Clinical Neurological Imaging on an Open Bore MRI System (MAGNETOM Espree)

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Introduction

An open bore MRI has the practical advantage of accommodating large or claustrophobic patients. Unfortunately, until the advent of the 1.5T MAGNETOM Espree, “open” was often synonymous with inferior image quality. However, in addition to the large 70 centimeter bore, the Espree system offers advanced coil combinations using the Total imaging matrix (Tim) and isocenter imaging technology. Together, these features allow for performance of advanced neuroimaging protocols in new clinical populations.

This article demonstrates neuroimaging applications routinely obtained in our clinical practice at Mallinckrodt Institute of Radiology, Washington University School of Medicine in St. Louis. The open bore Espree delivers excellent image quality, expanding the utility of MRI to meet diagnostic challenges increasingly encountered in clinical neuroimaging.

Methods

Image acquisitions were performed with a 1.5 Tesla MAGNETOM Espree scanner using the 12-channel head coil and spine coils of the Total imaging matrix. Our standard neurological examination on the Espree system includes common cross-platform protocols such as MPRAGE, FLAIR, T2 TSE, DSC Perfusion, and Spectroscopy. The Espree also enables advanced imaging sequences, including T2 BLADE [1], Susceptibility-Weighted Imaging (SWI) [2], and Diffusion Tensor Imaging (DTI) [3], for which we use the following protocols:

**T2 BLADE protocol:** TE = 102 msec, TR = 4000 msec, FOV = 240 mm, pFOV = 100%, slice thickness = 5 mm, base resolution = 384 (results in spatial resolution = 0.6 x 0.6 x 5.0 mm), average = 1, concatenations = 2, flip angle = 150°, BLADE coverage = 111%, echo spacing = 6.64 msec, turbo factor = 35, echo train per slice = 20, motion correction = ON, bandwidth = 362 Hz/pixel, acquisition time: 2:50 min.

**SWI protocol:** TE = 40 msec, TR = 50 msec, FOV = 240, pFOV = 100%, slice thickness = 2 mm, slices per slab = 72, base resolution = 256, phase resolution = 79%, slice resolution = 75% (results in spatial resolution = 1.2 x 0.9 x 2.0 mm), average = 1, flip angle = 15, PAT acceleration factor = 2, bandwidth = 80 Hz/pixel, spatial resolution = 1.2 x 0.9 x 2.0 mm, acquisition time 3:34 min.

**DTI protocol:** TE = 107 msec, TR = 3500 msec, FOV = 240 mm, pFOV = 100%, slice thickness = 5 mm, base resolution = 128, phase resolution = 100% (results in spatial resolution = 1.9 x 1.9 x 5 mm), averages = 3, Fatsat, phase partial Fourier = 6/8, PAT acceleration factor = 2, echo spacing = 0.92 msec, EPI factor = 128, BW = 1220 Hz/pixel, b-values = 0, 1000 sec/mm2, 12 diffusion directions. Average ADC map, trace-weighted map, FA map, and tensor data were created In-line. Acquisition time: 2:08 min.

Post-processing was performed with the Neuro 3D application package. The package has the capability of processing directional color encoded fractional anisotropy map (blue = SI direction, green = AP direction, and red = RL direction), tensor map, aligned tensor and anatomy data, aligned tensor and fractional anisotropy data, and tractography.

Clinical cases

The following examples demonstrate brain stem pathology which is difficult to image on a non-Tim system due to the position of the lesion and unavailability of coil combination. For brainstem cases examined on the MAGNETOM Espree system, the head coil and cervical spine elements were combined using the Total imaging matrix to better visualize the lesions. Tumor cases are also routinely examined on the Espree system. An example is also included in this article.
Patient 1

77-year-old man undergoing evaluation for stroke had a normal MRI (Fig. 1). The study exhibits tracts of the normal brain acquired with the 1.5T Espree system. DTI tractography processed with the Neuro 3D software resolves pontocerebellar fibers and the pyramidal decussation with great corresponding anatomic detail [4]. The DTI dataset is acquired in less than 3 minutes, and replaces the standard 3 direction DWI data for clinical stroke imaging on this scanner.

Figure 1
Normal brain MRI acquired on an open bore 1.5T MAGNETOM Espree using 12-channel head coil and spine coils of the Total imaging matrix:
A: Color fractional anisotropy of the normal pontine white matter.
B: Axial T2 BLADE demonstrates the corresponding pontine anatomy.
C: Normal DTI tractography showing the red pontine fiber shown in A (white arrow) and pyramidal decussation (red arrow). These images were generated from a 2 minute scan.
Patient 2

This 39-year-old obese woman presented for evaluation of dysphagia and left upper and lower extremity paresthesias. The brain MRI demonstrates a non-enhancing T2 hyperintense brainstem mass extending from the medulla into the cervical cord. Spectroscopy reveals a decreased NAA to choline signal ratio within this lesion. There is no diffusion restriction or T2* abnormality, but diffusion tractography is useful in demonstrating the spatial distortion by the expansile mass without infiltration of the longitudinal brainstem white matter tracts (Fig. 2).

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Following radiotherapy and four cycles of Temodar, the MRI study is repeated on the same equipment. There is no interval change (Fig. 3).

Patient 3

This 52-year-old man with a 12 months history of diplopia, dizziness and progressive left facial weakness presented to an outside facility where an MRI revealed a non-enhancing brainstem mass. A more detailed study on the Espree system demonstrates a mass arising in the posterior pons from midbrain to medulla with extension into the left middle cerebellar peduncle. Spectroscopy demonstrates decreased NAA and elevated choline signal within the lesion (Fig. 4).
Patient 4

This 30-year-old man was evaluated for six weeks of progressive nausea and headache. He was found to have an area of irregular, infiltrative T2 hyperintensity in the pons. T1 TSE sequence post contrast demonstrates regions of mild enhancement. Although the absolute value of choline is not elevated by spectroscopy, the signal ratio of NAA to choline is depressed. These imaging characteristics favor a glial neoplasm, which was confirmed by open biopsy of a spinal drop metastasis (Fig. 5).

A: Axial T2 BLADE and (B) Sagittal T2 TSE demonstrate the T2 hyperintensity of the mass. C: T1 TSE post contrast shows enhancement of the mass. D: Single voxel spectroscopy reveals the reduced NAA to choline ratio within the lesion (right) as compared to this patient’s normal tissue (left).
Patient 5
This 55-year-old man presented with a several week history of right-sided numbness, unsteadiness, and a mild comprehensive and expressive aphasia. Biopsy of a left thalamic glioblastoma multiforme resulted in parenchymal hemorrhage, necessitating open evacuation (Fig. 6). T1 TSE sequence post contrast exhibits avid enhancement at the margins of a centrally necrotic left thalamic mass. The medial portion of the tumor shows increased perfusion. DTI tractography demonstrates anterior deviation of the posterior limb of the left internal capsule (directionally encoded in blue), but the longitudinal fiber tracts remain intact. The transthalamic white matter tracts (yellow for clarity) are displaced radially with abrupt termination. Note inferolateral deviation of the optic radiations [5].

Conclusion
In addition to standard neuroimaging sequences, advanced applications such as spectroscopy, syngo BLADE, syngo SWI, and syngo DTI have produced excellent imaging results on the 1.5T MAGNETOM Espree system at Mallinckrodt Institute of Radiology, Washington University School of Medicine in St. Louis. The Espree scanner enables excellent clinical neurological diagnostics under the most difficult circumstances, including obese and claustrophobic patients as well as difficult imaging anatomy, such as brain stem masses. With the advances of Tim technology, the practicality of an open bore system no longer demands compromises in image quality.
DTI Tractography demonstrates anterior deviation of the intact posterior limb of the left internal capsule (directionally encoded in blue). The optic radiations of the left hemisphere (directionally encoded in green) are displaced inferolaterally. Transthalamic white matter (yellow for clarity) is displaced radially and disrupted at the rostral margin of the tumor.

References

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