abnormalities (due to ARIA-E and ARIA-H); ARIA-E, ARIA-edema/effusion; ARIA-H, ARIA-hemosiderin/hemorrhage; CT, computed tomography; MRI, magnetic resonance imaging.

1. Cummings J, et al. J Prev Alzheimers Dis 2022;9:221–230; 2, Cummings J et al. Alzheimers Dement. 2021;7(1):e12179 3. Barakos J, et al. J Prev Alzheimers Dis. 2022;9(2):211–220; 4. Cogswell PM, et al. AJNR Am J Neuroradiol. 2022;43(9):E19–E35.

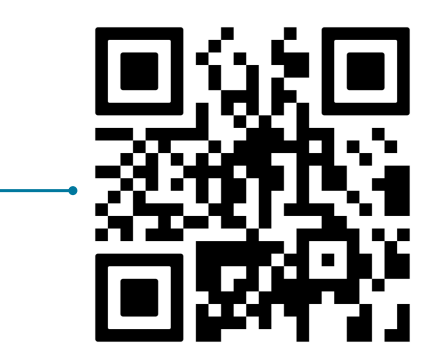




To access a growing repository of educational resources on ARIA, please scan the QR code or access the platform by the following link:

www.UnderstandingARIA.com

This information is intended for healthcare professionals only.



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