

Cohere Medical Policy -Magnetic Resonance Imaging (MRI), Brain

Clinical Guidelines for Medical Necessity Review

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Type: [X] Adult (18+ yo) | [X] Pediatric (0-17yo)

Table of Contents

Important Notices	2
Table of Contents	3
Medical Necessity Criteria	4
Service: Magnetic Resonance Imaging (MRI), Brain	4
Recommended Clinical Approach	4
Indications	4
Non-Indications	15
Level of Care Criteria	16
Procedure Codes (CPT/HCPCS)	16
Medical Evidence	
References	18
Clinical Guideline Revision History/Information	22

Medical Necessity Criteria

Service: Magnetic Resonance Imaging (MRI), Brain

Recommended Clinical Approach

Imaging analysis utilizing magnetic resonance imaging (MRI) brain can be performed alone or with magnetic resonance angiography (MRA) head based on clinical suspicion of disease presence or exclusion to direct value-based care. Contrast may or may not be necessary depending upon the clinical indication at the referring physician's request and the discretion of the supervising radiologist.

Medical Necessity Criteria

Indications

- → Magnetic resonance imaging (MRI), brain is considered appropriate if ANY of the following is TRUE¹⁻²:
 - ◆ The patient has **ANY** of the following neoplastic conditions (masses or mass-like conditions):
 - Brain parenchyma³; **OR**
 - Cranium⁴; **OR**
 - Meninges; OR
 - Meningioma; OR
 - Known or suspected parasellar (e.g., pituitary) tumors; OR
 - ◆ The patient has ANY of the following infections or infectious disorders⁵:
 - Suspected brain abscess or brain infection with ANY of the following:
 - Acute altered mental status; OR
 - Positive lab findings (e.g., elevated WBCs or abnormal CSF analysis); OR

- Follow-up assessment during or after treatment completed; OR
- Endocarditis with suspected septic emboli; OR
- Neurosarcoidosis with ANY of the following:
 - For initial evaluation with ANY of the following:
 - Suspected based on neurological sign(s), symptom(s), and lab work (e.g., angiotensin-converting enzyme (ACE], cerebrospinal fluid (CSF) analysis); OR
 - Known history of sarcoidosis with neurological signs or symptoms; OR
 - Follow-up of known neurosarcoidosis for ANY of the following reasons:
 - To assess treatment response; OR
 - Worsening signs or symptoms; OR
- The patient has a trauma-related condition, as indicated by ANY of the following⁶:
 - Assessment of unexplained post-traumatic neurological deficits; OR
 - Post-traumatic brain injury or persistent symptoms following initial imaging (e.g., dizziness, headache); OR
 - Post-concussive syndrome if persistent or disabling symptoms and MRI has not been performed; OR
 - Subacute or chronic traumatic brain injury with new cognitive and/or neurologic deficit; OR
- The patient has a vascular condition including ANY of the following:
 - Acute ischemic stroke or transient ischemic attack (TIA)⁷;
 OR
 - Known or suspected stroke with any acute, new, or fluctuating symptoms or deficits (e.g., sensory deficits, limb

- weakness, speech difficulties, visual loss, lack of coordination, or mental status changes); **OR**
- Suspected stroke with a personal or first-degree family
 history (e.g., brother, sister, parent, or child) of an aneurysm
 or known coagulopathy or on anticoagulation; OR
- Symptoms of TIA that are episodic with neurologic symptoms (e.g., sensory deficits, limb weakness, speech difficulties, visual loss, lack of coordination, or mental status changes); OR
- Evaluation of suspected acute subarachnoid hemorrhage (SAH); OR
- Follow-up for known hemorrhage, hematoma, or vascular abnormalities; OR
- Follow-up imaging of known cerebral cavernous
 malformations (CCM) should be done only to guide
 treatment decisions or to investigate new symptoms includes first-degree relatives of patients with one family
 member with a CCM should have a screening MRI as well as
 genetic counseling; OR
- Suspected central venous thrombosis; OR
- Screening for silent cerebral infarcts in early school-age children and adults with HbSS sickle cell disease or HbSβ0 thalassemia; OR
- Suspected temporal arteritis in a patient greater than 50 years old with ANY of the following:
 - Temporal headache; OR
 - Abrupt visual changes; OR
 - o Jaw claudication; OR
 - o Temporal artery tenderness; **OR**
 - o Constitutional symptoms; **OR**
 - Elevated erythrocyte sedimentation rate (ESR); OR

- Evaluation of neurological signs or symptoms in a patient with sickle cell disease; OR
- High stroke risk in sickle cell patients (2 to 16 years of age)
 with a transcranial Doppler velocity greater than 200
 cm/sec3⁸; OR
- **ANY** of the following chronic vascular diseases⁹:
 - Headaches with a family history of aneurysm/AVM,
 including ANY of the following 10-13:
 - One first-degree relative with a history of aneurysm/AVM if symptomatic or there is clinical concern for aneurysm; OR
 - Two or more relatives with a history of aneurysm/AVM (repeat imaging in 5 years); OR
 - Arterial or venous/dural venous sinus abnormalities such as ANY of the following:
 - Acquired thrombosis/occlusion; OR
 - ♦ Venous sinus stenosis; OR
 - Central nervous system (CNS) involvement in patients with known or suspected vasculitis or autoimmune disease with abnormal inflammatory markers or autoimmune antibodies; OR
 - Immunocompromised patients (e.g., transplant recipients, HIV with CD4 less than 200, primary immunodeficiency syndromes, hematologic malignancies) with focal neurologic symptoms, headaches, behavioral, cognitive or personality changes; OR
 - Evaluation of neurological signs or symptoms in vaso-occlusive disease (e.g., sickle cell disease, moyamoya, etc.); OR

- ◆ The patient has ANY of the following autoimmune or inflammatory conditions¹⁴⁻¹⁵:
 - Atypical trigeminal neuralgia, defined by ANY of the following symptoms¹⁶:
 - o Bilateral hearing loss; OR
 - o Dizziness/vertigo; **OR**
 - Visual changes; OR
 - Sensory loss or numbness; OR
 - o Pain greater than 2 minutes; **OR**
 - Pain outside trigeminal nerve distribution and progression; OR
 - Suspected multiple sclerosis (MS), and ANY of the following is TRUE:
 - For evaluation of patients with neurologic symptoms or deficits suspicious for MS with ANY of the following:
 - A clinically isolated syndrome (e.g., optic neuritis, transverse myelitis, or brain stem syndrome); OR
 - Recurrent episodes of variable neurological signs or symptoms not attributable to another cause; OR
 - To demonstrate dissemination in time for diagnosis (every 6-12 months); OR
 - For evaluation of known multiple sclerosis (MS) and
 ANY of the following is TRUE:
 - ◆ To establish a new baseline and ANY of the following is TRUE:
 - No recent imaging; OR
 - Postpartum; OR
 - 3-6 months after switching disease-modifying therapy; OR

- Before starting or switching disease-modifying therapy; OR
- 6-month repeat scan in patients with MRI
 disease activity that is not associated with new
 clinical symptoms on a routine follow-up scan
 (e.g., radiographically isolated syndrome); OR
- Every 1-2 years while on disease-modifying therapy to assess for subclinical disease activity, less frequently when stable for 2-3 years; OR
- New signs or symptoms suggested an exacerbation or unexpected clinical worsening;
 OR
- Progressive multifocal leukoencephalopathy
 (PML) surveillance for patients on natalizumab
 (Tysabri) and ANY of the following is TRUE:
 - 12 months after the start of treatment in all patients; OR
 - Brain MRI every 3–4 months for up to 12 months, in high-risk patients who switch from natalizumab to other therapeutics;
 OR
 - Further surveillance MRI scanning timing is based on risk, and ANY of the following is TRUE:
 - Annually, if the anti-JCV antibody is negative; OR
 - Every 3-4 months, if high-risk of PML occurrence and ANY of the following is TRUE:
 - Seropositive for JC virus and

- have been treated with natalizumab for greater than or equal to 18 months; **OR**
- High anti-JC virus antibody index values (greater than 0.9); OR
- Previously treated with immunosuppressive therapies; OR
- ◆ In the pediatric population, a similar scan frequency for disease and therapeutic monitoring should be used - the frequency should be increased (e.g., every 6 months) if the child has a highly active disease or when imaging will change management; OR
- ◆ Evaluation of **ANY** of the following miscellaneous pathologies:
 - Acute, new, or fluctuating neurologic symptoms or deficits, including ANY of the following:
 - Sensory deficits; OR
 - o Limb weakness; **OR**
 - Abnormal reflexes (pathological, asymmetric, hyperreflexia); OR
 - Speech difficulties; OR
 - o Visual loss; OR
 - o Lack of coordination; **OR**
 - o Mental status changes; **OR**
 - Brain structural abnormality identified or suspected on prior imaging; OR
 - Chronic disequilibrium with signs of cerebellar ataxia[™]; OR
 - Pathology involving the cranial nerve¹⁸; OR
 - Dizziness (including vertigo) with ANY of the following:

- Signs or symptoms suggestive of a CNS lesion (ataxia, visual loss, double vision, weakness, or a change in sensation); OR
- Progressive unilateral hearing loss; OR
- Risk factors for cerebrovascular disease with concern for stroke; OR
- Acute persistent vertigo with normal neurologic examination and HINTS examination is consistent with peripheral vertigo¹⁷; OR
- Acute persistent vertigo with abnormal neurologic examination or HINTS examination is consistent with central vertigo¹⁷; OR
- Chronic recurrent vertigo that is associated with unilateral hearing loss or tinnitus¹⁷; OR
- Chronic recurrent vertigo that is associated with other brainstem neurologic deficits¹⁷; OR
- Episodic vertigo with or without associated hearing loss or aural fullness¹⁹; OR
- Persistent vertigo with or without neurological symptoms (central vertigo)¹⁹; OR
- Epilepsy, known or suspected, and **ANY** of the following is **TRUE**:
 - New onset of seizures or newly identified change in seizure activity/pattern; OR
 - o Known seizure disorder without prior imaging; OR
 - Medically refractory epilepsy; OR
- Neurodegenerative disorders (congenital or acquired), including dementia²⁰⁻²¹; OR
- Movement disorders including ANY of the following^{20,22-24}; OR

- Acute onset of a movement disorder with concern for stroke or hemorrhage (brain MRI recommended over brain CT); OR
- For evaluation of Parkinson's disease with atypical features or other movement disorders (e.g., suspected Huntington's disease, chorea, parkinsonian syndromes, hemiballismus, atypical dystonia) to exclude an underlying structural lesion; OR
- Headaches as indicated by ANY of the following³:
 - In a pediatric patient with severe headaches who is less than 6 years old; OR
 - In a pediatric patient who is greater than or equal to 6 years old with severe headaches and a known underlying condition (e.g., sickle cell, cancer, immune deficiency); OR
 - Immediate imaging is required and the patient is considered high-risk with ANY of the following symptoms³:
 - Intracranial hypertension (e.g., papilledema, pulsatile tinnitus, visual symptoms worse on Valsalva); OR
 - Intracranial hypotension (e.g., positional, worse when upright, better when lying down); OR
 - ◆ ANY of the following "red flags":
 - Age of onset is 50 years or older; OR
 - Fever or neurologic deficit; **OR**
 - History of cancer or immunocompromise;
 OR
 - Increasing frequency or severity; OR
 - Posttraumatic onset with no prior imaging; OR

- Headache with new onset or pattern during pregnancy or peripartum period; OR
- Primary trigeminal autonomic cephalalgias (e.g., cluster headache); OR
- Persistent headaches that have failed physician-directed conservative treatment; OR
- Hearing loss (e.g., acquired sensorineural, mixed conductive, sensorineural)¹⁹; OR
- Acoustic neuroma including ANY of the following²⁵:
 - o Initial evaluation for suspected acoustic neuroma; OR
 - Monitoring of a known, symptomatic acoustic neuroma when surgery is not performed; OR
 - o Post-surgical management; OR
- Metabolic, nutritional, and dysmyelinating disorders²⁶; OR
- ANY of the following types of tinnitus²⁷:
 - Pulsatile tinnitus, unilateral or bilateral without retro tympanic lesion on otoscopy; OR
 - Nonpulsatile tinnitus, unilateral without hearing loss, neurologic deficit, or trauma; OR
- **ANY** of the following visual conditions²⁸:
 - Nontraumatic orbital asymmetry, exophthalmos, or enophthalmos; OR
 - Ophthalmoplegia or diplopia; OR
 - o Suspected orbital cellulitis, uveitis, or scleritis; OR
 - o Suspected optic neuritis; OR
 - Traumatic visual defect; OR
 - Vision loss with intraocular mass, optic nerve, or visual field defects; OR
 - Vision loss, nonischemic, with visual field defects; OR
- The patient requires ANY of the following preoperative, postoperative, or pre-treatment evaluations:

- Follow-up of treatment, including iatrogenic sequelae such as radiation necrosis²⁹; OR
- The patient has acquired conductive hearing loss (secondary to cholesteatoma or neoplasm) with suspected intracranial or inner ear extension, for ANY of the following purposes¹⁹:
 - Surgical planning; OR
 - Monitoring when surgery is not performed; OR
 - Post-surgical management; OR
- The patient has congenital hearing loss, total deafness, or is a cochlear implant candidate for **ANY** of the following 19:
 - Surgical planning; OR
 - o Post-surgical management; OR
- The patient has a known or suspected congenital abnormality with any acute, new, or fluctuating neurologic, motor, or mental status changes, including ANY of the following:
 - Craniosynostosis and other head deformities; OR
 - Hydrocephalus (congenital or acquired)^{3,30-31}; OR
 - Idiopathic intracranial hypertension; OR
 - Intracranial hypotension; OR
- Repeat imaging of a specific area or structure using the same imaging modality (in the absence of an existing follow-up guideline) is considered appropriate when ALL of the following is TRUE:
 - There is documented clinical necessity; AND
 - Prior imaging results of the specific area or structure, obtained using the same imaging modality, must be documented and available for comparison; AND
 - **ANY** of the following is **TRUE**:
 - A change in clinical status, such as worsening symptoms or the emergence of new symptoms, that

- may influence the treatment approach; OR
- The requirement for interval reassessment, which may alter the treatment plan; OR
- One-time follow-up of a prior indeterminate finding to assess for interval change; OR
- The need for re-imaging either before or after performing an invasive procedure.

Non-Indications

- → Magnetic resonance imaging (MRI), brain is not considered appropriate if ANY of the following is TRUE:
 - The patient has undergone advanced imaging of the same body part and for the same indication within 3 months, without being on treatment; OR
 - If contrast is used, history of anaphylactic allergic reaction to gadolinium contrast media with detailed guidelines for use in patients with renal insufficiency; OR
 - ◆ The patient has metallic clips on vascular aneurysms; **OR**
 - Incompatible implantable devices (e.g., pacemakers, defibrillators, cardiac valves); OR
 - Metallic foreign body in orbits/other critical area(s) or within the field of view and obscuring area of concern; OR
 - ◆ In the pediatric population, imaging is not indicated in simple febrile seizures or in idiopathic focal or generalized epilepsy with typical features of benign epilepsy with centro-temporal spikes (BECTS), childhood absence epilepsy (CAE), juvenile absence epilepsy (JAE), and juvenile myoclonic epilepsy (JME).

^{*}NOTE: MRI in patients with claustrophobia should be requested at the discretion of the ordering provider.

**NOTE: MRI in pregnant patients should be requested at the discretion of the ordering provider and obstetric care provider.

Level of Care Criteria

Inpatient or Outpatient

Procedure Codes (CPT/HCPCS)

CPT/HCPCS Code	Code Description	
70551	Magnetic resonance imaging (MRI) (e.g., proton), brain (including brain stem); without contrast material	
70552	Magnetic resonance imaging (MRI) (e.g., proton), brain (including brain stem); with contrast material(s)	
70553	Magnetic resonance imaging (MRI) (e.g., proton), brain (including brain stem); without contrast material, followed by contrast material(s) and further sequences	

Medical Evidence

Wangaryattawanich et al. (2023) review the imaging spectrum, clinical significance, and management of brain MRI. Rapid advancements have been made in imaging technology and improved accessibility. Consequently, radiologists discover incidental findings during brain MRI scans for unrelated reasons. These unexpected findings can range from clinically insignificant to requiring further investigation or treatment, leading to patient anxiety. Incidental findings encompass a diverse range, including asymptomatic brain infarcts, age-related white matter changes, microhemorrhages, intracranial tumors, intracranial cystic lesions, and anatomic variants.³²

Maas et al. (2022) analyze facets of managing traumatic brain injury (TBI), including imaging. An initial normal CT scan does not rule out the presence of structural traumatic abnormalities. Structural traumatic abnormalities seen on MRI (2–3 weeks post-injury) were observed in approximately 30% of patients with mild TBI who initially had a normal CT scan. Advanced MRI techniques, such as diffusion tensor imaging and volumetric analyses, can reveal further injuries that may not be discernible through visual examination of conventional clinical MR images. Emerging blood biomarkers, such as glial fibrillary acidic protein (GFAP), aid in refining decisions regarding the necessity of CT scans for patients with mild TBI or the requirement of an MRI if the initial CT scan is normal. Integrating biomarkers, quantitative CT, and MRI findings facilitates the identification of patients at-risk of persistent symptoms, enabling more tailored and frequent follow-up care.³³

Tekes et al. (2018) review the use of brain MRI in pediatric patients beyond shunted hydrocephalus. In the study period, 800 patients had undergone a previous ultrafast brain MRI scan. Patients diagnosed with ventriculomegaly, macrocephaly, or intracranial cysts were included (n=101). The findings support the use of ultrafast brain MRI to assess for these conditions. Given its radiation-free and sedation-free nature, ultrafast brain MRI may be appropriate as a primary screening neuroimaging modality for these indications.³⁴

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Clinical Guideline Revision History/Information

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Review History			
Version 2	8/2/2024	Annual review and policy restructure.	