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I am excited to be taking on this new role as Editor-in-Chief of MAGNETOM Flash. I would best describe myself as a student and a teacher. What inspires me is the desire to learn new things and apply that learning to my work. I will continue the tradition of focusing on making a difference to our readers, by providing information, education and context.

With changing healthcare economics, attention to patient outcomes and efficiency is critical. MAGNETOM Flash helps to inform on innovations, products and new applications – for example, Daniel Fischer’s XR 80/200 gradient* article provides insight in hardware changes. Marc-André Weber and Jürgen Biederer have contributed an article on applications in clinical practice. That will help align patient needs. The patient is the focus and accurate understanding of the clinical applications is important. It is important to define the right ‘context’ when applying this information? Because we all must grow in our understanding of the clinical applications. It is important for our readers, by providing the context to you, our readers.

Tradition of focusing on making a difference to our readers, by providing the context to you, our readers. While change is permanent, consistency and context are critical. In that regard, my predecessor Editor-in-Chief Milind Dhamankar, M.D. inspires me is the desire to learn new things and apply that learning to my work. I will continue the tradition of focusing on making a difference to our readers, by providing information, education and context.

Sincerely,

Milind Dhamankar, M.D.
Editor-in-Chief

We are looking forward to hearing from you.

Sincerely,

Milind Dhamankar, M.D.
Editor-in-Chief

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Bernhard Schuknecht, M.D.; Krisztina Baràth, M.D.

Diagnostic, Vascular and Interventional Neuroradiology, Medizinisch Radiologisches Institut MRI Zürich, Bethanien Hospital, Zürich, Switzerland

**Functional MRI (fMRI)**

Functional magnetic resonance imaging (fMRI) is based on the blood oxygenation level dependent (BOLD) effect. BOLD contrast rises with neural activation and depends on the variables blood flow and oxygenation. Augmentation of regional cerebral blood flow (CBF) is accompanied by a local increase in oxygen extraction due to a rise in the cerebral metabolic rate of oxygen consumption. The ferrous iron in deoxyhemoglobin adopts paramagnetic properties contrary to the

fMRI and DTI of a brain tumor: function and structure

1A (1A) Brain tumor (oligodendroglioma) (arrow) in the subcentral gyrus/inferior parietal lobule on the left shown on a FLAIR sequence.  
1B (1B) Activation of the primary motor cortex (thin arrow) close to the tumor (arrow) during finger tapping. Red = movement on the right, Green = movement on the left.  
1C (1C) Activation of the motor speech area and the sensory speech area (thin arrows) far away from the tumor (arrow). The DTI sequence (1D–E) shows the deviated U-fibers (thin arrow) due to the tumor (arrow). The Broca- and Wernicke areas are connected through the arcuate fibers, which makes a combination of fMRI and DTI for the preoperative planning necessary.  
1F (1F) The thinning of the left arcuate fibers indicates an infiltration of the speech-network.
diamagnetic effect in oxyhemoglobin. This results in a susceptibility induced field shift and leads to an increase in T2* value and a signal increase in T2* gradient echo or echo planar imaging (EPI) studies. The high sensitivity of EPI sequences enables real time visualization of activation of different parts of the brain in comparison to a resting state. In a functional MR paradigm the patient is asked to alternatively perform tasks or is stimulated to trigger several processes or emotions. Threshold functional MRI activation maps are spatially addressed (Fig. 1A) and therefore typically overlaid in colour on a high resolution anatomical T1 MP RAGE or SPACE FLAIR MR image (Figs. 1B, C).

fMRI was first employed for mapping of neuronal activation during visual perception tasks and motor activation in the early nineties. Since then fMRI has evolved into an invaluable research tool, e.g. in cognitive neuroscience or neuropharmacology as well as a routinely applied clinical MR technique. Clinical applications comprise mainly neurosurgical planning in order to determine resection margins in relation to functional ‘eloquent’ areas. With fMRI brain functional plasticity may also be visualized in recovery after stroke or surgery. Furthermore fMRI can be used to evaluate specific brain functions in a range of neuro-degenerative diseases. Despite existing limitations of fMRI, a significant increase in spatial and temporal resolution has been achieved in order to improve localization of specific brain function and individualize treatment.

**Diffusion Tensor Tractography (Diffusion Tensor Imaging, DTI)**

DTI is a non-invasive MR technique to study brain tissue composition and architecture in vivo. DTI is based on the concept that water diffusion is anisotropic in organized tissues. Cerebral white matter is composed of axons, myelin sheaths and neuroglia which are aligned to form fiber tracts. DTI provides information on water diffusion properties regarding the extent of diffusion anisotropy and its orientation. A pair of magnetic dephasing and rephasing field gradients is successively applied in distinct directions in 3D space. The resulting images exhibit signal attenuation in the direction of the applied gradient which is proportional to water diffusivity. The largest principal axis of the diffusion tensor aligns with the predominant fiber orientation in an MRI voxel. A diffusion tensor or the mean diffusivity may be estimated from as few as 6 diffusion-weighted images acquired along non-collinear directions and 1 minimally = T2-weighted \((b_0)\) image. From the diffusion tensor, the DTI indices derived include the mean diffusion coefficient (ADC), calculated eigenvalues and eigenvectors and an index of diffusion anisotropy, e.g. fractional anisotropy (FA). To achieve a high signal-to-noise ratio DTI employs more than the minimum of 6 diffusion-weighting angulations and/or acquires repeated measurements of diffusion directions. Despite substantial improvement in technique, spatial resolution of DTI in the order of 1–4 mm per dimension, reflects limitations imposed by using a macroscopic technique to visualize microscopic restrictions.

DTI is performed in combination with fMRI for preoperative localization of fiber tracts in proximity to a lesion (Figs. 1D–F), and for preoperative differentiation of white matter tract infiltration from displacement in patients with low grade glioma in particular. Following hemorrhagic or ischemic stroke and trauma DTI has become an important tool to assess white matter tract involvement beyond findings by morphologic imaging. This holds also true for white matter alterations in patients suffering from neurodegenerative and movement disorders, and to visualize secondary neurodegeneration in inflammatory conditions such as multiple sclerosis.

**Diffusion-Weighted Imaging (DWI)**

Diffusion-weighted MR imaging renders microscopic molecular motion of water visible within tissues. The anisotropic nature of diffusion in the brain can be visualized by comparing images obtained with DWI gradients applied in three orthogonal directions. DWI consists of an echo planar spin-echo T2w pulse sequence. Alternative DWI sequences are based on a single-shot gradient or single-shot fast spin-echo technique, ‘line-scan’ and spiral DWI technique. The signal intensity obtained by DWI corresponds to the signal on T2w images decreased by a signal amount that is related to the rate of diffusion in the direction of the applied gradients. Pathology therefore is reflected by high signal on diffusion-weighted images and by decreased signal on apparent diffusion coefficient (ADC) maps.

While DWI is highly sensitive in reflecting the physical properties of diffusion, the observation of restricted water diffusion is relatively non-specific for pathology. Restricted diffusion is the earliest detectable MR sign of ischemia. However diffusion changes are also seen with infection, inflammation and neoplasms. In an abscess restricted diffusion and low ADC values are attributed to increased fluid viscosity and higher cellularity present in pus. Metastases and tumors may have a similar appearance to an abscess on morphologic images but present with normal diffusion and high ADC values in areas of necrosis.

DWI is gaining increasing importance in oncology both for the initial diagnosis and as a sensitive tool to assess tumor response to treatment. Low ADC values in a lesion are an indicator for a malignant compression fracture in the spine as well as for malignant lymphadenopathy in the neck. In cerebral gliomas ADC values inversely correlate with the grading of gliomas, and low values indicate the higher grade component in a ‘mixed’ glioma. Lower ADC values in the edema of gliomas compared to metastases are an early sign of brain infiltration beyond macroscopic visible margins. Restricted diffusion in the early postoperative phase is more likely to correspond to ischemia and reparative changes, while after 6–8 weeks may indicate true progression or pseudoprogression. Increasing ADC values in a solid lesion like a glioma, lymphoma or metastasis are a more sensitive...
parameter to chemo-radiotherapy treatment response than the contrast enhanced T1 images. Metastases and glial tumors can be differentiated by low ADC due to their high grade cellularity (metastases) and by areas of high ADC due to necrosis (glial tumors) (Figs. 2A–D).

**Perfusion-MRI (PWI = Perfusion-weighted imaging)**

Perfusion MRI relies on two different techniques: dynamic contrast enhanced (DCE) perfusion MRI consists of a T1w 3D FLASH sequence which is used to depict cerebral microcirculation and is a direct measure of vascular permeability. On the other hand dynamic susceptibility contrast (DSC) MR perfusion is based on a gradient-echo echo planar sequence. DSC provides a measure of vascularity, microvascular density or relative cerebral blood volume (rCBV) and therefore is complementary to DCE perfusion. Both sequences require a bolus of contrast media injected into a peripheral vein.

**Diffusion in glioma and metastasis: cellularity and tissue ultrastructure**

Contrast enhanced MRI does not reliably distinguish a glioblastoma multiforme (2A) from a metastasis (2B). Focal high ADC values (bright) indicate necrotic components in a glioma (arrow 2C). Low ADC values represent a solid metastasis (2B) as in this case or a lymphoma with dense cellularity (2D). ADC values are lower within the edema of a glioma due to microscopic infiltration which is usually not present within the edema of a metastasis.
Perfusion MR in combination with diffusion-weighted imaging has been routinely applied in the setting of acute stroke. The initial perfusion – diffusion mismatch correlates with the ischemic penumbra, the tissue at risk and the extent of the definite infarct size. In brain neoplasms vascular proliferation and neoangiogenesis are hallmarks of differentiation to higher grades of malignancy. rCBV values as obtained by DSC show a significant correlation with tumor grade (Figs. 3 A–C), microvascular density and in case of increase predict malignant transformation. Frequently, changes of these parameters during cytostatic, anti-angiogenic and radiation therapy (Figs. 4A–C) precede tumor volume reduction. DCE is applied to assess the degree of tumor angiogenesis and vessel permeability. Dynamic contrast enhanced perfusion (DCE) in combination with dynamic susceptibility contrast (DSC) perfusion (Fig. 5B) and MR spectroscopy (Fig. 6C) is applied to distinguish treatment related effects (pseudoprogression or pseudoresponse) from true progression and true response respectively.

Perfusion and diffusion in brain tumor imaging: vascularisation, neo-angiogenesis, capillary permeability – cellularity vs. extracellular space

![Perfusion MRI of anaplastic glioma with high rCBV within the component of higher malignancy in the posterior part (arrow 3A) with restricted diffusion (3B) and low ADC values (3C) as additional signs of a highly cellular anaplastic component.](image)

**3** Perfusion MRI of an anaplastic glioma with high rCBV within the component of higher malignancy in the posterior part (arrow 3A) with restricted diffusion (3B) and low ADC values (3C) as additional signs of a highly cellular anaplastic component.

**Treatment monitoring: decreased vascularity**

![Cerebellar metastasis of renal cell carcinoma with subtle peripheral vessels indicated on the contrast enhanced T1w image. DSC perfusion MRI depicts markedly increased vascularity within the entire metastasis with high rCBV values before (4B) and with decreased rCBV values under radiotherapy (4C).](image)

**4** (4A) Cerebellar metastasis of renal cell carcinoma with subtle peripheral vessels indicated on the contrast enhanced T1w image. DSC perfusion MRI depicts markedly increased vascularity within the entire metastasis with high rCBV values before (4B) and with decreased rCBV values under radiotherapy (4C).
ASL Arterial Spin labelling

MR imaging with ASL is an alternative method to measure perfusion. Contrary to dynamic contrast enhanced (DCE) and dynamic susceptibility contrast (DSC) perfusion MRI, ASL uses electromagnetically labelled arterial blood water as a freely diffusible intrinsic tracer. In clinical applications, the ASL technique proved reproducible and reliable to assess cerebral blood flow (CBF) despite a relatively low signal intensity-to-noise ratio. ASL has been evaluated in various pathological states including cerebrovascular and neurodegenerative disease and for the assessment of glioma grading and tumor angiogenesis. Quantitative assessment of blood flow in gliomas by ASL yielded results and reproducibility comparable to DSC perfusion MR imaging.

MR Spectroscopy (MRS)

MR spectroscopy provides a measure of brain metabolic composition or chemistry. Each metabolite appears at a specific ppm (Fig. 6 A–C), and each one reflects specific cellular and biochemical processes. N-Acetyl-Aspartat (NAA) is a neuronal marker and decreases with any disease that adversely affects neuronal integrity. Creatine provides a measure of energy stores. Choline is a measure of increased cellular turnover and is elevated in tumors and inflammatory processes. Lactate is a marker of oxygen deficiency, lipids of tissue necrosis and myoinositol of granulation tissue (gliosis).

Indication of MRS:
- differential diagnosis of low-grade and high-grade tumors
- monitoring under radio-chemotherapy
- differentiation of recurrent tumor from secondary necrosis due to therapy

Susceptibility-Weighted Imaging (SWI)

SWI is a modified 3D gradient-echo technique which exploits the susceptibility differences between tissues and uses the phase image to detect these differences. The magnitude and phase data are combined to produce an enhanced contrast magnitude image which is exquisitely sensitive to products of blood-deterioration (DeoxyHB, MetHB and Hämosiderin) (Fig. 7 A–D).

Indications of SWI:
- micro-bleedings in hypertension and vascular dementia
- superficial bleedings in amyloid angiopathy
- hemorrhagic contusion and diffuse axonal injury after brain trauma
- iron deposition in neurodegenerativ diseases

Treatment related effects: low perfusion – low cellularity

5A T1w contrast enhanced image one year following resection and chemo-radiotherapy of a glioblastoma. Recurrence and treatment related necrosis cannot be reliably differentiated based on morphology (5A). Perfusion MRI depicts low rCBV values (5B) contrary to high rCBV values indicating recurrence in a high grade glioma (compare to Fig. 3A). High ADC values (5C) denote low cellularity indicative of a necrosis confirmed by subsequent histology.
MR Spectroscopy: tumor progression vs. pseudoprogression

6 (6A) Normal spectrum. (6B) Spectrum of an Oligodendroglioma with high Cholin as a marker of a high proliferation-rate (thick arrow) and with low NAA as a marker of lost neuronal integrity (thin arrow). (6C) Spectrum of a therapy-induced necrosis with low Cholin and NAA and with high Lactate as a sign of oxygen deficiency (thick arrow).

Iron deposition after trauma, in hypertension and in amyloid angiopathy

7 (7A) Circumscribed cortical gliosis (arrow) after brain injury without visible micro-bleeding on the T2-weighted image. (7B) Same patient with the same localization with well visible micro-bleedings (arrows) on SWI. (7C) Hypertensive small micro-bleedings in the basal ganglia and in the thalamus. (7D) Amyloid angiopathy with a large intraparenchymal bleeding on the left and with multiple superficial micro-bleedings on the right (arrows).

Advanced imaging in cerebrovascular diseases and vascular lesions

Contrast enhanced MR Angiography (MRA) is a well established complementary examination to Doppler sonography and in most cases replacement for digital subtraction angiography by producing high-quality static images.

Time resolved MR Angiography

The latest MRA technique is time resolved or 4D MRA with a time resolution of <0.7 s. This method is capable of capturing the dynamic filling of vessels, thus demonstrating the arterial, capillary and the venous phase of the cerebral circulation, similar to digital subtraction angiography (DSA). Hemodynamic changes caused by arterial stenoses and occlusions can be well
detected and dural arteriovenous fistulas (Figs. 8A, B) or arteriovenous malformations can be easily diagnosed. 4D MRA can contribute to the preoperative evaluation and characterization of a tumor by its degree of vascularization (Figs. 9A–F).

Indication of 4D MR Angiography
- hemodynamic changes due to an arterial stenosis or occlusion
- dural arteriovenous fistulas
- vascular malformations of the brain, face or neck

4D MRA of a dural arteriovenous fistula – high temporal resolution

Advanced MR imaging of inflammatory demyelinating diseases of the central nervous system

Advanced MR techniques have revolutionized the recognition and characterization of white matter disease. MR imaging is required to depict the presence and location of inflammatory lesions within the optic nerve, the brain and spinal cord. The morphology and distribution of lesions along the ventricular lining and perimedullary veins, in juxtacortical and infratentorial location may permit a tentative diagnosis. The acuity of lesions is assessed based on the presence of contrast enhancement or by alterations in the composition of neurometabolites in MR spectroscopy when large and unusual lesions or diffuse white matter changes are present. Follow-up examinations by high resolution MR sequences are of major importance to depict ‘dissemination in time’.

3D Fluid-Attenuated Inversion Recovery Sequence (FLAIR SPACE)

provides a higher spatial resolution with 0.8 to 1 mm isotropic slices and inherent contrast resolution in comparison to T2-weighted (Fig. 10A) and proton den-
The similar location, morphology and signal intensity of a paraganglioma (glomus tumor) and neurinoma (arrows: 9A, C) makes the differentiation sometimes challenging. The 4D MRA however, demonstrates the difference clearly: a paraganglioma shows an early and strong tumor filling (9B) and early washout in the venous phase (9C) whereas a vagal neurinoma (9D) exhibits no macroscopic vascularization in the arterial (9E) and venous (9F) phase.

3D Double Inversion-Recovery (DIR) sequence

is characterized by a high sensitivity to depict cortical (Fig. 10D) and deep grey matter involvement due to high signal of the lesions relative to the low white and grey matter signal intensity. 3D SPACE DIR sequence is equivalent to 3D FLAIR SPACE to depict white matter infra- and supratentorial lesions. The sequence appears slightly superior to recognize focal and generalized cortical grey matter volume loss in the vicinity of subcortical lesions and as a sign of neurodegeneration respectively.

4D MRA for demonstrating tumor perfusion: diagnostic and preoperative information

Paraganglioma

Schwannoma

9 The similar location, morphology and signal intensity of a paraganglioma (glomus tumor) and neurinoma (arrows: 9A, C) makes the differentiation sometimes challenging. The 4D MRA however, demonstrates the difference clearly: a paraganglioma shows an early and strong tumor filling (9B), and early washout in the venous phase (9C) whereas a vagal neurinoma (9D) exhibits no macroscopic vascularization in the arterial (9E) and venous (9F) phase.
Axial T2w (10A) and PD images (10B) hardly allow recognition of demyelinating lesions indicated by arrows within the forceps minor of the corpus callosum and in intracortical location within the subcentral gyrus. Images are acquired with 3.5 mm contiguous slices. Improved delineation of the corresponding lesions and significant more demyelinating plaques in the 3D FLAIR (10C) und 3D SPACE DIR sequence (10D) acquired at 1.0 and 1 mm slice thickness respectively.

Inflammatory demyelinating disease: new 3D sequences yield higher diagnostic accuracy, improved clinical and imaging correlation, and more precise follow-up assessment.

Contact
Prof. Bernhard Schuknecht, M.D.
Diagnostic, Vascular and Interventional Neuroradiology
Medizinisch Radiologisches Institut MRI Zürich
 Bahnhofplatz 3
 8001 Zürich
 Switzerland
image-solution@ggaweb.ch
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Imaging of Diseases of the Cranial Nerves: Tips and Tricks

Bernd F. Tomandl1; Norbert Sommer2; Patrick J. Egan1; Tibor C. Mitrovics1

1Dept. of Radiology and Neuroradiology, Christophshof Hospital, Göppingen, Germany
2Dept. of Neurology, Christophshof Hospital, Göppingen, Germany

Introduction

The intention of this article is to alert readers to common problems and pitfalls concerning magnetic resonance imaging (MRI) for pathologies of cranial nerves. It also provides an introduction to helpful sequences and post-processing techniques. There are numerous reports about imaging of the cranial nerves that show the capabilities of sub-millimeter heavily T2-weighted images, like the CISS or balanced FFE-sequences for the visualisation of the anatomy of the cranial nerves in the living body [1]. However, these sequences will only show the course of the cranial nerves within the basal cisterns, and whilst this is helpful in clinically-suspected cases of neurovascular compression symptoms [2, 3], in most other cases more information is needed to find the cause of cranial nerve palsy. To familiarise the reader with cranial nerve imaging, examples of the normal anatomy as well as typical pathological cases are shown in this article. Most images were acquired with a 1.5T MAGNETOM Avanto (Siemens Healthcare, Erlangen, Germany). All cited references are available online for free.

Anatomy and MR sequences

Imaging of diseases of the cranial nerves requires good knowledge of the course

1A

Frontal (1A) and rear (1B) view of the brain stem show the origin of the cranial nerves and their relation to surrounding arteries. The pictures were created from MR data. Figure courtesy of K. E. W. Eberhardt, Werneck, Germany and Peter Hastreiter, Erlangen, Germany.
Table 1: Course and function of the cranial nerves (CN).

<table>
<thead>
<tr>
<th>CN</th>
<th>Name</th>
<th>Course and Function</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>Olfactory nerves</td>
<td>Part of the brain. Responsible for the sense of smell. They run in the anterior base of the skull; their fibres exit the skull through the cribriform plate. These nerves are frequently injured in skull-base fractures.</td>
</tr>
<tr>
<td>II</td>
<td>Optic nerves</td>
<td>Part of the brain; surrounded by CSF and the dura. Responsible for the sense of vision. Frequent diseases affecting the optic nerves include multiple sclerosis and pituitary adenomas, as well as meningiomas and gliomas of the optic nerves.</td>
</tr>
<tr>
<td>III</td>
<td>Oculomotor nerves</td>
<td>Responsible for eye movement. The relatively large oculomotor nerves also control pupillary constriction. While CN III and VI are easily identified on CISS images, the trochlear nerve—being the only one of the cranial nerves to leave the brain stem at its back below the quadrigeminal plate—is often difficult to find due to its tiny size [11]. The course of the abducens nerves is interesting. They leave the brain stem below the pons, enter a duplication of the dura at the clivus (Dorello’s canal), and finally enter the cavernous sinus after crossing the petrosphenoidal ligament (Gruber’s ligament). Imaging of an affliction of an abducens nerve should therefore include contrast-enhanced T1w images of the clivus with fat saturation. Acute palsy of the oculomotor nerve is frequently (15%) related to an intracranial aneurysm of the internal carotid artery at the origin of the Pcom.</td>
</tr>
<tr>
<td>IV</td>
<td>Trochlear nerves</td>
<td></td>
</tr>
<tr>
<td>VI</td>
<td>Abducens nerves</td>
<td></td>
</tr>
<tr>
<td>V</td>
<td>Trigeminal nerves</td>
<td>They divide into three branches: the V1 (ophthalmic), V2 (maxillary) and V3 (mandibular) branch. They leave the skull through the superior orbital fissure (V1), the foramen rotundum (V2) and the foramen ovale (V3). They transmit sensations from the face, scalp and teeth, including chewing. Neurovascular compression of the nerve at the entrance zone to the brain stem may result in trigeminal neuralgia.</td>
</tr>
<tr>
<td>VII</td>
<td>Facial nerves</td>
<td>The facial and vestibulocochlear nerves both enter the internal auditory meatus. The facial nerves are responsible for the sensation of taste from the anterior 2/3 of the tongue as well as the motoric innervation of face muscles. Acoustic schwannomas and meningiomas are frequent pathologic findings affecting these nerves.</td>
</tr>
<tr>
<td>VIII</td>
<td>Vestibulocochlear nerves</td>
<td></td>
</tr>
<tr>
<td>IX</td>
<td>Glossopharyngeal nerves</td>
<td>The glossopharyngeal, vagus and accessory nerves leave the skull through the jugular foramen. CN IX is also responsible for involuntary blood-pressure reflexes, cardiac and respiratory sensing, contraction of the pharynx and the swallowing reflex. Some of these signals overlap with CN X, which is more involved in reflexes and vital functions. They can be involved in skull-base tumors or dissections of the carotid artery.</td>
</tr>
<tr>
<td>X</td>
<td>Vagus nerves</td>
<td></td>
</tr>
<tr>
<td>XI</td>
<td>Accessory nerves</td>
<td></td>
</tr>
<tr>
<td>XII</td>
<td>Hypoglossal nerves</td>
<td>The hypoglossal nerves exit the skull through the hypoglossal foramen. They control the movement of the tongue by innervating three of the four muscles. The fourth muscle is under the control of CN X.</td>
</tr>
</tbody>
</table>

Parts of table 1 are courtesy of Alicia Mae Prater, http://suite101.com/article/the-cranial-nerves-a105837.

of the particular involved cranial nerve. An overview of the cranial nerves is given in table 1 and figure 1. A more interactive and entertaining introduction to cranial nerve anatomy and function is given by Barbara Liang on the wisc-online site (http://www.wisc-online.com/Objects/ViewObject.aspx?ID=AP11504). Large cranial nerves are visible even on standard MR images: The ophthalmic, optical, trigeminal, oculomotoric as well as the facial and vestibulocochlear nerves can be readily identified on 4 mm T2w images (Fig. 2). The smaller nerves are more difficult to see and thin section images are required. As an example how the anatomic course of a cranial nerve influences the choice of MR sequences, the course of the sixth cranial nerves (abducens nerves) is as follows: The nerves leave the brain stem below the pons, enter a duplication of the dura at the clivus (Dorello’s canal), and finally enter the cavernous sinus after crossing the petrosphenoidal ligament (Gruber’s ligament) [4]. While CISS-images will only show the intracisternal course of the nerve (Fig. 3), other sequences are needed to exclude an infarction in the pons or a tumor along the course of the nerve. Imaging of an acute affliction of an abducens nerve should therefore always include diffusion-weighted...
How-I-do-it

2A 4 mm T2w TSE MRI. Some cranial nerves are always visible: (2A) T2w TSE with fat saturation in a patient with multiple sclerosis. Hyperintense right optic nerve indicating optic neuritis (white arrow). The left optic nerve is normal (arrowhead). In addition the olfactory nerves are visible (red arrows). (2B) The oculomotor nerves (arrows) crossing the space between the posterior cerebral artery and the superior cerebellar artery. (2C) The trigeminal nerves at their origin (red arrows) and the facial and vestibulocochlear nerves within the internal acoustic canal are clearly visible (white arrows).

3 Multiplanar reformations from 0.7 mm CISS images in oblique sagittal and coronal planes show the intracisternal course of the abducens nerves (yellow arrows) and the origin of Dorello’s canal (red arrows).

images (DWI) of the pons and (if no infarction is visible) contrast-enhanced T1w images of the clivus (sagittal plane) and cavernous sinus (coronal plane) with fat saturation [5]. Time-of-Flight (TOF) MR angiography (MRA) can be helpful to detect aneurysms in the cavernous sinus. Always remember that this technique is not sufficient to exclude intracranial aneurysms [6]. Often T1-weighted 3D-sequences after administration of gadolinium (e.g. Magnetization Prepared Rapid Gradient Echo (MPRAGE)) are very helpful as they allow for intensive post processing of the data (Fig. 4) [7].

Post processing

Usually the cranial nerves follow an oblique course through the basal cisterns. Therefore 3D post processing with multiplanar reconstructions (MPR) and/or volume rendering is often helpful to get a clearer delineation of the course of a specific cranial nerve and its vicinity.

The following case report will make that more clear:

A 46-year-old woman attended her physician after she developed acute ptosis and double vision especially when looking to the left side (Fig. 5). Clinical examination revealed an oculomotor palsy. The normal anatomy of the third
T1w MPRAGE after intravenous administration of Gadolinium-DTPA in a patient with multiple meningiomas and an assumed acoustic schwannoma. (4A) MPR in three orthogonal planes allows for optimal delineation of the tumor within the internal acoustic canal. In addition a volume rendered image shows the brain surface. (4B) Thin section MIP images (15 mm) and low opacity volume rendering allow good delineation of both the large intracranial arteries and veins.

Acute right oculomotor palsy in a 46-year-old woman. (5) Intracisternal course of the left oculomotor nerve (black arrow) demonstrated on a volume rendered image (6A) of the brainstem and an oblique sagittal view from 0.7 mm CISS data (6B). The nerve (black arrow) leaves the midbrain (*) between the posterior cerebral artery (white arrow) and the superior cerebellar artery (white arrowhead). After crossing the basal cisterns it enters the cavernous sinus.
How-I-do-it

The cranial nerve is shown in figure 6. MRI was performed including DWI and TOF-MRA without result. As the oculomotor palsy did not improve within 4 weeks the patient was sent to our hospital. She brought her images on a CD and we reviewed the images including post processing of the TOF-MRA that was done on a Siemens MultiModality workplace (Leonardo). While it was very difficult to see the lesion on the initially produced maximum intensity projection (MIP) reconstructions that included the whole volume data (Fig. 7) it was rather easy to detect the aneurysm on 15 mm thin MIP images in 3 planes and even easier on volume rendered images (Fig. 8). This is a good example why whole-volume MIP imaging is not very helpful if we want to see more than just the big arteries [8]. Of course it is mandatory to review the source images before any kind of 3D-imaging is done [9]. In our hospital we use routinely thin section MIPs of 15 mm section thickness in sagittal axial and oblique coronal planes where the coronal plane is reconstructed parallel to the basilar artery to get a clear visualization of the 2 vertebral arteries and the basilar artery and its branches (Fig. 9). Using this type of reconstruction makes it easy to delineate aneurysms in the vicinity of the cranial nerves. It is important to know that about 15% of acute oculomotor palsy cases are caused by intracranial aneurysms that are usually located at the distal intracranial internal carotid artery at the origin of the posterior communicating artery [10]. The course of the nerve roughly parallels the course of the posterior communicating artery (Fig. 6). It is very important to perform the MRA with high quality in these patients. We must be aware that TOF-MRA cannot exclude an aneurysm because slow flow within the aneurysm may lead to non-visualisation within the flow sensitive sequences so that other imaging modalities like contrast-enhanced MRA, CTA or even DSA are sometimes necessary.
References


Summary

Before performing MRI of a patient with cranial nerve palsy make sure that you know the course of the particular cranial nerves. Use thin section CISS-sequences to see the intracisternal course of the nerves. Use fat suppressed T1w images after contrast administration to visualize pathology within the skull base. Use 3D MPRAGE for a variety of reconstructions. Don’t rely on whole-volume MIP images from TOF-MRA. Routinely use the excellent post-processing tools that come with all Siemens scanners, such as MPR, thin section MIPs and volume rendering. This will ensure that you will not miss important findings.
3T DTI in Patients with Glaucoma. New Approaches for Data Analysis & Clinical Implications

Tobias Engelhorn1; Georg Michelson2; Joachim Hornegger3; Arnd Doerfler1

1Department of Neuroradiology, University Erlangen-Nuremberg, Germany, 2Department of Ophthalmology, University Erlangen-Nuremberg, Germany, 3Department of Computer Science, University Erlangen-Germany, Germany

Background
Glaucoma is responsible for approximately 10% of cases of blindness throughout the world and thus the third leading cause of blindness with over 8 million cases each year [1]. Glaucoma is considered a nervous system-based degenerative disease that is only partially influenced by ocular factors [2]. Moreover, neuronal degeneration involving all parts of the central visual pathways was documented at autopsy in patients with advanced glaucoma and severe visual field loss in both eyes [3]. Central neuronal degeneration can be assessed non-invasively using diffusion tensor imaging (DTI). To assess the optical pathways, DTI can exclusively depict the optic nerve (3rd neuron) and optic radiation (4th neuron) [4–6]. In addition, subtle changes can be assessed by calculation of fractional anisotropy (FA), which measures the orientation coherence of diffusion and provides information about axonal integrity [7]. The aim of the study presented was to evaluate whether 3T DTI with volumetric analysis of the optical pathway and calculation of diffusion coefficients can assess the (early) spread of glaucomatous damage within the central nervous system and to determine whether DTI-derived changes correlate with disease severity determined by established ophthalmological tests in patients with glaucoma. In addition, user-independent fast and robust assessment, new automatic segmentation approaches and a framework for a voxel-based morphometric analysis of the optic radiation were developed [8–14] and will be described.

Methods
The Clinical Investigation Ethics Committee of the University of Erlangen-Nuremberg approved the study protocol. Fifty patients (18 men, 32 women; mean age 52.2 ± 12.6 and 60.0 ± 16.9 years, respectively) diagnosed with damage of the optic nerve head or visual disturbance, were randomly selected for magnetic resonance imaging (MRI) and subsequent DTI. Fifty age-matched patients without glaucoma undergoing MRI because of headaches, dizziness, or stenosis of the nasolacrimal duct served as controls (22 men, 28 women; mean age 54.0 ± 14.2 and 61.4 ± 15.1 years, respectively). In these patients, increased intraocular pressure (IOP), optic nerve head atrophy or visual disturbances were excluded by an ophthalmological examination. Available additional examinations were referred to for evaluation, including Heidelberg Retinal Tomography, automated perimetry, spatial-temporal contrast sensitivity (frequency doubling test), nonmydriatic fundus images and optical coherence tomography (OCT) with measuring of the retinal nerve fiber layer (RNFL) thickness in glaucoma patients and controls.

Magnetic resonance imaging
MRI was performed using a 3T high-field scanner (MAGNETOM Trio with Tim; Siemens Healthcare AG, Erlangen, Germany) with a gradient field strength up to 45 mT/m (72 mT/m effective). Anatomical data were obtained in a T1-weighted three-dimensional magne-
tization-prepared rapid gradient-echo sequence (MPRAGE, TR 900 ms; TE 3 ms; FOV 23 x 23 cm; acquisition matrix size 512 x 256 reconstructed to 512 x 512; reconstructed axial plans with 1.2 mm slice thickness). DTI was performed in the axial plane with 4 mm slice thickness and no inter-slice separation using a single-shot, spin-echo, echoplanar imaging diffusion tensor sequence covering the whole visual pathway (TR 3400 ms; TE 93 ms; FOV 23 x 23 cm; acquisition matrix size 256 x 256; number of signal averages, 7; partial Fourier acquisition 60%). Diffusion weighting with a maximal b-factor of 1000 s/mm² was carried out along 15 icosahedral directions complemented by one scan with $b = 0$.

Data sets were automatically corrected for imaging distortions and coregistered.
in reference to T1-weighted MPRAGE images. These and further calculations, such as determining the independent elements of the diffusion tensor, deriving the corresponding eigenvalues and eigenvectors, and reconstructing and volume rendering fibers, were performed using dedicated software (Neuro 3D; Siemens Healthcare AG).

**Data analysis & postprocessing**

For manual segmentation by two experienced neuroradiologists (T.E. and A.D.) and quantification of the optic radiation, the seed regions for the tracking algorithm were selected on the coregistered T1-weighted images that were overlaid on the DTI data (Fig. 1). As a start region, an area consisting of approximately 12 to 18 voxels was chosen covering the LGN [5]. Therewith, fiber tracts emerging from the LGN into the primary visual cortex were investigated. The resulting fiber pathways were evaluated visually for integrity and accuracy of the reconstructed fibers (i.e. the course on the coregistered anatomic T1-weighted data was compared to the known anatomy of the visual pathway). In addition, volume rendering of the LGN and fiber tracts was performed by outlining the fourth neurons on each DTI slice by hand to calculate rarefaction.

**Automated segmentation of the optic fibers**

For automated segmentation and quantification of the optic radiation, a Mathcad-based software program was used. Sample-automated segmentation of the optic radiation on an axial brain slice. The left side shows the segmented optic radiation on a non-diffusion-weighted image (b = 0). The corresponding diffusion direction color-coded image weighted with FA is shown on the right side. The structure of the part of the visual pathway representing the main fiber bundle of the optic radiation and the LGN is clearly captured. From El-Rafei et al. Magn Reson Imaging 2011 [10].

Different regions-of-interest (ROI, red circles in the right visual pathways) that were used for the evaluation of fractional anisotropy in patients with glaucoma and controls in the left and right visual pathways: intraorbital part of the optic nerve (ON) (A, ROI 1), intracranial part of the ON (B, ROI 2), optic chiasm (C, ROI 3), lateral geniculate nucleus (D, ROI 4), and optic radiation (E & F, ROI 5–7). From Engelhorn et al. Acad Radiol 2012 [11].
(Parametric Technology Corporation, Needham, MA, USA) was developed (Figs. 2, 3). The program selects slices in which optical radiation is present by making use of the pattern recognition, head organ dimension (i.e., before actual image processing), location of the optic radiation in relation to the head size, and area of the largest green concentration (horizontal direction) as a starting point. Furthermore, the region-of-interest is selected for both sides. The green channel is extracted for the red-green-blue image and threshold operation in the green channel of every selected image. The white pixel represents green values above a computed threshold value, which is based on the image’s histogram. The optic radiation is segmented on the basis of the binarized image. The resulting area is calculated with a simple pixel per pixel summation. In the next step, the volume is calculated as area times slice thickness. To receive the overall optic radiation volume, the different slice volumes are summed.

**Qualitative analysis of the optic fibers using diffusion coefficients**

For qualitative assessment of the optic fibres, calculation of fractional anisotropy (FA) and radial diffusivity (RD) maps was performed with a dedicated software package (Neuro 3D; Siemens Healthcare AG). Regions of interest (ROIs) of approximately 9 mm² (range, 6–12 mm²) were depicted on the FA maps superimposed over the T1-weighted three-dimensional MPRAGE sequence described above. The following ipsilateral and contralateral ROIs were used for evaluation of FA (Fig. 4):

- **ROI 1:** the intraorbital part of the optic nerve directly after the eyeball
- **ROI 2:** the intracranial part of the optic nerve directly before the optic chiasm
- **ROI 3:** the lateral part of the optic chiasm
- **ROI 4:** the lateral geniculate nucleus (LGN)
- **ROI 5:** the optic radiation directly after the LGN
- **ROI 6:** the optic radiation at the level of the posterior horn of the lateral ventricle
- **ROI 7:** the optic radiation directly before its cortical spread

Statistical analyses were performed using PASW¹ release 18.0 (SPSS, Inc, Chicago, IL, USA).

**Results**

The automated segmentation and quantification of the optic radiation was feasible in all patients and controls. The evaluation matched 94% with manual evaluation for all patients and Cronbach’s α was > 0.81 for calculation of the manually and semiautomatically derived volumes. The mean evaluation time of semiautomatic segmentation including the time for data transfer was significantly shorter (38 minutes) compared to manual segmentation (91 minutes).

As result of DTI, age-related atrophy of up to 30% of the optic radiation was detected in healthy patients (Fig. 5) and there was high correlation between a decrease in the volume of the fourth neuron of both sides with increasing age ($r = 0.82$) (Fig. 5).

44% of glaucoma patients had at least a slight rarefaction of the optic radiation, i.e., averaged rendered volume was < 85% compared to controls. In these patients, the averaged rendered volume of the optic radiation of both sides was reduced to $67 \pm 16\%$ compared to controls (Fig. 6).

Measurement of FA in ipsilateral and contralateral ROIs was feasible in all test patients. A comparison of age-related rarefaction of the optic radiation (fourth neuron) in glaucoma patients and healthy subjects is shown (Fig. 6).

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¹ PASW: Predictive Analytical Software, a trademark of SPSS, Inc.
### Table 1

<table>
<thead>
<tr>
<th>Localization</th>
<th>Patients with Glaucoma</th>
<th>Controls</th>
</tr>
</thead>
<tbody>
<tr>
<td>ROI 1</td>
<td>0.36 ± 0.11</td>
<td>0.42 ± 0.13</td>
</tr>
<tr>
<td>ROI 2</td>
<td>0.48 ± 0.15*</td>
<td>0.66 ± 0.12</td>
</tr>
<tr>
<td>ROI 3</td>
<td>0.32 ± 0.16</td>
<td>0.35 ± 0.17</td>
</tr>
<tr>
<td>ROI 4</td>
<td>0.21 ± 0.09</td>
<td>0.24 ± 0.07</td>
</tr>
<tr>
<td>ROI 5</td>
<td>0.40 ± 0.16*</td>
<td>0.57 ± 0.13</td>
</tr>
<tr>
<td>ROI 6</td>
<td>0.48 ± 0.17</td>
<td>0.64 ± 0.11</td>
</tr>
<tr>
<td>ROI 7</td>
<td>0.44 ± 0.22*</td>
<td>0.53 ± 0.20</td>
</tr>
</tbody>
</table>

FA: fractional anisotropy; ROI = region-of-interest. *Significantly lower FA values compared to controls (p-value < 0.05). From Engelhorn et al. Acad Radiol 2012 [11].

---

Schematic of the analysis framework according to El-Rafei et al. (6). The system analyzes the diffusion tensor images of the optic radiation to produce localization maps showing regions with significant differences between glaucoma and control groups. The schematic illustrates the different steps including optic radiation identification and configuration, registration and statistical analysis.

Subjects. Mean evaluation time was 32 minutes per patient. Cronbach’s α at the 95% confidence interval for FA was 0.990 (right side) and 0.886 (left side). In controls, there was high correlation between a decrease in FA of both sides with increasing age (r = 0.78). The overall percentage change in FA in control subjects between 45 and 83 years of age was 33.7 ± 10.7%. This difference was significant (P < 0.05). Right-side and left-side FA measurements in controls and glaucoma patients were similar and differed only in small ranges up to 5%. Hence, the measurements for both sides were averaged and are displayed in Figure 7.

FA was significantly reduced in the intra-cranial part of the optic nerve (3rd neuron of the visual pathways) and in all three sections of the optic radiation (4th neuron of the visual pathways), thus in all sections with a more or less straight axonal direction. In contrast, there was no significant decrease (but a slight tendency toward a decrease) in FA in the intraorbital optic nerve, the optic chiasm, and the LGN. The explanation for this finding is most likely the crossing of axons in the optic chiasm and the reconnection of axons in the LGN, resulting in a change of the straight axonal direction with subsequent decrease in FA [4, 15], whereby ineluctably, movement of the eyeball subsequently results in movement of the intraorbital optic nerve, downgrading the quality of FA analysis in this section of the visual pathways by motion artifacts. This limitation could be overcome with retrobulbar anesthesia if needed.

In addition, we could establish a framework for automatic voxel-based morphometric analysis of the optic radiation using DTI (Figs. 8, 9) that can be also used in other forms of neurodegenerative brain disease: glaucoma patients have increased radial diffusivity (RD) and mean diffusivity (MD) significant voxels with a main concentration in the proximal part of the optic radiation. The proposed analysis provides a robust framework to capture the significant local changes of the optic radiation due to glaucoma.

Correlation between MRI and ophthalmic examination
Correlations were found for the presence/absence of rarefied optic radiation and the stage of optic nerve atrophy (Fig. 10). One-tailed analysis of data revealed a correlation of the stage of optic nerve atrophy with the presence of rarefied optic radiation (Kendall tau-b = 0.272; P < 0.05). The homonymous visual field defect scores were related to the corresponding rarefaction of the optic radiation (P < 0.05). Analogously to volumetric analysis, there was high correlation between decrease in FA measured and the extent of optic nerve atrophy and spatiotemporal contrast sensitivity of the retina (r > 0.81). Aside, there was high correlation between atrophy of the retinal fiber layer measured with optical coherence tomography (Fig. 11) and a decrease in FA and an increase in RD (Fig. 12).
68-year-old female patient, OD/OS with primary open angle glaucoma, in OS a parapapillary bleeding of the optic nerve head. (10A) In both eyes (OS [left] > OD [right]) the automated perimeter showed predominantly superior visual field defects due to a loss of axons of the 3rd neuron. (10B) The frequency doubling test indicated impaired spatial-temporal contrast sensitivity primarily in OS in the superior and temporal area as well as nasal near the center. (10C) Typical signs of glaucomatous optic nerve atrophy were recorded by a non-mydriatic fundus camera that is in OS a small rim area, smaller inferior rim than temporal and a parapapillary bleeding (arrows). (10D) DTI shows intact optic radiation in a healthy 67-year-old woman without any visual disturbances. (10E) DTI reveals significant rarefication of the optic radiation compared to the age-matched control (arrows). DTI, diffusion tensor imaging; OD, right eye; OS, left eye. From Engelhorn et al. Acad Radiol 2011 [9].
11 49-year-old male patient with primary open angle glaucoma and significant reduction of optic radiation (11A). Optical coherence tomography (OCT) demonstrates correlating reduction of retinal nerve fiber layer (RNFL) of both eyes, i.e. the 1st and 2nd neurons of the optic pathways (11B, C).

12 Correlation of averaged fractional anisotropy (FA) and radial diffusivity (RD) within the optic radiation of glaucoma patients with OCT-derived retinal fiber layer (RNFL) thickness.RNFL reduction results in decrease of FA and increase of RD.
Conclusion and clinical implications

Using DTI at 3 Tesla, the intracranial optic fibers were assessed in vivo non-invasively by volumetric analysis and by calculation of diffusion coefficients within the optic nerve and the optic radiation.

Based on volumetric analysis, we could demonstrate that
1) there is ongoing physiological atrophy of the optic radiation with increasing age and
2) that there is significant pronounced atrophy of the optic radiation in glaucoma patients with
3) good correlation to the extent of optic nerve atrophy whereas morphological MRI sequences do not show pathological findings.

Aside, FA measurement in different sections of the visual pathways is feasible, and
4) there is also significant physiological decrease in FA with increasing age;
5) compared to age-matched controls, FA is significantly decreased in the intracranial part of the optic nerve and in the optic radiation in glaucoma patients; and
6) there is a high correlation between the decrease in FA and the severity of optic nerve atrophy and retinal impairment. In contrast to volumetric analysis, there is a significant change in FA even at the very early stage of glaucoma.

Hence, DTI with calculation of diffusion coefficients is a sensitive tool for early diagnosis of glaucoma and therapy monitoring. In addition 7) the approach for automatic segmentation of the optic radiation and analyzing diffusion coefficients seems to be robust and is clearly faster compared to manual segmentation.

References


Contact

Prof. Tobias Engelhorn, M.D.
Department of Neuroradiology
University Hospital Erlangen
Schwabachanlage 6
D-91054 Erlangen
Germany
Phone: +49-9131-8544824
Fax: +49-9131-8536179
 tobias.engelhorn@uk-erlangen.de

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Case Study: Murray Valley Encephalitis (MVE)

Jan Neal

Benson Radiology, Adelaide, Australia

Introduction

Murray Valley Encephalitis (MVE) is caused by infection with a flavivirus belonging to the Japanese encephalitis (JE) antigenic complex, which also includes St. Louis encephalitis (StLE) and West Nile (WN) virus. The MVE virus occurs in Australia, New Guinea, and probably islands in the eastern part of Indonesia [1]. MVE virus is believed to be maintained in a natural cycle involving water birds and Culex annulirostris mosquitoes. The natural transmission cycle of the JE group involves infection of a mosquito vector alternating with viral amplification in a variety of vertebrates. Human disease is incidental to this cycle. Only one in 1,000 to 2,000 infections results in clinical illness resembling JE [1]. They are neurotropic viruses, which cause illness with headache, fever, vomiting followed by drowsiness, mental confusion and in severe cases there may be hyperactive reflexes, spastic paresis, seizures, coma and death. These viruses are becoming increasingly important globally as their geographic ranges steadily increase [2].

MR imaging (MRI) is more sensitive than computed tomography (CT) for detecting MVE-associated abnormalities such as changes in the thalamus, basal ganglia, midbrain, pons and medulla. When clinically correlated these changes can be specific to JE but not very sensitive. Mortality among hospitalized patients is about 20–30%. About half of the survivors have residual neurological deficits [2]. There is no vaccine for MVE virus. Prevention relies on mosquito control and avoidance of mosquito bites. There have been no human cases of MVE infection in South Australia for over 30 years.

1 T2-weighted axial images. Abnormal bilateral and symmetric hyperintense signal to central deep grey matter structures including the thalami, substantia nigra, dorsal midbrain and pons.
FLAIR coronal images showing abnormal hyperintense signal as seen in Figure 1. T1w images show corresponding hypointense signal.

Diffusion b1000 trace images (3A) and ADC images (3B). These images do not show restricted diffusion. Therefore this is not an acute infarct.
Patient history
A 26-year-old male was admitted to intensive care unit with respiratory failure. MRI head is requested.

Sequence details
MRI examination was performed on a Siemens 1.5T MAGNETOM Avanto system, using the head coil. Standard pre and post contrast sequences were acquired. Pre contrast PD/T2-weighted axial, T1w axial, T1w sagittal, FLAIR coronal, diffusion-weighted scans, MR venography and post contrast T1w axial and coronal images were acquired.
Imaging findings

MRI of the brain showed grossly abnormal appearances of the thalami, substantia nigra and dorsal aspect of the brainstem. Given the lack of restricted diffusion and no enhancement a tumor or Herpes Simplex virus (HSV) encephalitis were ruled out. Remainder of the brain was normal. Differential diagnosis would include metabolic processes (osmotic myelinolysis). Correlating to the patient’s clinical history it was not felt likely in this case. So an underlying infective aetiology particularly flavivirus infection such as MVE is favored. [3]

Conclusion

MRI is an excellent clinical tool to aid in the diagnosis of MVE. The classic MR appearance of this virus can assist in expediting the diagnosis. Blood testing must be done in interstate laboratories and can take some time for results to be returned.

References

3 From the radiology report by Dr. S. Knox, Consultant Radiologist, Benson Radiology.

Contact

Jan Neal, Grad Dip MRT
Benson Radiology
Adelaide
South Australia
jan.neal@bensonradiology.com.au

After 5 weeks the patient remained ventilated with no signs of neurological improvement. Repeat MRI Brain and c. spine performed. This showed no improvement, there has, in fact, been significant progression of gross abnormal signal intensity with new involvement of the globus pallidus and medial temp lobe.
Clinical Neurology

Neuroimaging of Stroke. The Complementary Roles of CT and MRI

Karl-Olof Lövblad; Vitor Mendes Pereira

Hôpitaux Universitaires de Genève, Department of Diagnostic and Interventional Neuroradiology, Geneva, Switzerland

Abstract

Cerebral ischemia or stroke is now considered a treatable medical emergency. This has created the need for powerful diagnostic tools for its determination with accuracy. Computed tomography (CT) and Magnetic resonance imaging (MRI) have known great technological improvements that have paralleled the clinical successes with treatment modalities. While overall equivalent, both techniques have their pros and cons and tend to be more and more accepted as complimentary techniques that may be used either alone or sequentially depending on the question that is being asked. This paper deals with their use alone and in combination for the diagnosis.

Introduction

Cerebral ischemia and its resultant pathology, stroke, is one of the leading causes of morbidity and mortality in industrialized countries. This is in part due to the fact that it shares common causes with cardiovascular diseases such as arteriosclerosis among others. While stroke was for a long time considered an untreatable entity for which there was no treatment and where the only outcome at best was a long and difficult period in rehabilitation, over the last two decades, parallel advances in both therapies and imaging have provided the clinician with new options when confronted with these patients. The main objective when confronted with a patient who has acute signs referable to a possible stroke is to exclude another possible pathology that could cause identical symptoms [1].

In the emergency situation, imaging has to perform many roles while being short and not interfering with the time window: it has to exclude another pathology (primarily hemorrhage), detect ischemia, demonstrate occlusion, show hypoperfusion and eventual penumbra and finally determine hemorrhagic risk and eventual outcome. Since its introduction in the 70’s, computed tomography has always played an important role in the management of patients with neurological diseases [2, 3]. Initially for patients with acute symptoms it was mainly used to exclude a very clear cause of neurological dysfunction such as cerebral hemorrhage. However, as knowledge and experience with the use of CT has increased, it has proved itself to be a very powerful tool in experienced hands (or eyes) [4]: one can see early signs of loss of differentiation between white and grey matter as well as signs of sulcal effacement that will eventually precede hypodensity; hypodensity will correspond to the accumulation and increase in water content in the tissue. Frank hypodensity when encompassing more than 33% of the MCA territory

1A Patient with left-sided weakness: there is a right-sided hyperintensity on the DWI image corresponding to ischemia (1A) along with a decrease in the ADC (1B), as well as hypoperfusion (1C). The difference between the diffusion lesion and the perfusion area corresponds to the penumbra.
was found to be a contra-indication to the administration of thrombolysis since there was a subsequent increase in mortality [5].

After the publication of the initial NINDS thrombolysis trial [6], as well as the ECAS trial [7], intravenous thrombolysis slowly became accepted as a standard of care for patients with acute stroke. It had the main drawback that therapy was initially considered to be effective only 3 hours after symptom onset (considered to be the ‘therapeutic window’). This had two impacts. The first was to look for a way of not just diagnosing but of certifying that one was treating a patient with acute stroke: thus began a new era for neuroimaging to become a marker for acute stroke. The second was to search for more effective drugs and to try to expand the therapeutic window.

Recent evaluation of thrombolysis data has shown that for rtPA at least the therapeutic window is now 4.5 hours [8]. Also there is mounting evidence that mechanical non pharmacological may be the preferred tool for drug trial where very often an accurate assessment of ischemic lesion volume is absolutely necessary in order to assess drug safety and efficacy [19].

After years of debate and controversy, it has finally been established that MRI can demonstrate hemorrhage probably as well as CT and is better at demonstrating ischemic lesions with DWI (Fig. 2). The penumbra concept, which has evolved substantially from its initial description, has been a cause of some controversy. Indeed, it was first a situation in which neuronal dysfunction was caused by decreased blood flow and thus cellular damage [20]. Since it was somewhat unclear in its scope and exact definition the area was called the penumbra. From this initial description of this entity, the clinical concept has evolved towards an imaging-driven one where the penumbra is seen as an area of hemodynamically impaired brain and not one of synaptic dysfunction. However since it is the hemodynamics that are at its basis, both kinds of penumbra – overall, or at least from a clinical point-of-view – can be considered the same in order to have a working hypothesis towards implementation of treatment. It is for this reason also that one prefers to speak about tissue at risk and not just about penumbra.

The imaging revolution

During the 90’s the first real revolution was the clinical implementation of echoplanar imaging capable scanners [10] that would thus allow diffusion [11] and perfusion imaging [12] techniques to move into the clinical arena. This led to the capacity of performing whole brain datasets demonstrating both diffusion lesion and perfusion deficit [13] (Fig. 1). Diffusion imaging relies on the capacity of a diffusion sequence to demonstrate changes in water distribution between the intra- and extra-cellular compartments that may occur in stroke [14]: on the so-called diffusion-weighted images (DWI) this is associated with a signal increase that can be quantified by assessing the maps of the apparent diffusion coefficient (ADC) that are usually generated at the same time. Initially this led to great enthusiasm due to the high sensitivity [15] and specificity of DWI for stroke and it was even hoped that it might constitute the first step towards the equivalent of an ECG for cerebral ischemia [16]. A very simple MR protocol consists of a T2-weighted image, a T2*-weighted image a Time-of-flight (TOF) MR angiography (MRA) sequence and the diffusion and perfusion sequences, all of which can be done in 20 minutes or less. Computed tomography had also made some advances since its debut with the development of spiral scanning [16]. While this was initially applied to do examinations of the chest and abdomen, it set the standard for future potential uses in imaging of the neck vessels. Here it was the additional development of multi-slice scanners that allowed one to cover larger and larger areas in shorter times, thus enabling both implement accurate angio-CT of the head and neck vessels as well a brain CT perfusion imaging [17].

Due to a certain facility of use, CT became the leading technique. Indeed, along with some difficulties in its use (magnetic fields, claustrophobia, many sequences), MRI was also considered more difficult to read when it came to the detection of hemorrhage in inexperienced hands. Also, since its use is slightly less time consuming, it has been widely adopted for screening patients with acute stroke that are enrolled in clinical trials or even in clinical routine of thrombolysis protocols.

However, magnetic resonance imaging, due to its capacity to provide a better visual delineation of acute and especially small infarcts [18] (Fig. 2), has become a preferred tool for drug trial where very often an accurate assessment of ischemic lesion volume is absolutely necessary in order to assess drug safety and efficacy [19].

Also there is mounting evidence that mechanical non pharmacological may be the preferred tool for drug trial where very often an accurate assessment of ischemic lesion volume is absolutely necessary in order to assess drug safety and efficacy [19].

After years of debate and controversy, it has finally been established that MRI can demonstrate hemorrhage probably as well as CT and is better at demonstrating ischemic lesions with DWI (Fig. 2). The penumbra concept, which has evolved substantially from its initial description, has been a cause of some controversy. Indeed, it was first a situation in which neuronal dysfunction was caused by decreased blood flow and thus cellular damage [20]. Since it was somewhat unclear in its scope and exact definition the area was called the penumbra. From this initial description of this entity, the clinical concept has evolved towards an imaging-driven one where the penumbra is seen as an area of hemodynamically impaired brain and not one of synaptic dysfunction. However since it is the hemodynamics that are at its basis, both kinds of penumbra – overall, or at least from a clinical point-of-view – can be considered the same in order to have a working hypothesis towards implementation of treatment. It is for this reason also that one prefers to speak about tissue at risk and not just about penumbra.
Initially the MR penumbra concept was based on a simple visual subtraction between the hypoperfused area and the smaller central initial DWI lesion. While rather imprecise and probably inexact, it provided the clinicians with an initially functioning model to use to monitor disease and interventions.

When considering the ischemic brain, one point that has always been considered elusive, at least with CT and MRI, has been the capacity to detect collateral flow. Indeed, since from a physiological point-of-view, a drop in blood flow is very quickly followed by neuronal death, perfusion is maintained at least by a collateral system of present. This is an observation that is very often made by interventional neuroradiologists, and an area where planar (CT and MRI) imaging has failed somewhat. However it is becoming increasingly evident that at least a partial image of collateral flow can be seen on post-contrast CT images where very often dilated collateral vessels will appear or on MRI where arterial spin labeling (ASL) seems to be able to demonstrate at least in some instances the presence of collateral circulation. This may be especially relevant in cases that arrive at the hospital beyond the therapeutic window: indeed, in any case, based on imaging there may still be a potential tissue at risk, or little signs of secondary hemorrhage so that these patients may indeed benefit from the use of a non-pharmacological intervention. The typical case will be a patient arriving with symptoms after 5 hours where intravenous rTPA is thus contraindicated but where MRI reveals only a small lesion with a surrounding hypoperfusion: this candidate may be amenable to clot extraction and/or stenting based on imaging findings.

The strengths of CT and MRI

A strength of magnetic resonance imaging is clearly its capacity to demonstrate even small lesions that will be often invisible or only visible to the trained eye.

Magnetic resonance provides whole brain coverage with both diffusion and perfusion sequences; additional techniques such as ASL and susceptibility-weighted imaging (SWI) can provide unique information on the ischemic brain. Very often MRI can provide a diagnostic result in patients where the initial CT is not conclusive (Fig. 3). MRI also remains non-ionizing, which is of prime importance if one considers patients that require frequent follow-ups. This is why, even in patients who have had an initial CT, MRI is to be preferred for follow-up: on the second day DWI will be able to demonstrate the exact extent of the lesion and thus allow one to obtain a better delineation of the final lesion extent (Fig. 4). As already explained, arterial spin labeling is a technique using labeled blood as an endogenous tracer: this has been shown to produce images of cerebral blood flow but also shown able – at least in part – to obtain images containing information about collateral flow [21] (Fig. 5); this could be of advantage.

Patient with acute posterior fossa symptoms. The CT is not conclusive and does not show any clear signs of hypodensity (3A, B) while there is non-visualization of the basilar artery (3B). The diffusion image (3C) shows a clear ischemia in the central pons but no signs of hemorrhage on the T2*-weighted image (3D). Angio-MRI confirms occlusion of the basilar artery (3E). Based on the presence of ischemia and occlusion but no hemorrhage it was decided to go to angiography and intervention (3F). After intervention there was reperfusion of the basilar artery (3G).
in cases of strokes again beyond the therapeutic window and where the outcome could be improved by some kind of treatment. SWI, due to its inherently strong T2* weighting, could help to demonstrate the well known venous stasis that is present in severe cases of stroke [22]; it can also demonstrate small bleeds better, and there have been some instances where it has been shown to demonstrate changes referable to the altered oxygenation in these tissues [23] (Fig. 6). The use of higher fields such as 3T has enabled an improvement in perfusion techniques and the implementation of SWI and ASL clinically [24]. In addition, new high-resolution angiography techniques should allow one to investigate the vessels both intra- and extracranially (Figs. 7 and 8).

A CT is very often more accessible since there is no need to screen often unconscious or uncooperative patients for metallic implants; the room can also be entered more quickly by an emergency team that does not have to remove pens, scissors and wallets from their pockets...
before placing the patient on the table. Also, until recently the gantries of CT scanners gave the team a better view of the often unstable and sometimes intubated patients so that they could be monitored more safely during the examination.

The typical situation where CT and MRI are both going to be necessary are the cases of less acute situations, where an initial CT and CT angiography (CTA) shows a lesion maybe not amenable to thrombolysis but which necessitates a more thorough work-up because of the underlying disease. This is the case in carotid disease, where patients either present with a stroke or with stroke-like symptoms. The initial CT may initially be normal despite the patient having certain clinical signs of acute stroke: in these cases going to MRI can be very helpful. This will be the case for small cortical lesions. It will typically be the case in patients with severe carotid stenosis, where CT has often been done since CTA will provide the best delineation of vascular calcification, but where MRI can provide two additional findings: the demonstration of small ischemic emboli in the brain and local demonstration of the presence of plaque hemorrhage (Figs. 9 and 10).

**Future perspectives**

Whenever one considers the etiologies of stroke, local carotid plaque vulnerability is going to be a key factor. Thus, the idea of performing any kind of ‘molecular’ imaging that would allow to determine which plaque is more ‘active’ and more prone to eventually seed emboli is very seductive. This could be done either by MRI using either SPIO contrast materials, or even by conventional contrast media or some combination of a nuclear medicine technique with a radiological method (CT or MRI) to produce combined hybrid imaging.

**Conclusions**

From a radiological point of view we are now at a crossroads where we deploy two advanced techniques, CT and MRI. These techniques are equally useful since they have complementary pros and cons: MRI is more sensitive to ischemia, but CT can more clearly demonstrate hemorrhage; CT angiography may provide a more correct lumino- graphic effect and demonstrate plaque calcification better. Both techniques can be utilized in first intention with almost comparable results, but this depends often on local logistics. However, in complex vascular situations it becomes increasingly clear that one needs both
Patient with an intracranial stenosis and a frontal opercular infarction on the left. The DWI image shows a left frontal lesion (7A) and the high resolution MRA images show the stenosis (arrows).

Patient with a tight stenosis of the left ICA, as well as cortical lesions (8A–C). The carotid artery is calcified on CTA (8D, E) and on MRI there is a hemorrhage visible in the plaque (8F–H).
9A Patient with a high-grade carotid stenosis on the right, seen on CTA. On the diffusion images one can clearly see the many ischemic cortical lesions (9A–D) that can only be seen retrospectively as a slight cortical swelling on the CT (9F).

10 Patient with acute stroke. There is hypoperfusion on the perfusion CT but not clearly any new lesion visible (10A). On T2 there are extensive alterations in the brain parenchyma (10C). On DWI one can see that there is a recent ischemic cortical area that explains the new symptoms (10D). Angiography revealed a carotid stenosis (10E) and CT after angiography showed luxury perfusion in the ischemic territory with contrast accumulation but no blood (10F).
the information from the CT, which can better exclude hemorrhage and demonstrate vascular calcifications, and from the MRI, which has an inherently higher contrast to demonstrate small cortical lesions that will explain the patient’s status.

Magnetic resonance is also mandatory for follow-up examinations: not only does it allow one to determine exact lesion extent, but it will also help determine prognosis based on this finding. Indeed, it has been demonstrated that both acute and late ischemic lesion volumes do correlate with clinical status [25].

The progresses and improvements in imaging that have been made in the last two decades have been enormous and have allowed us not just to improve diagnostic quality but also to gain insights into the disease processes that are ongoing so that therapy can be planned optimally and the follow-up can be as precise as possible. Thus, any stroke service should not rely only on CT or only MRI but use both in an optimized and optimal way in order to improve patient outcomes and safety.

References


Implication of Brain Susceptibility-Weighted Imaging

Yu-Kun Tsui1,3; Fong Y Tsai1,2; Chi-Yuan Chen1; Kwo-Whei Lee4

1Imaging Research Center, Taipei Medical University, Taipei, Taiwan.
2University California Irvine Medical Center, Orange, Ca. USA.
3Chi-Mei Medical Center, Tainan, Taiwan.
4Chang-Hua Christian Hospital, Chang-Hua, Taiwan.

(1A) SWI showed no high signal at left transverse sinus and (1B) no contrast in the left transverse sinus is seen on contrast enhanced MRI. (1C) MR venography showed poor visualization of left dural sinus, it cannot distinguish between atresia or thrombosis. (1D) Venous phase of DSA confirmed left transverse sinus thrombosis as seen in SWI. (1E, F) In another case contrast-enhanced T1-weighted MRI showed contrast-enhanced right transverse sinus. (1G) Coronal T2-weighted MRI showed iso-intensity signal in right sinus and (1H) low signal on SWI, these two findings are compatible with chronic thrombosis and confirmed by (1I) venous phase of DSA.
Introduction

Susceptibility-weighted imaging (SWI) is an MR imaging technique sensitive to magnetic susceptibility effects. It uses both magnitude and phase information from a high-spatial resolution three-dimensional GRE pulse sequence [1–3]. While phase post processing accentuates the magnetic properties of different substances, SWI is sensitive in detecting intravenous deoxygenated blood as well as extravascular blood products. It was originally referred to as high-resolution blood oxygen level-dependent (BOLD) venography [2, 4], but due to its broader application in detecting other diamagnetic or paramagnetic substances including iron, calcification, deoxygenated hemoglobin content etc., it is now referred to as SWI [1, 5]. SWI has been used in studies of vascular pathologies, trauma, arteriovenous malformation, tumor, multiple sclerosis, stroke, and other brain disease [6, 7]. The clinical application of SWI has been demonstrated in both adult and pediatric populations [8, 9].

The principles of SWI have been extensively described by Reichenbach and Haacke et al. [1, 10]. The sensitivity to susceptibility effects is maximized by using a long-TE high-resolution fully flow-compensated three-dimensional GRE sequence with filtered phase information in each voxel both to enhance the contrast in magnitude images and to add the susceptibility differences between tissues as a new source of information [1, 10–12]. After the imaging acquisition, the final processed SWI magnitude images are obtained after merging magnitude and phase images [10]. Due to combining both magnitude and phase information, the final processed SWI magnitude images can provide a benefit in detecting signal-intensity changes arising from both T2 and susceptibility differences between tissues. The processed SWI magnitude images can further generate into thick miniIP images to demonstrate vasculature, tortuous structures, and the continuity of vessels or lesions across slices. In processed SWI magnitude images, deoxyhemoglobin (deoxy-Hb), being paramagnetic with four unpaired electrons, generates magnetic fields that additively combine with the external magnetic field [2, 13], and presents as an intrinsic contrast agent appearing hypointensity in venous structure. On the other hand, arteries are hyperintense because of time-of-flight effects and lack of T2* effect [10, 14]. Therefore, the use of SWI in distinct and simultaneous evaluation of the arterial and venous systems of brain and related vascular pathologies is conceivable and possible and the application of SWI has been widely expanded [7, 15, 16, 31–36].

Vascular malformations

Several types of vascular malformations with slow or venous flow have been shown to be better visualized with SWI, including developmental venous anomaly (DVA), cerebral cavernous malformation (CCM), telangiectasia and Sturge-Weber Syndrome (SWS) [5, 6, 9]. SWI not only offers improved sensitivity but can also depict vascular structures that are invisible on conventional T2* GRE images.

DVA

DVA is the most common type of vascular malformation. DVA consists of radially arranged venous complexes converging to a centrally located venous trunk, which drains the normal brain parenchyma [17]. SWI can demonstrate the whole structure clearly and has been proven to provide better detection of venous structures than conventional T2* imaging. However, please note that in some DVA lesions with relatively higher blood flow, no obvious medullary veins of DVA may be shown on SWI [18].

CCM

CCM comprises 5%–13% of all the central nervous system vascular malformations [19]. CCM lesions can be detected in routine MR imaging when they have calcification or have previously bled. However, if CCM lesions are intact and have not bled, they may be almost invisible except for a faint or ill-defined nonspecific blush of enhancement after contrast administration. In addition, lack of flow-related signal intensity makes CCM undetectable using conventional MR angiographic techniques. Because of the possibility of blood stagnation phenomenon and chronic microhemorrhages, CCM lesions contain deoxy-Hb and hemosiderin and become very dark, thus easily detected on SWI, especially in the case of tiny lesions. The high degree of SWI sensitivity in assigning the number of CCM lesions is significantly superior to that of T2-weighted fast spin echo and GRE sequences [20].

Telangiectasia

Telangiectasia is a low-flow vascular malformation with low signal intensity on SWI and is typically small, ranging from several millimeters to 2 cm in size [21]. Telangiectasia lesions are mostly found in the pons, but are less well imaged with conventional MR sequences [22]. They may occur sporadically or may be associated with syndromes (e.g., hereditary hemorrhagic telangiectasia) or may occur as a result of endothelial injury, such as radiation-induced vascular injury, particularly in children who have received cranial irradiation [9]. SWI is a useful adjunct to conventional MR imaging in diagnosing telangiectasia.

Sturge-Weber syndrome (SWS)

SWS is a neurocutaneous disorder characterized by cutaneous angioma, glaucoma, and leptomeningeal venous angiomatosis. The leptomeningeal angiomatosis is associated with loss of normal cortical venous drainage results in abnormal venous drainage through the deep venous system, which may lead to progressive venous stasis and chronic hypoxia. The typical imaging findings include cerebral hemiatrophy and cortical ‘tram-track’ calcification resulting from chronic venous ischemia. The pial angioma is often easily seen with contrast-enhanced imaging. SWI can detect microstructural changes in cortex and white matter, as well as deep venous collaterals in children with SWS, and may be useful to objectively assess...
microstructural abnormalities at an early stage of SWS when interventions have the best chance to prevent irreversible neurocognitive sequelae [23].

Vascular pathologies with arteriovenous shunting
As previously mentioned in this article, venous structure are hypointense due to the presence of deoxyhemoglobin and arteries appear hyperintense due to time-of-flight effects and lack of T2* effect in processed SWI images. Therefore, abnormal hyperintense signal within the veins in processed SWI images may indicate the presence of lesion with high-flow arteriovenous shunting, such as arteriovenous malformation (AVM) and dural arteriovenous fistula (DAVF) [15, 16]. The accuracy of the SWI sequence in the detection of arteriovenous shunting can be further improved by performing post-contrast SWI studies [24].

Arteriovenous malformation (AVM)
AVM is a congenital vascular malformation that results in abnormal direct arteriovenous shunt. Imaging studies, including MR and intra-arterial digital subtraction angiography (DSA), are essential to identify and define exactly the different vascular components before embarking on therapy such as surgery, embolization, or radiosurgery. The use of SWI may be beneficial in the visualization and delineation of very small AVM lesions without hemorrhages and may help to detect these lesions at an early stage prior to bleeding [25]. SWI is also accurate for the detection of arteriovenous shunting in AVM lesions and may offer a noninvasive alternative to angiography in screening for, or follow-up of, treated lesions [15].

Dural arteriovenous fistula (DAVF)
The clinical features of DAVF range from headache, vertigo, or tinnitus to neurologic deficits or a life-threatening intracranial hemorrhage [26]. However, the clinical and neuroradiologic diagnosis of DAVF may be challenging. In processed SWI images, abnormally increased signal intensity in venous structures can result from high flow and rapid shunting of oxygenated blood via fistula. Due to the ability of SWI to detect and delineate intravascular deoxygenated blood and venous structures, as well as prominence of venous vasculature due to the prolonged passage time of intracranial blood, venous engorgement, and possible venous congestion caused by DAVF, can also be identified by SWI [16, 27]. The use of SWI may be helpful in early diagnosis and prompt treatment in DAVF.

Cerebral venous sinus thrombosis
Cerebral venous sinus thrombosis (CVST) is an unusual condition which is difficult to diagnose due to its variable modes of onset and wide spectrum of symptoms and signs [28]. In the present series, the major female predominance, age of onset, frequency of various onset and different clinical manifestation are those classically reported [29]. CVST usually presents nonspecific lesions, including hemorrhage, edema or infarction. Because of its sensitivity to susceptibility effects, SWI is of additional diagnostic value for clot detection in CVST in conjunction with conventional MR sequences and MRV, particularly in the acute phase of thrombosis in cortical CVST [30]. Because of the ability to detect signal-intensity changes coming from both T2 and susceptibility differences between tissues, processed SWI images can demonstrate not only the CVST, but also the extent of parenchymal edema and even hemorrhage that can occur after venous thrombosis leading to infarction. Furthermore, in some cases with chronic CVST, SWI should be incorporated in the evaluation of chronic occluded dural sinuses which often misleads as patent sinuses due to the enhancement in post-contrast images.

Stroke
Acute infarction with or without hemorrhage occurs due to thromboembolism, arterial stenosis or other entity. Acute thromboembolism accumulates high intracellular deoxy-Hb, which causes T2* shortening secondary to paramagnetic susceptibility differences making the thromboembolism visible on SWI. SWI can also add information about the acute stroke in detecting acute hemorrhage and in detecting cerebral amyloid angiopathy (CAA) with micro- or macro-hemorrhages. Therefore, SWI is useful in the assessment of acute stroke. The ability of SWI to detect hemorrhages may influence the subsequent treatment decision making for patients with acute stroke. To differentiate microbleeds and calcification in the infarct area, an analysis of phase image may be helpful. The susceptibility effects due to blood clots and blood stasis with deoxy-Hb detected in SWI can help locate the intra-arterial thromboembolism [31]. This may be useful in planning various treatment options and in assessing the extent of infarct [32]. In addition to its usefulness in the assessment of acute infarct, SWI also has the potential to predict stroke evolution. SWI can provide comparable information on mean transit time (MTT) and is an alternative to perfusion-weighted imaging (PWI) for the assessment of ischemic penumbra. The mismatch between SWI and diffusion-weighted imaging (DWI) can be a marker for ischemic penumbra and a predictor of stroke evolution [33]. The detection of deoxygenated blood and deep medullary veins in infarct region by SWI offers the possibility to assess tissue viability [9]. Because of the ability to demonstrate the abnormal visibility of transcerebral or deep medullary veins, SWI also has potential to evaluate the possibility of further hemorrhagic transformation in acute stroke patients treated by intravenous thrombolysis [34]. Therefore, SWI may be useful to select acute ischemic stroke patients for endovascular therapy. Further investigation may validate these implications of SWI for acute stroke.

There are two essences of imaging studies for acute stroke: speed and accuracy. Modified MR imaging including DWI, SWI and FLAIR, may be useful for acute stroke. However, the image acquisition may take too long for examining acute stroke patients due to motion from agi-
(2A) DWI showed high signals at left basal ganglia area. (2B) FLAIR showed small area of infarct at left mid-temporal cortex. (2C) SWI showed thrombus in left MCA.

(2D, E, F) SWI showed diffused low signals of left cerebral arterial distribution as extensive deoxygenated arteries.

(2G) DSA showed occlusion of left M-1. (2H) Micocatheter placed in left MCA for thrombolysis. (2I) DAS showed successful recanalization of left MCA.

(2J, K, L) Follow-up CT showed more infarctions with some hemorrhagic transformation.
T2-weighted MRI showed diffuse cerebral edema, optic nerve sheath swelling and transtentorial herniations, however, those findings cannot confirm imaging evidence of brain death. (3D–F) SWI showed extensive deoxygenated intracranial arteries and veins it indicates no intracranial circulation. (3G) CT angiography confirmed no intracranial circulation as imaging evidence of brain death and compatible with the findings of SWI.
tation and mental status changes. In our study, by adjusting the parameters of SWI including phase resolution, voxel size, matrix size and partial Fourier, reasonable diagnostic quality of SWI images can be obtained with shorter scanning time to 2 minutes and 14 seconds (with base resolution to 150 × 192 with voxel size of 1.5 × 1.2 × 2.0 mm and phase resolution 70%). This change can facilitate the application of SWI for acute stroke patients.

Mismatch PWI/DWI may indicate potential presence of penumbra, however, it cannot identify irreversible brain damage from ischemic stroke. Diffused deoxygenated middle cerebral artery on SWI indicates irreversible brain tissue with extensive oxygen extraction from severe ischemia. Recanalization was achieved, but the patient had progressive infarction (Figs. 2G–I). When imaging, SWI should incorporate evaluation for thrombolytic therapy to avoid complication and reduce the use of interventions in patients with very poor prognosis [33–35].

**Brain death**

In the case of patient with brain death, SWI show all intracranial vessels, including arteries and veins, as hypointensity and prominent cortical and deep cerebral veins. This change could reflect a combination of increased oxygen extraction, arterial and venous stasis, and/or possible venous dilation secondary to release substances after cell death. This finding most likely indicates a higher level of intravascular deoxygenated blood and diffuse hypoxia-ischemia change due to stopped blood flow with blood stasis, and may correspond to conventional angiographic and nuclear medicine imaging findings [36]. This initial experience of detecting prominent hypointense signals in intracranial arteries on SWI in patients with brain death may indicate absence of intracranial arterial circulation and thus may enable definitive diagnosis of brain death [36].

**Trauma**

**Traumatic brain injury (TBI)** is a major cause of morbidity, mortality and lost years of productive life throughout the world [37]. CT is the primary imaging modality for the assessment of TBI. However, TBI patients with **diffuse axonal injury** (DAI) often exhibit tiny or punctuate hemorrhages in the deep subcortical white matter and are not routinely depicted by brain CT or conventional MR imaging sequences. SWI is 3–6 times more sensitive than conventional T2*-weighted gradient-echo (T2*GE) sequence in detecting DAI hemorrhages [38], especially for lesions in the brainstem [6]. Furthermore, other traumatic intracranial hemorrhages can also be shown in SWI to establish the degree of injury more accurately.

**Fat embolism syndrome** (FES) is an uncommon but serious complication of traumatic injury and is frequently diagnostically challenging because first assessment by brain CT is usually normal. FES includes a triad: hypoxia, neurologic abnormalities, and petechial skin rashes [39]. In the brain, FES causes microinfarcts, vasogenic edema, and petechiae [40]. Conventional MR imaging has been reported to effectively visualize microinfarcts and vasogenic edema in FES in brain, but SWI can show diffuse microhemorrhages resulting from vascular injury due to toxic effect of local free fatty acid release or microinfarctions. In a case of DAI, the abnormal neurological features appear immediately post injury, whereas in FES neurologic abnormalities appear generally after orthopedic intervention or hours after trauma [41].

**Other implications for SWI**

**Assessment of brain tumor.** The advances of MR imaging have provided diagnostic, staging, morphologic, metabolic and functional information of brain tumor. The development of SWI
improves the ability to evaluate not only internal vascular architecture and hemorrhages, but also intratumoral calcification, which may not be detected in other MR sequences. These internal findings may be useful for diagnosis, differential diagnosis, and staging of brain tumors [42]. SWI can also be used in indentifying the hemorrhagic change or telangiectasia which occurs after radiation therapy [9]. SWI may assist to differentiate primary or secondary lesions by a deep medullary vein inside of tumor. It may be an important tool to diagnose the type of tumor.

Assessment of multiple sclerosis

Although the mechanism of iron accumulation in brain tissue is not yet clear, however, this phenomenon is observed in many neurodegenerative and inflammatory diseases, including multiple sclerosis (MS). Local accumulation of iron due to disruption of the blood-brain barrier and accumulation of iron-rich macrophages is found in MS brains [43]. SWI can indicate the lesions that may not be seen in conventional MR sequences because of its sensitivity in detecting iron deposition. Furthermore, because of the ability to present cerebral venous architecture, SWI can show the veins draining the MS lesions and also can help to diagnose chronic cerebrospinal venous insufficiency (CCVSI) in MS patients [44, 45].

Venous hypertension

Pseudotumor cerebri (PTC), or idiopathic intracranial hypertension, is a syndrome associated with multiple clinical conditions. The elevation of intracranial venous pressure may be a universal mechanism in PTC with different etiologies [46, 47]. This elevated venous pressure leads to elevation in CSF and intracranial pressure by resisting CSF absorption [47]. Sometimes these high pressures appear to be secondary to central venous hypertension but more often they seem to be the result of lesions obstructing cerebral venous outflow. The diagnosis of idiopathic intracranial hypertension is essentially one of exclusion of known causes of raised intracranial pressure [48]. CVST, in particular, must be excluded before idiopathic intracranial hypertension can be diagnosed [49]. In patients with the appropriate clinical phenotype of intracranial hypertension, SWI with conventional MR sequences can help to exclude the possibility of intracranial hypertension caused by CVST or other intracranial abnormality [7].

Hyperperfusion syndrome

Cerebral hyperperfusion syndrome (CHS) is characterized by ipsilateral headache, hypertension, seizures, and focal neurological deficits after carotid endarterectomy, carotid stenting, extracranial-intracranial bypass, or other revascularization procedures [50]. Intracranial hemorrhage will develop in up to 40% of these patients and has 36% mortality. The incidence of the syndrome varies, reflecting the range of mild to severe (intracranial hemorrhage) symptoms. MR imaging abnormalities include white-matter edema, focal infarction, and local or more overt hemorrhage. The hypersignal intensities along the involved cerebral hemisphere on SWI are suggestive of increased oxygenation and hyperperfusion. The possibility that SWI demonstrates increased oxygenated blood and perfusion in the areas of CHS may provide a rapid and convenient method to diagnose CHS [7]. Further investigation and study correlated with perfusion-weighted MR imaging, single-photon emission computed tomography (SPECT), or positron emission tomography (PET) are necessary to confirm the utility of SWI in detecting CHS.

Conclusion

The use of SWI to detect hemorrhage and vascular architecture has been documented and discussed in many studies.
SWI is useful in the improvement and provision of additional diagnostic information for intracranial diseases. SWI may expand to many other implications for brain pathology. Thus SWI should be a routine component of MRI studies of all cerebral disorders.

References

6A, B SWI showed high signals along left cerebral veins. (6C) showed left jugular vein stenosis. (6D) Venous phase of DSA showed left transverse sinus thrombosis. (6E) Venous pressure was still very high after thrombolysis and angioplasty of left jugular vein. (6F) showed stenosis was also seen at distal left jugular vein. (6G) Balloon angioplasty was performed. (6H) Partial dilatation of left distal jugular vein.
35 Tsai FY,Kao HW, Tsui, KY, Chen CY. Identifying irreversible brain tissue damage and penumbra. presented at 8th World Stroke Congress Oct 10-13th, Beasilia, Brazil.

Contact
Fong Y Tsai, M.D., FACR
Imaging Research Center
Taipei Medical University
250 Wunchi Street
Taipei,11031
Taiwan
fttsai@uci.edu or fttsai@tmu.edu.tw
Pseudoprogression and Pseudoresponse: Imaging Challenges in the Assessment of Post Treatment Glioma

L. Celso Hygino Cruz Jr.; Nina Ventura

Clinica de Diagnostico por Imagem (CDPI, DASA), Rio de Janeiro, Brazil

Glioblastoma multiforme is the most common primary malignant type of brain neoplasm in adults and is associated with a dismal prognosis. The current standard of care is surgical resection followed by radiation therapy (RT) and concomitant and adjuvant temozolomide (TMZ) chemotherapy [1]. Bevacizumab, an anti-vascular endothelial growth factor (VEGF) agent, has been recently approved for recurrent glioblastoma.

With the standardization of treatment around surgery/RT/TMZ and the current use of bevacizumab, certain patterns that were not previously noticed are beginning to emerge. These changes in magnetic resonance (MR) imaging can

1 Pseudoprogression. 59-year-old male, GBM treated with surgery followed by chemotherapy and radiotherapy. Just one month after finishing the treatment the lesion increased in size. The chemotherapy was continued and the lesion decreased in the follow-up exams.
have an impact on individual patient care and on clinical trials of new therapies (alguma referencia).

**Pseudoprogression**

Pseudoprogression is a subacute treatment-related reaction, usually associated with asymptomatic patients [2]. Shortly after completion of RT, predominantly within the first 3 months, patients with high-grade brain tumors can present with an increase in contrast-enhancing lesion size, mimicking tumor progression, followed by subsequent improvement or stabilization without any further treatment [3, 4].

**Pseudoresponse**

Pseudoresponse is a rapid decrease in contrast enhancement observed in recent high-grade glioma treatment trials after administration of antiangiogenic agents such as bevacizumab and cediranib, a VEGF receptor tyrosine kinase inhibitor [5]. These agents produce a high response rate and 6-month progression-free survival, but with rather modest effects on overall survival.

**Mcdonald Criteria**

The Macdonald Criteria [6] are currently the most widely used guideline for assessing response to therapy in patients with high-grade gliomas. According to the Macdonald Criteria, tumor progression is considered to have occurred when an increase of 25% in the size of the contrast-enhancing lesion is observed. There are important limitations to these criteria, since they address only the contrast-enhancing component of the tumor, which is nonspecific and may not always be considered a true surrogate of tumor response.

**Pathophysiology**

Pseudoprogression is found to correspond to gliosis and reactive radiation-induced changes without evidence of viable tumor [6]. It may represent an exaggerated response to effective therapy, involving early changes to the vascular endothelium and the blood-brain barrier (BBB), causing new or increased contrast enhancement on MR imaging examinations. Most importantly, some studies have found an association between the incidence of pseudoprogression and increased survival; perhaps pseudoprogression represents an active ‘inflammatory’ response against the tumor [4]. The early decrease in contrast enhancement seen in pseudoresponse suggests a change in vascular permeability, with a ‘normalization’ of the BBB, rather than a true tumor reduction, as being the underlying cause of the improvement [7]. Normalization of the BBB and subsequent reduction in the vasogenic edema can result in an improvement of symptoms [8, 9].

**O6-Methylguanine DNA MGMT promoter**

The methylation status of the methyltransferase (MGMT) promoter has been shown to be a potent prognostic factor in patients with GBM; cells that are deficient in MGMT have shown an increased sensitivity to TMZ. Furthermore, MGMT promoter status may predict pseudoprogression in ~90% of patients with methylated glioblastoma [10], due to higher sensitivity to treatment [4] and an approximately 60% probability of early true tumor progression was observed in un-methylated MGMT promoter tumors [11]. Thus, we can speculate that methylated MGMT may be a good indicator of therapeutic response and better prognosis, as an increased overall survival rate has been observed in these patients.
Advanced MR imaging techniques

No single imaging technique has been validated to recognize and adequately establish a diagnosis of pseudoprogression [12] and the diagnosis should depend on follow-up scans until an improved method is established. Dynamic susceptibility-weighted contrast-enhanced perfusion MR imaging (DSC) is a surrogate marker for angiogenesis and has been used to assess brain tumor treatment response with high sensitivity for distinguishing residual/recurrent neoplasm from radiation brain injury [13–15]. Permeability DSC is also a potential new tool for differential diagnosis between pseudoprogression and true tumor progression. Although no prospective study has examined this hypothesis, preliminary results with this new technique seem very promising, and a number of clinical trials are underway to better delineate the performance of all of the above techniques.

Other techniques, such as MR spectroscopy, diffusion-weighted imaging (DWI) and diffusion tensor imaging (DTI) have been assessed to differentiate tumor progression and/or residual tumor from necrosis [11, 16]. In short, none of them provides sufficient information for differential diagnosis between pseudoprogression and true tumor progression.

Management

Pseudoprogression may influence the clinical recommendation to continue with adjuvant chemotherapy rather than the change to a second-line therapy for recurrence [17]. Currently, the only method of distinguishing pseudoprogression and early progression of disease is to perform follow-up examinations of the patients, since conventional MR imaging is unable to differentiate them and alternative techniques have not yet been validated in prospective trials [4, 11]. Analysis with follow-up conventional MR imaging examinations allows such a distinction because the changes related to pseudoprogression decrease in size. If a post chemoradiotherapy follow-up MR imaging examination demonstrates complete or partial response or stable disease (i.e. smaller or stable tumor enhancement), maintenance of chemotherapy is typically continued. When enlargement occurs, then the treating...
Conclusion

Pseudoprogression and pseudoresponse are abnormalities that have been described following high-grade tumor treatment, and remarkably both appear to be associated with future favorable patient outcome. Both phenomena appear to be best diagnosed through follow-up scans because no established method of imaging is yet capable of yielding a definitive diagnosis of true tumor versus enhancement changes due to other reasons. DSC and other methods appear promising but require further testing in the multicenter setting.

After failure of the concomitant radio and chemotherapy, second line treatment with antiangiogenic drug was indicated. Just after the initiation of anti-VEGF therapy, a reduction in the contrast enhancing area was seen. Besides a continuing reduction in the enhancing portion of the lesion, an expansion is observed in the FLAIR sequence. However, diffusion images demonstrate an area of restricted diffusion, which may correspond to areas of tumor dissemination, and which may lead to new areas of contrast enhancement.

physician does face a dilemma. If pseudoprogression is suspected, perhaps based on MGMT status and/or very early changes in imaging features in the first months post treatment, ongoing chemotherapy with TMZ might be continued, with close monitoring. In clinically symptomatic patients, more options must be considered, including cessation of therapy, addition of anti-VEGF treatment, or even surgery, since identical symptoms can be observed in patients with true tumor progression and patients with pseudoprogression [8].

Contact
L. Celso Hygino Cruz Jr.
Clinica de Diagnostico por Imagem (CDPI,DASA)
Centro Medico Barrashopping
Av das Americas 466, Sl 325
Rio de Janeiro
Brazil
celsohygino@hotmail.com
References
Imaging and Spectroscopy at 9.4 Tesla: First Results on Patients and Volunteers

R. Pohmann¹; G. Shajan¹; J. Hoffmann¹; J. Budde¹; G. Hagberg¹²; O. Bieri³; S. Bisdas⁴; U. Ernemann⁴; M. Weigel⁵; Ph. Ehses¹²; J. Hennig⁶; G. Chadzynski¹², K. Scheffler¹²

¹MRC Department, MPI for Biological Cybernetics, Tübingen, Germany
²Dept. Biomedical Magnetic Resonance, University of Tübingen, Tübingen, Germany
³Division of Radiological Physics, Department of Radiology, University of Basel Hospital, Basel, Switzerland
⁴Dept. of Neuroradiology, University of Tübingen, Tübingen, Germany
⁵Department of Radiology Medical Physics, University Medical Center Freiburg, Freiburg, Germany

Introduction

9.4 Tesla (T) is currently the highest field strength to have been successfully used for imaging and spectroscopy in humans, and the Max Planck Institute for Biological Cybernetics in Tübingen is probably the first site to have enrolled patients for an ethically-approved study of brain tumors in November 2012. Since 2008 when the first human MR imaging (MRI) / MR spectroscopy (MRS) measurements at 9.4T began in Tübingen, 261 healthy subjects have volunteered to participate.

The big step from 3T to 7T has clearly shown the advantages and problems at very high fields. Most prominent are challenges related to the transmitted radio frequency (RF) field in terms of specific absorption rate (SAR) and transmit homogeneity, as well as increased susceptibility effects. The next step from 7T to 9.4T is somewhat smaller and, accordingly, further increased RF inhomogeneity and susceptibility variations have been observed, although these are qualitatively similar to those effects reported at 7T. So far, in our experience...

1A (1A) Photograph of the transmit array hardware. Integrated TR switches with preamplifiers enable the coil to be used in the transceive mode as well. 1B (1B) Photograph of the 31-channel receive-only array. 1C (1C) Final RF configuration with the receive array assembled inside the transmit array.
the most striking differences between 7T and 9.4T are the even shorter RF wave length and the significantly increased susceptibility effects. The short wave length and thus locally very confined excitation patterns actually increase the flexibility and effectiveness of parallel transmit techniques and lead us to the design of transmit arrays with 16 independent loops. The sensitivity to local susceptibility changes is also highly increased, as demonstrated in T2*-weighted BOLD experiments, and effects related to oriental anisotropy become clearly visible.

In this report we show examples of different imaging techniques and spectroscopy that has been measured during the last two years at 9.4T. The goal was to test the feasibility of basic imaging sequences, such as gradient echoes, and to apply advanced techniques like TrueFISP and hyperechoes with reduced SAR. Quantitative comparisons of signal strength, relaxation times and contrast can be found in the referenced papers.

**Dedicated RF coils for 9.4T**

In addition to the static magnetic field strength (B₀), RF coil performance is a
dominant factor in determining the sensitivity of an MR experiment. The principal RF engineering challenge at high \( B_0 \) field is to achieve homogeneous excitation across the volume of interest. As the wavelength in tissue approaches sample dimensions, constructive and destructive interference of the electromagnetic field causes field concentration, but also signal voids, resulting in an inhomogeneous MR image \([1]\). At 9.4T, even in human brain MRI, severe field dropouts appear in the lower temporal lobe and cerebellum if the coil is driven in the conventional circularly polarized (CP) mode, making it a challenging task to achieve whole-brain excitation. While the traveling wave approach is able to extend the imaged volume into the lower regions of the brain with often acceptable homogeneity \([2]\), static and dynamic RF shimming approaches are the most promising techniques to mitigate the \( B_1^+ \) field inhomogeneities. They require an array of transmit elements with the possibility to individually control the amplitude and phase of the current to each of these coil elements. An additional degree of freedom can be obtained by arranging the transmit elements in multiple rows to extend the longitudinal coverage and more importantly, to correct the inhomogeneities in all three directions \([3]\). While transceiver coils, which combine multi-channel transmit and receive in the same array elements, are a popular design for ultra-high field applications \([4]\), both signal-to-noise ratio (SNR) and parallel imaging performance can be improved by using a separate array of receive coil detectors that closely follows the contours of the anatomy \([5]\). Parallel imaging benefits particularly from the shorter wavelength at ultra-high field strength because of reduced inductive coupling to the farther coil elements, resulting in distinct coil sensitivities and hence effective sensitivity encoding \([6]\). Our approach for human brain imaging at 9.4T combines separate transmit and receive arrays to maximize the receive sensitivity together with the ability to modulate the transmit field in three dimensions.

The transmit coil consists of 16 elements arranged in two rows of eight elements each \([3]\). The lower row elements are rotated by 22.5° with respect to the upper row and all adjacent coil elements.
are inductively decoupled. For reception, 31 elements are arranged symmetrically in 4 rows on a close-fitting helmet for maximum sensitivity. A combination of inductive decoupling and geometric overlap is used to minimize the inductive coupling between the coil elements [7]. The transmit and receive arrays and the final setup are shown in Figure 1. Unlike for the closely-coupled transceiver arrays, subject-specific adjustments of the transmit array are not required. Furthermore, the RF circuitry and low noise preamplifiers required for the receive elements are closely packed in the coil housing, thus providing a simple and fast setup, comparable to clinical routine examinations. Due to the dual row design of the transmit array, the entire brain, including the cerebellum, can be imaged. This is demonstrated in the sagittal image in Figure 2, for which slice selective static phase shimming was applied to optimize the transmit phase.

High-resolution GRE, TrueFISP and susceptibility-weighted imaging

Due to the high sensitivity towards variations in the magnetic susceptibility at 9.4T, improved and even novel contrast mechanisms can be achieved. The pulse sequences required to exploit these advantages are based on gradient-echo techniques and are relatively insensitive towards SAR limitations and inhomogeneities of the transmit field. T2*-weighted images (Fig. 3A) already show excellent image contrast and can replace T1 or T2-weighted imaging techniques that suffer from serious SAR limitations and B1-dependent contrast variations in many applications. In addition, these techniques allow for fast acquisition with parameters that are optimized for maximum SNR. Thus, using flip angles close to the Ernst-angle and carefully adjusting bandwidth and echo time makes it possible to reach high spatial resolutions in acceptable scan times. The image in Fig. 3A was acquired with a voxel volume of 40 nl (voxel size 0.2 x 0.2 x 1 mm³, echo time 20 ms, repetition time 28 ms, 21 slices) within around 15 min, using the above described combination of a 16-channel transmit and a 31-channel receive array. Even higher contrast-to-noise ratio can be reached by using the image phase as completely B1-independent contrast parameter. Phase images (Fig. 3B) show good contrast between gray and white matter, but are also able to depict intracortical structures or oriented fibers. In addition, phase imaging at high fields is used to detect changes in iron or myelin content [8, 9]. At 9.4T, the high sensitivity of phase imaging makes voxel sizes below 20 nl feasible within reasonable scan times of around 20 minutes. Finally, magnitude and phase information are combined in susceptibility-weighted imaging (SWI, Fig. 3C) to specifically emphasize susceptibility variations as in venous blood. In a comparison to images acquired at 3T, an SNR gain of almost a factor of nine was found [10]. In addition, the high sensitivity towards susceptibility variations made it possible to distinguish venous structures at a considerably smaller size. These data show that anatomical imaging with high resolutions, great contrast and high image quality is possible at 9.4T without the limitations due to SAR or B1-inhomogeneity. Modulations of the magnetic susceptibility, especially, can be detected with high accuracy with simple gradient echo sequences at 9.4T. Additionally, the strong susceptibility effect can help to improve the spatial specificity of spin-echo EPI-based functional MRI studies [11]. Successful balanced SSFP imaging is commonly hampered by its prominent
sensitivity to local frequency offsets (off-resonances) that may lead to pronounced signal voids, typically appearing as dark bands in the image. Since the major cause of such local off-resonances are susceptibility-related frequency shifts that are expected to scale linearly with the main magnetic field, successful banding artifact reduction becomes a key issue at ultra-high field MRI with balanced SSFP.

Here, we demonstrate in vivo balanced SSFP imaging of human brain at 9.4T. Signal voids with balanced SSFP become apparent for frequency offsets close to about 1/(2TR) or -1/(2TR) or multiple thereof, where TR is the repetition time of the balanced SSFP imaging protocol. As a result, the appearance of banding artifacts with balanced SSFP is directly related to its repetition time and can thus be mitigated by a reduction by ultra-fast imaging, typically associated with TR close to about 1 ms. For a TR ~1.6 ms (see Fig. 4), banding artifacts are expected for local frequency offsets as far as about ±300 Hz, and artifact-free balanced SSFP imaging becomes available and can be demonstrated even at ultra-high fields. SAR constraints that become more severe at ultra-high fields and the exceptional short TR used with ultra-fast imaging limit the maximum flip angle for balanced SSFP imaging for in vivo applications to about 10°. For the typical low T2/T1 << 1 for tissues and with respect to an estimated T2/T1 ~1/30 for brain tissue at 9.4T, however, optimal balanced SSFP imaging becomes shifted towards the low flip angle regime (~20°), approaching the limits set by SAR. Moreover, for balanced SSFP only a marginal contrast is expected between normal appearing gray and white matter tissue due to the highly similar T2/T1, being in contrast to the experimental results, as shown in Fig. 4. Similar to what was observed at low fields, the appearance of a prominent contrast between gray and white matter is likely due to magnetization transfer effects.

**Time-of-flight imaging in patients and volunteers**

Time-of-flight (ToF) angiography aims to visualize the cerebral arteries and relies on the in-flow effect: continuous excitation suppresses the local signal while in-flowing blood remains unsuppressed and strongly contributes to the final image contrast. The most popular approach is selective 3D excitation of a slab with limited extent. The image contrast is determined by the slab thickness, the velocity of the blood flow across the slab and by the sequence repetition time, TR. The goal is to avoid saturation of the flowing blood, allowing it to cross the entire slab within just one or a few TR times. ToF angiography at ultra-high magnetic field strengths benefits from long tissue T1 times that lead to improved background suppression. However, SAR restrictions impose severe limits on the reduction of tissue signal that may be achieved, and the suppression of signals in the cerebral veins, obtained by additional pulses, also becomes challenging. At 7T, the use of VERSE (variable-rate selective excitation, [12]) pulses has proven beneficial to attain both goals: a high signal contrast in arteries only within SAR limits [13, 14]. VERSE consists of a shaped RF pulse applied together with a time-varying gradient. In order to reduce SAR without excessive time penalty, pulse segments with high RF power are prolonged in time, while segments with low power requirements are shortened. The excitation bandwidth of VERSE is therefore effectively varied from segment to segment, making it sensitive to off-resonance effects, such as chemical shift differences and B₀ inhomogeneities. Since such effects tend to increase with the magnetic field strength, we decided to use the TONE RF pulse with a 100% flat top shape for excitation rather than VERSE. The sole modification of the standard clinical...
syngo MR B15 sequence was to allow a variable duration of the excitation pulse. Slab-wise imaging (2.4 cm axial FOV) with a 3D gradient echo sequence, a TR of 20 ms, a GRAPPA factor of 4 and 32 reference lines, and an in-plane voxel size of 0.5 × 0.5 mm with a slice thickness of 0.4 mm was used. The echo time (3.8; 4.55; 10 ms), flip-angle (FA 15–36°) and the duration of the excitation pulse (1024–4096 μs) were varied. The acquisition time for each slab was between 1 ½–3 ½ min.

Post-processing consisted of removing the scalp signal originating from the subcutaneous fat using BET (Brain Extraction Tool, FSL$^3$), and intensity correction, obtained after thresholding out the angiographic information and fitting the background signal with a 2nd order polynomial. Maximum Intensity Projections were performed across 40 mm. We found that a reasonable flip angle at 9.4T is 20–30° to optimize the signal for blood that remains within the slab for about 5 excitations.

The improved contrast-to-background ratio that we expected was not achieved with the standard TONE pulse due to SAR restrictions. Even in the absence of pulses for the suppression of venous blood or magnetization transfer, we came close to the SAR limits (95–99%) with low flip angles of 14–16°. Under these conditions, the venous signal was prominent and the image contrast was poor (Fig. 5A). By increasing the duration of the RF pulses, the flip angle could be increased to 24°–32°, yielding an improved image contrast (Fig. 5B). Nevertheless there was no room for additional venous suppression pulses, not even by the reduced flip angle approach that has been successfully introduced at 7T [15]. At 9.4T, not only the T1 but also the T2 relaxation times are changed. The T2 in arterial blood is similar to tissue (about 40 ms) while in venous blood it is substantially shorter (5–9 ms dependent on the fractional oxyhemoglobin content [16]). In line with these observations we found that increasing the TE from 4.5 to 10 ms was sufficient to suppress the venous blood in the sagittal sinus without substantially compromising the visibility of the arteries (Fig. 5C).

**Hyperecho imaging**

Turbo spin echo (TSE, RARE, FSE) sequences [17] find widespread application in clinical routine MRI, the main reason being that they combine the diagnostically relevant T2 contrast with the robust signal behavior of spin echo refocusing. T2 contrast is highly sensitive to a broad variety of pathologies such as inflammation and is an indispensable tool for examination of various pathologies of the CNS. Spin echo refocusing facilitates a relative insensitivity to susceptibility and field inhomogeneity, i.e. B$_0$ effects. Additionally, TSE sequences produce a high amount of stimulated echo contributions that are T1-weighted. Since T1 >> T2 for most biological tissues, particularly at ultra-high fields, these stimulated echo contributions partially counterbalance the signal loss from refocusing flip angles deviating from 180° due to B$_1$ variations. All in all, these reasons seem to make TSE sequences the ideal technique for clinical imaging at (ultra-)high fields.

However, a major drawback is the high RF power deposition caused by the multiple refocusing RF pulses, which can lead to severe tissue heating. Thus, the RF power deposition, usually quantified in terms of SAR has to be strictly limited. Since SAR increases quadratically with field strength, the same image acquired at 9.4T requires almost ten times the SAR as at 3T. Therefore, the fast multiple refocusing of magnetization with high turbo factors in modern TSE sequences severely hampers their use at ultra-high fields.

Common ways of mitigating SAR problems include reducing the number of slices, shortening the echo train via a smaller turbo factor, increasing the repetition time TR, changing the shape of the RF pulses (e.g. using a Gaussian shaped pulse) and increasing the duration of the RF pulses. No matter which (combination of) option(s) is chosen, some compromise has to be made in

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Three out of five slices from the hyperTSE experiment performed at 9.4T in Tübingen.
Whereas the Hyperecho-scheme makes use of symmetry relations between the flip angles to reinstate the full signal at some given point of time (echo) within the TSE train [18], TRAPS allows an almost free variation of refocusing flip angles along the TSE train after a preparation of magnetization close to the so-called static pseudo steady state (SPSS) [19, 22, 23]. In the SPSS, the echo intensity always stays close to the optimal value that can be reached for a given flip angle. Any loss of coherence would lead to signal fluctuations and inevitable signal loss, resulting in image artifacts. Since the variation of signal in k-space directly influences the point spread function (PSF), flip angle variations can also be used to improve the PSF. Thus, TRAPS, which offers the unique possibility to vary flip angles in a flexible way in hyperTSE sequences, allows both considerable SAR savings as well as the optimization of image properties for TSE [19, 20, 21, 24]. Methods that employ variable instead of constant refocusing flip angles along the TSE train have been introduced in order to allow SAR reductions while maintaining a T2-like contrast and high SNR. These novel TSE sequences are usually referred to as hyperTSE-, Hyperecho-TSE-, and smooth transitions between pseudo steady states (TRAPS)-sequences [18, 19, 20, 21]. Generally, high signal for the echoes encoded for the center of k-space is maintained, thus, warranting a high SNR for the reconstructed image. The signal in the peripheral parts of k-space may be reduced slightly due to lower flip angles; however, strong stimulated echo contributions still maintain sufficient signal levels at ultra-high fields due to T1 >> T2, as mentioned above.

Whereas the Hyperecho-scheme makes use of symmetry relations between the flip angles to reinstate the full signal at some given point of time (echo) within the TSE train [18], TRAPS allows an almost free variation of refocusing flip angles along the TSE train after a preparation of magnetization close to the so-called static pseudo steady state (SPSS) [19, 22, 23]. In the SPSS, the echo intensity always stays close to the optimal value that can be reached for a given flip angle. Any loss of coherence would lead to signal fluctuations and inevitable signal loss, resulting in image artifacts. Since the variation of signal in k-space directly influences the point spread function (PSF), flip angle variations can also be used to improve the PSF. Thus, TRAPS, which offers the unique possibility to vary flip angles in a flexible way in hyperTSE sequences, allows both considerable SAR savings as well as the optimization of image properties for TSE [19, 20, 21, 24]. A great emphasis in TRAPS lies on the flip angle train, which has to be properly adapted to any changes of the protocol settings that influence the timing or the view ordering / reordering of the TSE sequence. ‘Unwise variations’ can lead to suboptimal image and SAR properties. Basically, there are two recipes to calculate suitable flip angles for TRAPS:

1. Setting the flip angles directly via some support points defined by functions, etc. This AUTO-TRAPS approach has the advantage that the flip angle courses can be adapted in a flexible way to consider protocol changes made by the user, and the AUTO-TRAPS recipe allows a quite direct control of SAR [21].

2. More sophisticated refocusing flip angle schemes can be obtained by calculating the flip angles from a predefined signal course or echo intensity course along the echo train [24]. This method has the advantage that dedicated image properties can be set via the predefined signal course [24]. However, there is no unique solution for the flip angles and it is more complex to adapt the settings to any protocol changes made by the user.

For our proof of concept study at 9.4T, a flexible AUTO-TRAPS approach was used [21]. The TRAPS based flip angle method was implemented in a self-made TSE sequence within the IDEA platform syngo MR B15. A T2-weighted TSE protocol with the following parameters was...
set up: FOV 220 × 175 mm², 5 slices with 1 mm slice distance, voxel size 0.58 × 0.58 × 1.0 mm³, turbo factor 17, TE 48 ms, TR 8.94 s. Due to additional T1-weighted stimulated echo contributions, hyperTSE sequences (as all low flip angle TSE sequences) show a reduced T2 contrast compared to a TSE 180° sequence at a given echo time. In order to compensate for this effect, TE was increased from the originally desired value [20]. Here, please note that T2 relaxation times are considerably shorter at 9.4T than at 3T or even 1.5T. Thus, a comparatively low TE already gives a substantial T2 contrast.

The used hyperTSE protocol demonstrated a relative SAR of 33.7%, i.e. a SAR saving of 66.3% compared to a conventional TSE sequence. This allowed the acquisition of 5 slices instead of only 1. Fig. 6 shows the first hyperTSE images from a patient at 9.4T. The used AUTO-TRAPS based hyperTSE demonstrates very good imaging behavior and suggests that hyperTSE sequences are highly suitable for acquiring ultrahigh resolution images at 9.4T that offer promising diagnostic value.

Proton spectroscopy in patients and volunteers

Chemical shift imaging (CSI) data were collected with a Stimulated Acquisition Mode sequence (STEAM) with the following parameters: TE 20 ms; TR 2000 ms; TM 11 ms; FOV 160 × 160 mm²; volume of interest (VOI) 60 × 60 × 10 mm³ and 40 × 40 × 10 mm³ (volunteer and patient respectively); voxel size 10 mm³ isotropic; spectral bandwidth 4000 Hz. In order to keep the reference voltage below the hardware limits, the 90° h-sinc RF pulses (sinc pulses with 4 side lobes) within the standard STEAM sequence had to be replaced by 90° hermite pulses. Additionally, their RF bandwidth has been increased to 3100 Hz. This allowed minimizing the influence of chemical shift displacement. In order to determine the reference voltage necessary for the STEAM sequence, CSI acquisition was preceded by an Actual Flip angle Mapping (AFI) scan [25, 26]. Figure 7 shows spectra acquired from a healthy volunteer (31 years old). In this case, an 8-channel transmit coil combined with a 24-channel receive array was used for data acquisition. Here, the spectra were acquired from a superior region of the human brain. The VOI (yellow square, Fig. 7A) was placed in the axiplane, parallel to the line between the anterior and posterior commissures (ac-pc line). A typical spectrum from healthy brain tissue (localization within the VOI marked with blue square) is presented in Fig. 7B. The following brain metabolites have been identified: myoinositol (Ins, marked with 1), methylene (CH₂) and methyl groups (CH₃) of creatine and phosphocreatine (Cr and Pcr, #2), overlapped peaks of glutamine and glutamate (Gln and Glu, #3), taurine (Tau, #4), choline containing compounds (tCho, #5), aspartate (Asp, #6), N-acetylaspartate (NAA, #7), Gln (#8), Glu (#9), γ-Aminobutryic acid (GABA, #10), N-acetylaspartylglutamate (NAAG, #11) and macromolecules (#12).

Improved spectral resolution and SNR make it possible to differentiate between metabolites which are typically overlapped at lower field strengths, particularly the resonances between ~4 and ~3.2 ppm, and ~2.9 and 2.1 ppm (Ins, Glu, Gln, GABA, NAA and NAAG). Those findings are in agreement with results described in previous studies [27, 28]. Analysis of the spectra acquired from the entire VOI (Fig. 7C) shows that the spectra from the central 4 × 4 voxel region have excellent quality. Specialized sequences [29, 30] and parallel transmit techniques [31] will further improve the spectra by reducing the chemical shift displacement, which is still significant for some of the peaks (tCho, Cr and NAA), as well the dependence of the signal amplitude on the spatially inhomogeneous transmit field. Spectra acquired from the brain of a 43-year-old patient with clinically confirmed oligodendroglioma are shown in Fig. 8A. MR scanning showed the presence of an extended tumorous mass in the right hemisphere, pressing on the cerebral midline that was deviated towards the left hemisphere. In the antero-posterior direction the tumor extended from the frontal to the parietal lobe, and in the crano-caudal direction from the hand area in the primary motor cortex, across the lateral fissure into the temporal cortex. The tumor texture was heterogenous and showed calcifications, confirmed by CT scans, indicating an oligodendroglioma.

Here all the CSI data were collected with a 16-channel transmit coil combined with a 31-channel receiving helmet. Spectra were collected at two different locations: within the lesion (Fig. 8A), and contralaterally, in healthy tissue (Fig. 8B). Analysis of the acquired spectra revealed a strong decrease in the signal of tCho and an increase in signals of tCho, Glu, Gln, and Ins within the lesion (Fig. 8A) compared to the contralateral side (Fig. 8B). Observations made here are in agreement with the previous clinical studies performed at lower magnetic fields (1.5 and 3T) [28, 29]. Furthermore, our results suggest an increase in signals of Tau and scyllo-inositol (Scylo) within the tumor tissue (Figs. 8A and 8B, marked with yellow arrow). According to the latest research those play an important role in distinction between different tumor types or grades [32, 33] and may serve as prognostic markers [34]. However, the accurate detection of Scylo and Tau with 1H MRS at lower magnetic field strengths could be difficult as they are strongly overlapped with the neighboring resonances of tCho and Ins.

Summary

Although inhomogeneity in B₁ and B₀ is further increased at 9.4T compared to 7T resulting variations in signal strength across the image is acceptable and comparable to 7T. We currently investigate the possible gain in SNR going from 3T to 7T and to 9.4T, and preliminary results indicate the expected linear...
increase. However, a solid comparison is quite difficult due to very different coil geometries and technologies used at these different field strengths. Macroscopic homogeneous $B_1$ and $B_0$ fields are the prerequisite for the study of microscopic susceptibility changes within tissue, which are significantly enhanced at ultra-high fields. Thus, one of the most important steps in the near future in our lab will be the implementation and application of parallel transmit concepts and corresponding RF coil technology, as well as systems to improve and control $B_0$ homogeneity such as dynamic shimming with high-order shim inserts.

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The 7T and 9.4T systems are investigational devices. The products mentioned herein are still under development and not commercially available yet. Their future availability cannot be ensured.

The Brain Extraction Tool, FSL is not a Siemens Healthcare Product. Siemens bears no responsibility for this product including, but not limited to, its regulatory status.
References:


Improved Echo Planar Diffusion-Weighted imaging of the Head and Neck using syngo ZOOMit

Philipp Riffel, Henrik J. Michaely

Institute of Clinical Radiology and Nuclear Medicine, University Medical Center Mannheim, University of Heidelberg, Germany

DWI as a cancer biomarker

Echo-planar imaging (EPI) is a single-shot MR imaging (MRI) technique based on the rapid acquisition of a train of sign-alternating gradient echoes [1]. EPI sequences offer high signal-to-noise ratio (SNR) and imaging speed (volume coverage) which render them attractive for functional neuroimaging and diffusion-weighted magnetic resonance imaging (DWI). Imaging biomarkers are important tools for the detection and characterization of cancers as well as for monitoring the response to therapy. DWI is rapidly gaining popularity for the assessment of oncologic and non-oncologic pathologies. DWI depends on the microscopic motion of water. This motion, called Brownian motion, is due to thermal agitation and is highly influenced by the cellular environment of water. Thus, findings on DWI could be an early harbinger of biologic abnormality [2]. Once a technique primarily used in neuroradiology, diffusion-weighted MRI is already being incorporated into general oncologic imaging practice because of its many clinical advantages. Recently, Padhani and Koh have alluded to the promising future of DWI as a cancer biomarker in tumor staging with an improved tissue characterization (differentiating benign from malignant lesions), in monitoring response to chemotherapeutic agents, and in differentiating post-therapeutic changes from residual active tumor [2].
EPI-DWI in the head and neck

DWI is potentially useful in the evaluation of head and neck lesions [3, 4] and ADC measurements have suggested that this imaging method may be useful for their characterization [4]. Although results of initial studies were both challenging and promising, one drawback was the technical difficulty in assessing diffusion by echo-planar DW imaging. Applications of EPI may be affected by inherent problems, such as ghosting artifacts, as well as by geometric distortions and signal losses caused by chemical shift and susceptibility differences. These effects degrade the achievable image quality and limit the spatial resolution. This is especially valid for the head and neck area. Due to the presence of dental work, as well as adjacent air and bone, it may be difficult to obtain precise ADC measurements in lesions located in these regions. Therefore in the majority of studies only lesions larger than 1 cm were evaluated. Currently the determination of accurate ADC values in smaller tumors and lymph nodes seems to be unreliable [3-5]. Therefore, DWI techniques that are less sensitive to susceptibility artifacts are mandatory for the evaluation of head and neck lesions.

Solution approaches

Physically, the above mentioned artifacts are caused by phase distortions, which increase with longer echo time and mimic the encoding of spatial information during image reconstruction. Therefore, fundamental to overcoming these limitations is the use of shorter echo trains [6]. There are several possibilities for reducing these artifacts and conventional DWI has seen many improvements. Latest developments include, for example, read-out segmented EPI [7]. In combination with navigator-based phase corrections and re-acquisition, read-out segmented EPI can effectively reduce susceptibility artifacts [8] and is now commercially available as syngo RESOLVE. Nevertheless, this approach may increase acquisition time and is best applied in the imaging of non-moving organs. A complementary way of decreasing the echo train is to reduce the field-of-view (FOV) along the phase-encoding dimension of the image while simultaneously avoiding image aliasing. First attempts used spin-echo and stimulated-echo EPI sequences with orthogonal RF pulses to excite an inner-volume FOV and then reduce the number of k-space lines required for image reconstruction [9]. A major drawback of these techniques is the restricted volume coverage due to neighboring sections becoming presaturated by one of the RF excitations [6]. With the clinical availability of parallel RF transmit coils and the potential to utilize its spatial information in an array during RF transmission, it is now possible to move beyond the uniform slice-select excitation and to generate spatially-tailored RF pulses. Two-dimensional spatially-selective RF excitation pulses for
single-shot echo-planar imaging combined with reduced FOV – i.e. zooming – in the phase-encoding direction leads to a decreased number of acquisition k-space lines and significantly shortens the length of the EPI echo train [6, 10]. The numerically calculated two-dimensional RF pulse provides, in addition, superior flip angle homogeneity, since it is optimized using the B1 field information of the two independent RF channels. To summarize, zoomed imaging provides largely improved homogeneity with decreased distortion artifacts, opening new clinical applications. We have investigated the applicability of zoomed diffusion imaging of the neck, to examine its potential for reduced susceptibility artifacts in regions close to major air cavities and the associated clinical relevance.
Initial results of zoomed DWI in the head and neck

All examinations were performed on a 3T whole-body MR system (MAGNETOM Skyra, Siemens Healthcare, Erlangen, Germany) equipped with a two-channel fully dynamic parallel transmit array, termed TimTX TrueShape. The excitation of the standard diffusion-weighted EPI sequence was extended by the two-dimensional spatially-selective RF pulse using an echo-planar transmit trajectory. We found that syngo ZOOMit bears considerable potential for overcoming some of the inherent limitations of conventional EPI techniques. In particular, for a reduced FOV, zoomed DWI allows for improved image quality in terms of markedly reduced susceptibility artifacts. Zoomed images allow for an excellent correlation between the zoomed DW images and the morphological T2 TSE images (Fig. 1). This is especially obvious in regions prone to artifacts, e.g. in the tongue or the hard palate (Fig. 2).

As head and neck squamous cell carcinomas (HNSCC) are often located in those anatomical distributions, zoomed imaging could potentially improve diagnostic accuracy in the assessment of cancers in early stage in those areas. Additionally, due to the improved image quality with less distortions, zoomed images provide perfect delineation of cervical lymph nodes and an exact co-registration with morphological images. Figure 3 shows images of a patient with malignant lymph nodes. While the malignant lymph nodes could also be seen on the conventional diffusion-weighted images, the differentiation between lymph node and vessels could only be achieved with zoomed DWI. Due to the reduced distortion artifacts, the external carotid artery delivers a sharply delineated flow void, which allowed the radiologist to absolve the vessel of tumor infiltration.

The achieved improvements in image quality originate from both the shorter echo train (at a given TE) and the possible reduction of the TE. On the other hand, the improved image quality is at the expense of a smaller FOV and a lower SNR due to the decreased number of acquired echoes. To cover a larger FOV several images of spatially shifted fields-of-view may be composed (Fig. 4).

Conclusions
These initial data are encouraging as they demonstrate that zoomed imaging (syngo ZOOMit) may be a very robust method for DW imaging of challenging areas such as the head and neck, enabling more reliable imaging, particularly in areas close to major air cavities. With improved correlation with anatomy, zoomed DWI may assist in the detection and assessment of early cancers in regions where the application of conventional DWI may have been limited.

Such developments are crucial, especially given the need to counteract the much more pronounced susceptibility problems in high-field MRI systems at 3T and beyond.

References
Sodium (\(^{23}\)Na)-Imaging as Therapy Monitoring in Oncology – Future Prospects

Stefan Haneder, M.D.¹; Stefan O. Schoenberg, M.D.¹; Simon Konstandin, Ph.D.²; Lothar R. Schad, Ph.D.²

¹Institute of Clinical Radiology and Nuclear Medicine, University Medical Center Mannheim, Heidelberg University, Mannheim, Germany
²Computer-Assisted Clinical Medicine, Heidelberg University, Mannheim, Germany

Introduction

“Form follows function” – Louis Sullivan, 1896

Whilst this concept was originally applied to modern architecture, it could well become a highly appropriate maxim for future imaging and therapy concepts. Magnetic resonance imaging (MRI) has continually developed into a powerful, widely used diagnostic tool and offers the opportunity to expand traditional imaging concepts based on morphological information. In the future, the pure morphology will remain a central component of multimodal imaging, but will be flanked increasingly by functional approaches reaching far beyond the current imaging standards. In oncological therapy follow-up the drawback of relying on pure morphology is widely known, resulting, for example, from delayed morphological reflection of tumor regression. Consequently, the RECIST Working Group addressed this point in the context of the new RECIST1.1 criteria [1]: “A key question considered by the RECIST Working Group in developing RECIST 1.1 was whether it was appropriate to move from anatomic unidimensional assessment of tumour burden to either volumetric anatomical assessment or to functional assessment with PET or MRI. It was concluded that, at present, there is not sufficient standardisation or evidence to abandon anatomical assessment of tumour burden. The only exception to this is in the use of FDG-PET imaging as an adjunct to determination of progression. [...]”

This statement contains several basic implications for future MR strategies of therapy control. First, the internal radiology benchmark of functional MR sequences seems to be nuclear medicine approaches, already fixed in guidelines as PERCIST 1.0 [2]. Second, further multi-center, international studies are required to obtain reliable data for (functional) MR approaches. Third – not explicitly, but indirectly – the radiology community should not abandon the assessment of new functional approaches and should try to implement them in clinical settings. The arsenal of current functional MR imaging approaches includes diffusion-weighted imaging (DWI), diffusion tensor imaging (DTI), arterial spin labeling (ASL), blood-oxygenation level dependent imaging (BOLD) and sodium (\(^{23}\)Na)-imaging. DWI [3] should be emphasized as a kind of paradigm shifting technique. Since the 1990s DWI has been performed for intracranial diseases and has contrib-

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**Figure 1**

1A Sequence design for 3D radial acquisition with global excitation. Absolute gradient values (\(\sqrt{G_1^2 + G_2^2} \)) are shown for standard radial acquisition with constant gradient readout (\(\|G\|_{\text{readout}}\)) and sampling density-weighted apodization (\(\|G\|_{\text{samp}}\)) for intrinsic filtering. Due to hardware restrictions the gradient adaption begins at \(t = t_0\) with gradient amplitude \(G_0\). The echo time TE is defined as the time interval between the middle of the RF pulse and the beginning of data acquisition with readout duration \(T_{RO}\). The k-space sample points are exemplarily shown in the \(k_{xy}\)-plane for standard radial (1B) and SDWA (1C) acquisition. Figure adapted from [18].
uted to the detection of early stroke. But in recent years the number of studies has increased substantially, evaluating this method in detection and characterization of lesions, especially in the field of oncology and the possibility of (early) tumor treatment response [4–8]. This remarkable success story cannot (yet) be applied to $^{23}\text{Na}$ imaging, but we are witnessing continuous development.

**Physiological and technical basics**

$^{23}\text{Na}$ ions play a fundamental role in human life and can be traced – similar to protons ($^{1}\text{H}$) – ubiquitously in the human body. Fluxes of $^{23}\text{Na}$ ions in cells and across cell membranes are central part of many processes of cell activity. Up to 70% of the energy from adenosine triphosphate (ATP) hydrolysis is used for the Na+–K+–ATPase, which pumps three Na+ ions out of the cell while two K+ ions vice versa [9]. An extracellular concentration of $\approx 145$ mM and an intracellular concentration of $\approx 12$–$20$ mM [10] are maintained in healthy tissue by this pump mechanism, which leads to a mean tissue sodium concentration (TSC) of about 50 mM. Pathologic changes such as tissue injury, edema, or necrosis result in a degradation of Na+–K+–ATPase and hence in an increase in TSC from 50 mM up to 145 mM in case of cell burst [11].

From an electro-physiological point-of-view, some physical characteristics of $^{23}\text{Na}$ hamper the simple application for MR imaging. The sodium nucleus has a spin of $3/2$ and is therefore subject to quadrupolar relaxation resulting in a biexponential T2 decay with relaxation times of T2f* 0.5–8 ms and T2s* 15–30 ms for the fast and slow component, respectively [12]. Additionally, sodium MRI suffers from low in vivo concentration with a weak gyromagnetic ratio of only $1/4$ of that for $^{1}\text{H}$ resulting in an approximately 10-fold lower MR sensitivity compared to $^{1}\text{H}$ MRI with about 10,000-fold less signal. Consequently, signal-to-noise ratio (SNR)-efficient acquisition strategies with short echo times such as the radial scheme [13] are required. Many MR sequences were recently developed to acquire the k-space homogeneously yielding higher SNR: twisted projection imaging (TPI) [14], 3D cones [15], density-adapted projection reconstruction [16, 17]. Since filtering is usually applied to sodium images as a post-processing step, sampling density-weighted apodization (SDWA) with intrinsic filtering [18, 19] is preferred when using short readout durations (Fig. 1). Anisotropic 3D imaging sequences using cones [20] or twisted projection imaging [21] were recently developed for applications where anisotropic resolutions are needed (e.g., cartilage).

The technical developments of acquisition strategies and sequence design over the last decade were accompanied by MR hardware improvements. The trend to higher field strengths and stronger gradient systems was continued and led not only to a routine use of 3T MR scanners in patient care but to a growing number of 7T whole-body-MR installations worldwide. The electro-physiological characteristics of $^{23}\text{Na}$ predestine the implementation of higher field strengths. Complementary progression can be stated for coil design. Meanwhile multi-channel $^{23}\text{Na}$ coils are commercially available and experimental new designs – as a double-tuned two-port surface resonator for $^{23}\text{Na}$- and $^{1}\text{H}$-imaging [22, 23] – have been introduced.

**Oncologic therapy monitoring using $^{23}\text{Na}$ MRI – quo vadis?**

Taking into account the above-described technical developments over the last few years, a kind of renaissance of $^{23}\text{Na}$-MRI and the determination of tissue sodium content (TSC) can be stated. Feasibility of $^{23}\text{Na}$-MRI for *in vivo* imaging of physiological conditions has been demonstrated in various parts of the human body, e.g. kidney [24–26], cartilage and musculoskeletal in general [12, 27–29], brain [30, 31] heart [32–34] and prostate [35]. Initial translation from physiology to pathophysiology was correspondingly addressed in a broad spectrum of organs and different pathologies. The possibility of imaging of transplanted kidneys [36] and detection of renal changes after 3 dimensional con-

formal radiotherapy in a long-term follow-up in patients after gastric cancer [37], has been shown. In musculoskeletal imaging, for example, different cartilage repair approaches in the knee were evaluated with $^{23}\text{Na}$-MRI at 7T [38–40] and presented marked differences in comparison to native cartilage. Increased sodium concentrations were found in different brain tumors relative to normal brain structures [41, 42]. An up to threefold increase in TSC can be observed in human stroke [43] allowing monitoring of the progression of stroke pathophysiology [44, 45]. Surprising results revealed a study about relapsing-remitting multiple sclerosis at early and advanced stage. TSC was increased inside demyelinating lesions in both groups of patients, but TSC accumulation dramatically increases in the advanced stage, especially in the normal-appearing brain tissues, concomitant with disability [46]. Furthermore, $^{23}\text{Na}$-MRI provides a non-invasive solution to distinguish viable from nonviable myocardial tissue after myocardial infarction in an animal model [47] and in humans [48, 49]. TSC measurements have shown an increased signal mainly in nonviable myocardium after infarction due to loss of cell membrane integrity. Despite the never-ending discussion of necessity of separating intra- and extracellular $^{23}\text{Na}$ components, TSC offers a unique tool for measuring tissue viability noninvasively. The pathophysiology phenomena in almost all acute pathologies (stroke, myocardial infarction) are mainly based on the idea of changing $^{23}\text{Na}$ environments e.g. due to the loss of cell membrane integrity and the following adjustment of intra- and extracellular $^{23}\text{Na}$. Laymon et al. [50] described that cell membrane depolarization preceding the large degree of cell division in neoplastic tissue leads to an increase in the intracellular sodium concentration (ISC) and a concomitant rise in the total TSC. In human brain tumors, Ouwerkerk and co-workers showed that measured $^{23}\text{Na}$ changes within the tumors cannot only be attributed to alterations in $^{23}\text{Na}$ relaxation time, e.g. in the presence of surrounding edema, but reflect
real intrinsic changes of Na\(^+\)-K\(^-\)-pump function [41]. This research group concluded in the same work as prospect to therapy monitoring: "Therapies that alter tumor ion homeostasis or affect or destroy tumor cell membrane integrity are likely to generate changes that are observable with \(^{23}\)Na MR imaging and sodium concentration measurements. With these measurements, changes can be observed much earlier than the effects of anatomic remodeling." This idea of using \(^{23}\)Na as surrogate parameter for oncology therapy control is therefore not new and was among others also addressed by Thulborn et al. [42] in the field of management of brain tumors. The ability to quantify early effects of tumor therapeutic response using non-invasive \(^{23}\)Na-MR imaging approaches would have a major impact in clinical oncology. To date, clinical studies assessing these predicted potentials are missing, but first data, mainly derived from different animal models, apart from several tumor entities, have been published. In 2000, Kline et al. [51] detected significantly increased \(^{23}\)Na signal in mouse xenograft tumors propagated from human prostate cancer cell lines, 24h after administration of antineoplastics compared to baseline. Histopathological correlation of explanted tumors confirmed that chemotherapy reduced proliferation, inversely correlated with \(^{23}\)Na MRI response on a tumor-to-tumor basis. A logical development was the combination with another functional MR approach. Babsky et al. [52] performed \(^{23}\)Na-MRI and DWI in a mice model with subcutaneously-implanted radiation-induced fibrosarcoma (RIF-1) before, and daily for 3 days after, chemotherapy treatment. In contrast to the control group, in vivo MRI experiments showed an increase in both \(^{23}\)Na and apparent diffusion coefficient (ADC) in treated tumors, correlating to histological confirmed decreased cell density. After chemotherapy a chemical analysis showed an increased relative extracellular space and \([Na^+]\) concentration in treated tumors. Sharma and co-workers [53] evaluated at 4.23T the association between in vivo intracellular \(^{23}\)Na MRI intensities, immuno-biomarkers and histopathological features respectively, to monitor the early tumor response to chemotherapy using a rat xenograft breast tumor model. They concluded that intracellular \(^{23}\)Na MRI intensities possibly indicate chemosensitivity response in vivo associated with apoptosis and different pre-malignant features within 24 hours of exposure of cancer cells to anti-neoplastic Taxotere drug. Schepkin et al. [54] compared \(^{23}\)Na and DWI for their ability to detect early cellular changes in rats with subcutaneous 9L gliosarcomas treated with chemotherapy. Both imaging modalities were able to detect early changes (2 days post-treatment) in tumor cellularity continuing to evolve to a maximum after 8 days. Subsequent tumor shrinkage followed the functional parameters. The authors concluded that therapeutically-induced changes in \(^{23}\)Na and DWI were found to have similar dynamic and spatial changes and detect similar early cellular changes after treatment. The same research group demonstrated the sensitivity and applicability of \(^{23}\)Na and DWI as tools for dynamic assessment of tumor response to therapy [55]. They detected in a 9L rat gliosarcoma model, a correlation between tumor \(^{23}\)Na and DWI to gauge tumor response to therapy with varying doses of chemotherapy. In summary, all animal studies confirmed the possibility to detect tumor changes with \(^{23}\)Na imaging after oncologic therapy as correlation of treatment success. But the really astonishing finding of these preclinical studies is the indication that \(^{23}\)Na MRI could develop into an early predictor of therapy response within the first 24h. If these results were to be affirmed in human studies, it could lead to a major medical

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Image examples, including (2A) \(^{23}\)Na-MRI, (2B) morphological T2w HASTE (Half fourier-Acquired Single shot Turbo spin Echo) and (2C) a colored fusion of both sequences, of a 66-year-old male patient with lung cancer (stage IV adenocarcinoma). The lung cancer in the right lower lobe is marked with an arrow. Courtesy of PD Dr. Thomas Henzler, University Medical Center Mannheim, Heidelberg University, Germany
and economical impact on oncological therapy control and generate an early imaging tool for personalized therapy control. One of the first in vivo translation into human pathologies in oncology treatment monitoring was reported by Henzler et al. [56], who showed the feasibility of $^{23}$Na-MRI in patients with lung cancer (Fig. 2). In this feasibility study, three patients with stage IV adenocarcinoma of the lung were enrolled and multimodal examined. Data were available of $^{23}$Na- and non-contrast enhanced $^1$H-MRI, CT and $^{18}$F-FDG-PET-CT. One of the three included patients was chemo naive and examined before and after the initiation of combination therapy. Fusion of $^{23}$Na-MR images with $^1$H-MRI, CT and FDG-PET-CT was feasible in all patients and showed differences in solid and necrotic tumor areas. Between the two exams the signal intensity of the tumor as well as the ratio of signal intensity between the tumor and the CSF slightly increased indicating early therapy induced changes within the tumor. The authors concluded that $^{23}$Na-MRI is feasible in patients with lung cancer and could provide valuable functional molecular information regarding tumor viability, and potentially a treatment response. 

A second example was recently published by Layman et al. [50] from the University of Pittsburgh. This research group aimed in their feasibility study to implement and compare $^{18}$F-FLT positron emission tomography (PET), $^{23}$Na and morphological MRI at 3T in patients with glioblastoma multiforme. Two patients underwent repetitive scans at baseline (before therapy), at an early and a late follow-up time point after beginning therapy. Both functional methods were registered to the morphological MRI and calculated on a voxel-wise basis to address the heterogeneity of tumor physiology. Both – $^{18}$F-FLT PET and $^{23}$Na-MRI – independently presented changes of the tumor tissue varying in different regions, as a function of scan time point. But these initial results indicate that the two functional modalities may provide complementary information regarding tumor progression and response. The authors stated that, unlike $^{18}$F-FLT uptake, changes in sodium concentration occur without limitations from the state of the blood brain barrier. But the final value of $^{23}$Na MRI in these patients and the possibility to discriminate tumor progression from pseudoprogression requires additional patient data and outcome control. Undoubtedly $^{23}$Na MRI is an auspicious
functional technique, for which the preclinical animal and first in vivo human data show a huge potential in the field of oncology treatment. However, this technique clearly still requires a special and sophisticated technical setup, which up to now is only available in a select number of research centers worldwide.

**Integrated concepts of functional therapy +/- monitoring**

The great challenge for the coming years is to translate this additional diagnostic information into a more effective, less invasive therapy for the patient with fewer side effects and at the same time higher cost-effectiveness rendering it more affordable for the general health care system. This implies that imaging is specific for the mechanism of the disease and the target of the therapy on one hand and provides a complete picture of the systemic spread and thus the stage of the disease on the other. For this, the critical gap between modern molecular histopathology, molecular imaging and image-guided, minimally-invasive therapy has to be bridged and the results transferred from basic science into clinical routine. This challenge cannot be comprehensively addressed by a single research group, but requires the close interaction of scientists from multiple disciplines of medical imaging and from different types of academic institutions as well as industry in close proximity on a medical university campus. Currently, industry on campus proximity on a medical university campus in a long-term private-public patient-focused research on a single campus in Germany are specifically designed to support German Federal Ministry of Research (IOC) initiatives such as the one from the University Medical Center Mannheim, Heidelberg University for providing Figure 2. We thank PD Dr. Thomas Henzler, University Medical Center Mannheim, Heidelberg University for providing Figure 2.

**Conclusion**

Taking into account the three basic implications deduced in the introduction, \(^{23}Na\)-MRI undeniably has still a long way to go. But – also undeniably – \(^{23}Na\)-MRI clearly shows promise as an outstanding new approach for measuring tissue viability non-invasively. And its full potential is by no means exhausted. This technique can develop into a non-invasive avenue of therapy monitoring for a variety of diseases, particularly, but not solely, in oncological settings.

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We thank PD Dr. Thomas Henzler, University Medical Center Mannheim, Heidelberg University for providing Figure 2.

**References**

16. Nagel AM, Laun FB, Weber MA, Matthies C, Semmler W, Schad LR. Sodium MRI using a...
19 Stobbe R, Beaulieu C. Advantage of sampling density weighted apodization over postacquisi-
22 Wetterling F, Hogler M, Molkentin U, et al. The design of a double-tuned two-port surface reso-
23 Wetterling F, Corteville DM, Kalicycian R, et al. Whole body sodium MRI at 3T using an asym-
26 Wetterling F, Hogler M, Molkentin U, et al. The design of a double-tuned two-port surface reso-
27 Wetterling F, Corteville DM, Kalicycian R, et al. Whole body sodium MRI at 3T using an asym-
34 Thulborn KR, Davis D, Snyder J, Yonas H, Kassam A. Sodium MR imaging of acute and subacute stroke for assessment of tissue viability. Neuro-
35 Thulborn KR, Gindin TS, Davis D, Erb P. Comprehen-
sive MR imaging protocol for stroke management: tissue sodium concentration as a measure of tis-
36 Zaaraoui W, Konstandin S, Audoin B, et al. Distrib-
40 Laymon CM, Oborski MJ, Lee VK, et al. Com-
43 Sharma R, Kline RP, Wu EX, Katz JK. Rapid in vivo Taxoterore quantitative chemosensitivity response by 4.23 Tesla sodium MRI and histo-
immunostaining features in N-Methyl-N-Nitro-
47 Gerlinger M, Rowan AJ, Horswell S, et al. Intra-
48 Kunz T, Gupta R, Schonberg SO, Semmiller W, Kachelriess M, Bartling S. Real-time X-ray-based 4D image guidance of minimally invasive inter-

Contact Stefan Haneder, M.D.
Institute of Clinical Radiology and Nuclear Medicine
University Medical Center Mannheim
Heidelberg University
Theodor-Kutzer-Ufer 1–3
68167 Mannheim
Germany
Phone: +49/621/383 2067
Fax: +49/621/383 1910
stefan.haneder@umm.de
A novel 2D Pulsed Arterial Spin Labeling Protocol for Pediatric Cases with Brain Tumor

Yang Wang, M.D.¹; Chang Y. Ho, M.D.¹; Josef Pfeuffer, Ph.D.²

¹Center for Neuroimaging, Department of Radiology and Imaging Sciences, Indiana University School of Medicine, Indianapolis, IN, USA
²Siemens Healthcare, MR Applications Development, Erlangen, Germany

rCBF curve using different TI allows assessment of the rCBF considering pathological alteration in regional arterial transit time.
**Introduction**

Arterial spin labeling (ASL) is a non-invasive MRI technique that does not rely on ionizing radiation for measuring regional cerebral blood flow (rCBF). ASL can be applied as an attractive alternative method for measuring perfusion in pediatric brain tumor patients where it is desirable to minimize the use of ionizing radiation and Gd-based contrast agents. Compared with conventional dynamic susceptibility contrast (DSC) techniques, ASL can be quantitative enabling the measurement of global hypo- or hyper-perfusion as well as absolute perfusion changes in longitudinal studies. However, to date ASL has not been widely utilized in clinical practice. Quantification of rCBF using ASL perfusion MRI requires several experimental and physiological parameters to be properly accounted for. Variable bolus transit time and post-labeling delay are two confounding factors that may compromise the quantitative accuracy of perfusion estimates [3]. While the widely used PICORE Q2TIPS method of pulsed ASL (PASL) with a fixed temporal width tagging bolus enables quantitative estimates of rCBF from measurements taken at a single inversion time (TI), one of the key assumptions of Q2TIPS is that TI is sufficiently long for the trailing edge of the tagged bolus to have reached the imaging voxel [4]. Empirical observations suggest that transit time can be altered in different physiological and/or pathological conditions [1, 3, 5]. Therefore, we have developed a new PASL protocol that fits the single compartment model to multiple TI acquisitions to reduce artifacts caused by spatially variable bolus transit times.

**Theory**

PASL uses an inversion pulse and a tag saturation module to define the temporal duration of the tag (e.g. QUIPSS or Q2TIPS methods) [8], where a single compartment model can be used to estimate rCBF [2].

In this model $\Delta M$ is the signal difference between label and control images, $T_I$ is the total inversion time between the label pulse and the readout of the proximal slice, $T_i$ is bolus cut off time (also defines bolus duration), $\tau$ is the arterial transit time (ATT) referring to the time it takes the arterial blood to travel from the labeling site to the capillaries in the issue being imaged, $\alpha$ is the longitudinal relaxation time of blood, $M_0$ is the acquired map of equilibrium magnetization of arterial blood. This model assumes no exchange of labeled blood water into the tissue. Using this model, the rCBF map can be calculated using equation (1) [7], where $\alpha = \text{blood/tissue water partition coefficient}$, $\tau = \text{bolus cut off time}$ (also defines bolus duration).

$$f = \frac{\lambda \Delta M}{2aM_0 \tau \exp(-T_I/T_0)} \quad (1)$$

Moreover, we collected data using four different inversion times $(T_I)$ with a fixed bolus cut off time $(T_i)$, then fitted those four different datasets with the theoretical perfusion curve of the single compartment model, in order to estimate rCBF and regional ATT [3].

**Methods**

PASL data were acquired using PICORE Q2TIPS sequence on a 3T MRI scanner (MAGNETOM Verio, A Tim System, Siemens Healthcare, Germany). A 32-channel head receive coil was used to increase SNR. Scan parameters: 16 slices, 5 mm thickness, dist. factor 20%, matrix 64 × 64, field-of-view (FOV) 24 cm, 6/8 partial Fourier, BW 2298 Hz/Px, TR 3000 ms, TE 13 ms, $T_i$ 700 ms, four different $T_I$ (1200 ms, 1500 ms, 1800 ms, 2100 ms) were applied twice with one scan in ascending and another one in descending slice order, each scan with 32 acq. pairs plus one $M_0$ image, total scan time of the protocol about 11 min. Inline 3D Prospective Acquisition
Correction (PACE) was used during all PASL scans to minimize head motion artifacts [6]. In post-processing, perfusion-weighted images were calculated for each TI2 acquisition and combined from two scans with different slice order. Then four different perfusion-weighted images were fitted into the single compartment model to estimate ATT and rCBF, as described elsewhere [1, 3].

Case report
A 16-year-old male presented with slurred speech and blurred vision. MR morphological imaging shows a heterogeneously enhancing tumor centered in the left putamen with a final pathological diagnosis of primitive neuroectodermal tumor (PNET). Both PASL and DSC perfusion are concordant in demonstrating regional increased CBF within the tumor, consistent with a high grade neoplasm. Using our novel PASL protocol, we were able to generate different rCBF maps at different TI, and more precisely assessed the CBF, considering the pathological alteration in regional ATT (Fig. 1).

Conclusion
The development of a robust, quantitative ASL protocol is critical for estimation of CBF in research and clinical applications. A number of studies have measured multiple TI times for the same slices and demonstrated in healthy human brain that the model provides unbiased rCBF estimates in some regions, but in other regions considerable bolus dispersion and latency have been observed [3]. Moreover, acquisition of multiple TI will considerably increase the exam time and might not be feasible in some clinical applications. In this report, a novel PASL multiple-TI protocol is introduced using a 32-channel coil to preserve SNR at clinically acceptable measurement durations. Brain tumors remain the second most common tumor in the pediatric age group after leukemia. It is well known that malignant brain tumors typically cause neoangiogenesis due to increased metabolic demand, resulting in an increase in rCBF. By developing a robust, quantifiable ASL protocol tumor grading – and therefore clinical management – can be more accurately accomplished without the use of gadolinium injection. The novel PASL multiple-TI method enables us to more accurately estimate rCBF in pediatric tumor cases at clinically acceptable examination time. Future research is needed to determine whether this method may be useful for doctors in making treatment decisions and monitoring patients’ response to treatment.

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References

Contact
Yang Wang, M.D.
Center for Neuroimaging
Department of Radiology and Imaging Sciences
Indiana University School of Medicine
355 W. 16th Street, GH Suite 4100
Indianapolis, IN 46202-7176
USA
Phone: +1 (317) 963-7506
Fax: +1 (317) 963-7547
ywang1@iupui.edu

4MR scanning has not been established as safe for imaging fetuses and infants under two years of age. The responsible physician must evaluate the benefit of the MRI examination in comparison to other imaging procedures.

WIP – Works in progress: The new PASL protocol is currently under development; it is not for sale in the U.S. Its future availability cannot be ensured.

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We have been using the 1.5T MAGNETOM Aera for a year now and it is perhaps time to reflect on how the system is performing in a busy environment. My experiences with the MAGNETOM Aera are summarized in this article. Compared to our earlier 1.5T MAGNETOM Symphony a Tim system, MAGNETOM Aera’s new RF system – Tim 4G – introduces signal increase that is used to scan at higher resolution than before whilst keeping the same acquisition time or can be combined with parallel imaging for higher speed.

Section 1: fast brain imaging with Tim 4G and Dot

In our center we use eight sequences for brain evaluation in a 20-minute time slot that includes getting the patient in and out of the MRI suite. Figure 1 shows the details of the brain protocol we follow. The Brain Dot Engine helps us to acquire all brain examinations with consistent accurate coverage from the foramen magnum up to the vertex and from left to right or anteroposterior without fault, which is possible irrespective of the experience level of the operators. Moreover, Brain Dot Engine starts with an AutoAlign 3D localizer which is mapped to the Talairach space, and as a result sequence planning is automated according to the reference space independent of patient positioning / habitus.

This is especially convenient in follow-up examinations, such as in this case of demyelinating disease or tumor, resulting in a more reliable and comfortable comparison of the examinations at different time points (Fig. 2).

Sequence details of our 20-minute brain evaluation:
1. T2*w EPI, 23 slices, 5 mm in 9.7 sec, 0.9 x 0.9 x 5 (FOV 185 x 220, matrix 216 x 256) res
2. T1w sag SE, 4 mm slices in 2:09 min, 0.7 x 0.7 x 4 (FOV 290 x 223, matrix 246 x 320) res
3. T2w ax TSE, 25 slices, 4 mm in 01:34 min, 0.5 x 0.5 x 4 (FOV 171 x 221, matrix 256 x 384)
4. T2w FLAIR, 23 slices, 5 mm in 32 sec x 2 0.8 x 0.8 x 5 (FOV 211 x 211, matrix 205 x 256)
5. DWI EPI b50, b500, b1000, 69 slices in 1:30 min (1.2 x 1.2 x 5, FOV 216 x 231, matrix 174 x 188)
6. 3D TOF 3 slabs, 116 slices in 2:44 min, 0.4 x 0.4 x 0.5 mm isotropic (FOV 141 x 181, matrix 376 x 512)
7. T1w cor TSE, 22 4 mm slices in 1:34 min, 0.8 x 0.8 x 4 (FOV 174 x 200, matrix 213 x 256)
8. T1w ax SE 22 4 mm slices in 2:34 min, 0.9 x 0.9 x 5 (FOV 174 x 200, matrix 213 x 256)
Section 2: 16-channel MSK coils

The new RF system enables the use of 16-channel MSK coils like the Shoulder, Foot/Ankle and Hand/Wrist coils. The new wrist coil gives abundant signal and superb detail. The wrist coil has grown, in fact, to a 16-channel hand-wrist coil (Fig. 3), which is quite convenient for both the patient and the operators and helps faster setup times. Many patients have combined hand/wrist pathology and symptoms, and extended coverage is a huge benefit. Since it is easy to position the hand in the Hand/Wrist coil, examining the fingers has become very straightforward. Figure 4 shows a young butcher who suffered from a deep wound in the index finger. You still can notice the susceptibility artifacts on the coronal STIR and GRE (Figs. 4D, E). One week later the young man can no longer flex the distal interphalangeal joint (DIP) of the index finger. MRI nicely depicts the torn deep flexor retracted to the proximal phalanx in the tendon sheet (Figs. 4H, I). All these images have an in-plane resolution of 0.5 mm, a slice thickness of 2.3 mm or less and scanning times of 1.30 min or less thanks to the higher signal from the high-channel coil. Since this coil is so successful for fingers, it could also be used for toes. This is the flexibility with Tim.

Figure 5 shows images of a middle-aged man, who suffers from a “tingling pain sensation” in the forefoot intermetatarsal space III-IV and numb feeling in digit IV. A T2 hyperintense, T1 hypointense dumbbell shaped lesion bulging between the metatarsal heads III-IV is clearly demonstrated (Figs. 5A, B, C) confirming the clinical suspicion of Morton neuroma.

Section 3: reduction of susceptibility artifacts caused by metal

MRI is challenging in the presence of metal. MAGNETOM Aera came equipped with syngo WARP with high bandwidth TSE and TSE STIR protocols and an
Superb details in a case of trauma to index finger in 1:30 min per sequence.
optional VAT (View Angle Tilting) technique. In Figures 6–8 I share some examples of real day-to-day cases. Figures 7 and 8 show images of a female patient who has had osteosynthesis for tibial plateau fracture following a skiing accident 20 years ago. She had an MRI of the knee, after another skiing accident. WARP helped reduce susceptibility artifacts for better image quality and confidence in diagnosis.

Section 4: faster imaging with CAIPIRINHA

CAIPIRINHA (Controlled Aliasing in Parallel Imaging Results in Higher Acceleration) is a new parallel imaging technique from Siemens.

The new RF system and high channel coils (Body 18) provide the signal that is needed, whilst the parallel imaging with fourfold acceleration with CAIPIRINHA offers the speed required to maintain short breath-holds even at large coverage in the z-direction and thin slices. The short breath-holds are a real game changer especially in the case of elderly patients, but even young and fit patients benefit.

No trade-off between thick slice / large coverage or thin slice / partial coverage is necessary: we always scan full coverage with thin slices. Examination of the abdo-
9 CAIPRINHA PAT4 in 10 sec compared to PAT2 in 21 sec with same slice thickness and in-plane resolution.

10 With CAIPRINHA no trade-off between thick slice / large coverage or thin slice / partial coverage is necessary: we always scan full coverage with thin slices. The coverage is well seen in this screenshot (yellow dotted box top left, or yellow bold box top middle).
men and thorax benefits greatly from the Aera system. It all fits perfectly together. Here are some examples.

More signal in less scanning time!

Figure 9 demonstrates how, in a liver examination in the same patient examined with CAIPI4, the images with a breath-hold of 10 sec have more signal than the same images (same slice thickness and in-plane resolution) with iPAT2 and 21 sec breath-hold.

MRI examination of the thorax – although more rare – follows the same rules as the abdomen: sequences should be fast enough for the patient to cooperate. Figure 11 gives an example of a metastatic lung carcinoma where PET-CT was not able to exclude or confirm chest wall invasion. MRI was ordered and it turned out to be quite an easy job on MRI, having the short breath-hold CAIPIRINHA PAT 4 VIBE sequences. Even this patient, who clearly is in a bad condition,
Clinical Head-to-Toe Imaging

Coronal dynamic sella imaging with CAIPI PAT4 VIBE in 34 seconds.

Screenshot showing 1.41 min CAIPI VIBE Dixon for 3D shoulder imaging. High SNR due to the 16-channel MSK coils allows us to use these fast sequences.

cooperated and tolerated the exam well. Invasion in the lateral and dorsal chest wall is readily depicted in both axial and coronal enhanced images. Axial T2w BLADE and free breathing diffusion-weighted imaging (DWI) confirm the chest wall invasion with actual rib invasion and the malignant nature is clearly translated in low ADC values of the lesion (Fig. 12). Maybe we don’t have the fancy colors, like PET does, but we do have the diagnosis!

CAIPIRINHA can be used in other regions, too. Figure 13 shows images of a female patient who underwent a brain scan in search of a (post-traumatic?) cause of the headaches. A small mass in the sella was incidentally found. Coronal dynamic VIBE imaging provides a means to assess the perfusion of the sella and pick up smaller lesions e.g. adenomas as demonstrated in this examination. The VIBE sequence is pimped using CAIPIRINHA with 4-fold acceleration, providing 40 high res 1.5 mm slice thickness images every 34 seconds, and dramatically improving detection rate. The acquisition speed makes it possible to complete this examination within the same 20-minute time slot. At our center, shoulder examination is mostly done after arthrogram. This can cause some discomfort and the fast scanning of patients is the best option to avoid movement artifacts. syngo BLADE helps reduce motion artifacts. However CAIPIRINHA is a huge benefit here. We acquire a 3D VIBE Dixon fatsat with CAIPI PAT4 in 1:41 min. The screenshot in Figure 14 shows the details of our shoulder protocol.

Section 5: the Large Joint Dot Engine for shoulder imaging

The AutoAlign tool in the Dot Engine helps technologists to plan more quickly and accurately, with reduced operator-depen
Shoulder Dot screenshot showing Inline MPR planning and VIBE CIPI reformat.

Section 6: liver imaging with Abdomen Dot Engine

Dot has many smart tools that help improve efficiency, consistency, reproducibility and throughput. AutoAlign is one tool I mentioned above in Brain Dot. The Abdomen Dot Engine actually provides several operator-independent tools: with AutoAlign for liver, and AutoCoverage, you always have the complete anatomy covered, ensuring good quality images with fewer incidences requiring repeat scans. This is a significant benefit of Dot and it holds true when we image obese patients, too.
MAGNETOM Aera’s 70 cm open bore enables some larger patients to fit for the first time in their lives inside an MRI machine. But even if we did manage to fit such a patient inside the previous MRI scanner, we still ran into the problems of too much noise and of breath-holds that were too long to cover all the phase encoding steps needed in an obese patient. The increased signal available on the Aera tackles the noise problem and the CAIPIRINHA technique gives you a fourfold parallel imaging factor (without additional noise) resulting in acceptable breath-hold times (10.7 sec) even in large patients. Again, it all fits perfectly together!

Abdomen Dot enables easier, faster and more consequent – less operator-dependent – sequence planning and execution with AutoCommand tools where breath-hold commands are given by the system in a language the patient understands. The Abdomen Dot features ABLE (add-in for Automatic Breath-hold Liver Exams) where the system actually triggers on bolus arrival to ensure a pure arterial phase even in cirrhotic liver patients where timing of the bolus could be a challenge for less experienced technologists (Fig. 17).

Non-rigid liver registration of dynamic VIBE series is an automatic step done within the ABLE function to save all phases registered in the database, which enables faster and more accurate reading and reporting. Different contrasts scans (b50 – b800 DWI and ADC map & T2w BLADE) can be in similar anatomical positions as close as possible to the multi-phase dynamic scans due to smart AutoCoverage functionality in Abdomen Dot. The example of a cirrhotic liver in Figure 18 will clearly illustrate the efficiency in reading. This holds true within the scope of a single examination but becomes even more important when comparing examinations at different time points (Fig. 19). Having imaged and sent the images to the PACS in registered series turns out to be of great benefit for reading and reporting in follow-up examinations. This registration process allows for synchronized scrolling up and down simultaneously in the registered series even in a bare bone viewing system used by, for example, the referring physician. Figure 19 shows how convenient it was to reevaluate the liver on MRI in a case of invasive adenocarcinoma of the sigmoid colon. Comparing exams from 2011 and 2013 even on a bare bone viewing system: the Dot registration provides all the synchronization that is needed.

**Section 7: excellent images in the head/neck region**

Whereas in the old days fatsat could be problematic in the cervicothoracic region, TSE Dixon really assists with impeccable fatsat images and having the T2w or T1w TSE images for free.

Figure 20 gives a nice example of metastatic melanoma after gadolinium enhancement: no artifacts in the base of the neck! DWI has also improved due to the gradient power enabling short TE thus minimizing artifacts.

The next two examples of DWI in the neck highlight the importance of ADC maps. High signal on the high b-value DWI and low ADC value confirm the malignant nature of the metastatic melanoma. DWI can help in determining the nature of lesions, such as in the small benign mixed tumor (BMT) of the right parotid gland with high ADC value depicted in Figure 22.
Section 8: leveraging syngo.via

Presenting the roadmap to vascular surgeons has been made easier and better with syngo.via. The MIP and composing functionality helps us to integrate three 4D datasets for angiography of the aorta and lower limbs into one dataset and to present the roadmap to the vascular surgeons – something that we do in every case. Cardiac evaluation is done on syngo.via and, again, instead of having to leave my reporting system to go to the workstation, I simply push the syngo.via button and do my evaluations. Another area where I use syngo.via is for volume calculations. Volume calculating helps me out with every acoustic neuroma and for prostate (benign prostate hypertrophy). The inline registration in the Dot engines really helps with the multi timepoint follow up of 90% of cases: it gives me the series synchronized and sends them in that registered way to my non-intelligent, bare bone viewing station (the reporting
system). This synchronization-registration assists in most of the brain and abdomen cases. syngo.via helps in reading difficult cases, where I like to compare, for example, not only the registered axial slices but also coronal slices, the free breathing DWI and other contrasts. When this is required, I simply push the syngo.via button on my viewing station and get my patient opened up in syngo.via. It makes reading and reporting not only faster but also more accurate.

Conclusion

The new RF, high-channel coils, gain in SNR, faster techniques like CAIPIRINHA, new metal implant imaging possibilities, Dot features and syngo.via all are a perfect fit and the MAGNETOM Aera truly combines throughput and highest quality MR imaging in an optimized clinical workflow.

Contact
Johan Dehem, M.D
VZW Jan Yperman
Ieper
Belgium
johan.dehem@gmail.com

Further information
Visit us at www.siemens.com/magnetom-world to listen to Dr. Dehem’s talk on Highest Quality Imaging in an Optimized Clinical Workflow given during the lunch symposium at the 15th International MRI Symposium MR 2013 in Garmisch-Partenkirchen, Germany.

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Initial Experience with Whole-Body Diffusion-Weighted Imaging in Oncological and Non-Oncological Patients

Marcos Vieira Godinho, M.D. 1; Romulo Varella de Oliveira, M.D. 1; Clarissa Canella, M.D., MSc 1; Flavia Costa, M.D., MSc 1; Thomas Doring, MSc 1; Ralph Strecker, Ph.D. 2; Romeu Cortes Domingues, M.D. 1; Leonardo Kayat Bittencourt, M.D., MSc 1

1 Clínica de Diagnóstico por Imagem, Rio de Janeiro, Brazil
2 Siemens Healthcare, São Paulo, Brazil

Background
Initially used as a diagnostic tool for acute stroke, over recent years diffusion-weighted imaging (DWI) has developed a growing role in the diagnosis and follow-up of diseases not only in extra-cranial pathologies, but also as a study of the whole body in systemic diseases, providing a steady increase of indications for the technique. With a growing elderly population worldwide and the highest incidence of oncologic illnesses, use of whole-body diffusion-weighted imaging (WBDWI) in this setting is evolving.

We initially used WBDWI for the already established indication in oncological patients with breast, prostate, gastro-intestinal, hematological and other cancers, and then also decided to investigate its use in patients with auto-immune, inflammatory and infectious diseases.
The purpose of this article is to describe our examination technique, imaging protocols, interpretation of imaging findings and initial experience with oncological and non-oncological diseases, as well as the limitations of the study.

Methods

Since September 2011 we have recruited patients with oncological disease undergoing diagnostic or follow-up exams in our clinic with findings in conventional MRI suggestive of metastases, as well as patients with auto-immune or inflammatory systemic diseases and those with imaging findings that could be differential diagnosis for metastatic disease. In some cases the purpose was the staging of the disease before initial treatment, in other cases we did the follow-up of patients during therapy.

WDBDI was performed on a 1.5 T clinical scanner (MAGNETOM Aera, Siemens Healthcare, Erlangen, Germany) recently installed in our center using two 18-channel body phase array coils together with the combined 20-channel Head/Neck coil. Patients were examined in supine position and four stations of axial free-breathing DWI with short tau inversion recovery (STIR) fat suppression were obtained using the following parameters: echo time (TE) 79 ms, repetition time (TR) 19000 ms, inversion time (TI) 180 ms, b-values 0 and 800 s/mm², readout bandwidth 1502 Hz/pixel, 5 averages, field-of-view (FOV) 400 mm, slice thickness (SL) 5 mm, matrix size 128 × 128, voxel size 3.1 × 3.1 × 5.0 mm³, acquisition time per station 7.17 min.

The protocol also included four stations of axial T1-weighted (T1w) VIBE in- and opposed-phase images (TR 7.01 ms /TE1 2.38 ms /TE2 4.76 ms, FOV 450 mm, SL 5 mm, matrix 512 × 512, voxel size 0.9 × 0.9 × 5 mm³) and axial T2-weighted (T2w) HASTE images with fat suppression STIR (TR 600 ms /TE 81 ms, FOV 420 mm, SL 5 mm, matrix size 256 × 256, voxel size 1.6 × 1.6 × 5 mm³). Head, neck, chest, abdomen, pelvis and proximal thighs were studied. The four stations were composed inline, after the last station was acquired. No contrast media was administered and total duration of each exam was about 60 minutes.

So far, we have performed 4 exams with healthy controls, 9 exams with actual oncological patients (1 breast cancer, 1 small cell cancer with unknown primary site, 3 prostate cancers, 2 gastrointestinal neuroendocrine cancers, 1 plasmacytoma, 1 lymphoma) and 4 exams with non-oncological cases (1 prostatitis, 1 dermatopolymyositis, 1 spondylarthrosis, 1 chronic recurrent multifocal osteomyelitis).

Imaging findings

Our objective was to look for findings based on the principle that some neoplasms and their metastases have restricted diffusion because of their hypercellular nature. Inflammatory/infectious illnesses may also have foci of restricted diffusion along the body because some of them may produce abscesses. We always tried to detect lesions and to characterize their distribution. In other patients we also analyzed b0 images looking for the T2 shine-through effect, as explained further below.

For interpretation of WDBDI we used ‘raw’ data with inverted gray scale in b0 and b800 images, so lesions with restricted diffusion appeared as black foci in a white background (Figs. 4B and 6). In some cases, we also obtained the calculated b1400 images, improving the specificity of the findings, as in this case only lesions with real restricted diffusion were seen. Apparent Diffusion Coefficient (ADC) Maps were obtained for all patients and ADC values were measured. We also did three-dimensional maximum intensity projection (3D MIP) reconstructions using the Siemens composer tool in all cases and used T1w and STIR images for anatomical reference and fusion with WDBDI.

Control subjects

Before initiating the study with patients, we studied healthy subjects to test and adjust the protocols, to understand normal findings in the exams, considering the presence of imaging artifacts, problems with misregistration and intrinsic limitations of the study.

Normal brain tissue, spine, spleen and testicles had restricted diffusion, notably the latter. Even in control cases, small lymph nodes free of disease appeared with restricted diffusion, mainly in the neck, a finding which we consider as a limitation of the study (Fig. 1).

In some patients a mismatch between the head/neck and the thorax station, was observed. A problem most prominently visible in the ‘broken spine’ in sagittal MPR images and described by other authors before [1]. We also observed in some cases spatial variations in image intensity between the different stations which impaired the image interpretation of MPR images. It is worth to mention here that both the broken spine artifact and inhomogeneous signal can be corrected by using the new Diffusion Mode for Composing in software version syngo MR D13A.

We did not study the whole arms, because an increase in the size of the FOV would have been necessary, which could have reduced the resolution of the images.

Oncological patients

One of the most prominent features of the WDBDI were the study of metastatic bone disease, which was well depicted in most of the oncologic patients studied, with examples of all types of distribution (Figs. 2, 3 and 5–9), some confirmed by biopsy. In one patient with breast cancer the findings of bone scintigraphy were similar to that of WDBDI. In one of the patients (small cell cancer with unknown origin) when we compared the results of PET-CT scan with WDBDI, the last one showed a greater number of bone metastases (Figs. 7–9). The performance of WDBDI was even better in the case of diffuse bone marrow involvement when we analyzed it side by side with T1w in- and opposed-phase images and with fusion of both, to help in the differentiation of the lymphomatous infiltrate from hyperplastic red bone marrow (which could also present as black foci in the inverted gray scale of WDBDI), as there was no signal reduction in T1w opposed-phase compared with T1w in-phase images (Fig. 8). The reduced ADC values obtained in the lymphoma-
tous lesions also helped us to distinguish both.

Other kinds of metastatic spread could be shown, and enlarged lymph nodes were noted in patients with lymphoma, small cell and prostate cancers, the first one with disseminated nodal disease (Figs. 7 and 9), the second one with mediastinal nodal disease (Fig. 5) and the last ones with retroperitoneal spread (Fig. 3), corroborating the natural history of the diseases. One patient with relapsing prostate cancer had also enlarged mediastinal, axillary and supraclavicular lymph nodes, an uncommon finding most seen in advanced disease (Figs. 3C and 3D). However, the depiction of enlarged lymph nodes was not homogeneous along the body, because the performance of WBDWI to show metastatic lymph nodes in the mediastinum were worse in comparison with cervical, axillary, retroperitoneal and pelvic nodal spread, probably because of respiratory and cardiac motion artifacts and pulsation artifacts from thoracic arteries. In the patient with small cell cancer PET-CT showed more mediastinal disease than the WBDWI.

In one specific case, the patient came to our institution with shoulder pain and was submitted to MRI of the shoulder, showing lesions suspected for bone metastases. The study was complemented with WBDWI that showed much more bone lesions as well as retroperitoneal and pelvic nodal disease. The pattern and combination of bone and nodal disease, together with a mass with restricted diffusion infiltrating the prostate suggested the diagnosis of metastatic prostate cancer, confirmed by biopsy (Figs. 3A and 3B).

The detection of visceral metastases, mainly liver metastatic lesions, could be well done in 3 patients (1 small cell cancer and 2 neuroendocrine gastrointestinal tumors had multiple liver metastases), also showing us the natural history of the diseases. In the patient with small cell cancer most of the liver lesions had central necrosis without restriction of diffusion and a peripheral region of restricted diffusion (Fig. 5). No lesion was found in the central nervous system in any of the patients in WBDWI neither in other imaging sequences.

The patient with plasmacytoma was accompanied during the treatment (one DWI before and one after the beginning...
4. 70-year-old male, carcinoid tumor of small bowel. Magnetic resonance enterography (4A): small bowel lesion with contrast enhancement (green arrow). Axial b800 image (4B): multiple liver metastases (blue arrows). The primary site was not well depicted in this case.

5. 67-year-old male, investigating weight loss in the last six months. Biopsy of mediastinal enlarged lymph node has shown small cell cancer. No primary site was discovered. Reconstruction of PET-CT in coronal plane (5A) showing fluorodeoxyglucose uptake in liver (arrows) and bone lesions (arrowheads) and in enlarged mediastinal lymph nodes (asterisks). Fused T1w in-phase and b800 DWI coronal images (5B and 5C): liver lesions with markedly restricted diffusion some with central necrosis (arrows), bone metastases (arrowheads) and mediastinal enlarged lymph nodes (asterisks). Bone lesions were better depicted in WBDWI, but comparison was compromised because WBDWI was done two months after PET-CT. Comparison between ADC values of one liver lesion in the first (5D) and second DWI (5E) two months after treatment: increasing ADC value of the lesion in follow-up image. Also note there are more areas of cystic/necrotic degeneration along the liver after initial treatment.
6 46-year-old female, dorsal thoracic pain. Axial b800 image (6A) and calculated b1400 image (6B) showing expansive lesion with markedly restricted diffusion in dorsal vertebral body (arrows). Biopsy revealed plasmacytoma.

7 WBDWI of a 66-year-old female with recurrent lymphoma, presenting with diffuse enlarged lymph nodes and bone marrow infiltration. STIR (7A), b800 DWI (7B), ADC map (7C) and fused STIR and b800 DWI (7D), showing conglomerate of retroperitoneal and mesenteric lymph node masses (arrows) and restricted diffusion in the bone marrow of a lumbar vertebra (arrowhead).
8 T1w in-phase (8A) and T1w opposed-phase (8B) of the bony pelvis in axial plane (same patient as in figure 7), showing diffuse bone marrow hypointensity, without signal reduction in T1w opposed-phase images, indicating tumoral infiltration instead of red marrow hyperplasia (confirmed by bone marrow biopsy).

9 Fused T1w in-phase and b800 DWI coronal images (same patient as in figure 7), showing diffuse bone marrow infiltration (arrowheads), conglomerate of retroperitoneal and mesenteric lymph node masses, and enlarged cervical, axillary, mediastinal, hepatic hilum, mesenteric, retroperitoneal, common iliac and external iliac lymph nodes (arrows). Note the spleen in left hypochondrium (asterisk).

10 62-year-old male, HIV+, fever and malaise with prior negative prostate biopsy. 3D reconstruction from WBDWI showing prostatitis with slightly restricted diffusion in prostate (red arrow), associated with two tiny liver abscesses (black arrows), enlarged lymph nodes (green arrows) and multiple spleen abscesses with markedly restricted diffusion (red arrowheads).
of the treatment), where an increase of the ADC of the bone lesions was observed after two months of the first session of radiotherapy, with a subsequent improvement in the clinical symptoms. One of the patients with small cell tumor with unknown primary site was also followed up (one DWI before and one after the beginning of the treatment) and showed an increase in the ADC values in bone and liver lesions after one week of the second cycle of chemotherapy, also accompanied by clinical improvement (Fig. 5). Unlike other authors, we didn’t use histogram analysis of the ADC values [2].

**WBDWI beyond oncological applications**

During the study a patient presented at our institution with fever and malaise investigating liver, splenic and nodal lesions one month after a negative biopsy for prostate cancer. He was submitted to a conventional MRI of the abdomen and pelvis and we complemented the exam with WBDWI. There were multiple lesions suggested of abscesses with central region presenting hypointense signal in T1w and hyperintense signal in STIR images with markedly restricted diffusion in the liver and notably in the spleen. We could also note axillary lymph nodes, as well as pelvic and retroperitoneal enlarged lymph nodes, the latter with apparent central areas of abscess formation on conventional images. There were also foci with slightly restricted diffusion in the prostate (Fig. 10). The type and pattern of distribution of the lesions suggested the diagnosis of prostatitis with septic foci to lymph nodes and solid viscera, which was confirmed by laboratory exams. Adequate antibiotics were administered and there was clinical improvement of the patient initially. During the investigation of prostatitis it was also discovered the patient was HIV positive with Acquired Immunodeficiency Syndrome, which could have helped in the spread of the septic disease.

In rheumatic patients, we mostly used the T2 shine-through effect of WBDWI in the interpretation of images. The patient with dermatopolymyositis had decreas-
34-year-old male, diagnosis of spondyloarthritis, significant pain and movement restriction of the lumbar spine. Axial $b_0$ images with inverted gray scale (12A) and standard gray scale (12B) and STIR images (12C): edema near the spinous process of a lumbar vertebra, configuring enthesitis (arrows). The study was complemented with composed STIR images in sagittal plane (12D) depicting discrete edema in interspinal ligaments in lumbar spine (arrowheads).

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In the patient with spondyloarthritis, enthesitis near the spinous processes of lumbar spine and in interspinal ligaments could be well depicted (Fig. 12), a finding described in the literature [3]. Chronic recurrent multifocal osteomyelitis (CRMO) is an autoinflammatory disorder of children and young adults that is characterized by non-bacterial osteomyelitis presenting with multifocal bone pain secondary to sterile osseous inflammation, with a relapsing and remitting course. In the study of the patient with CRMO we also included in the protocol a station for the distal thighs and legs. This patient had marked bilateral edema in metaphyses around the ankles and the knees adjacent to the growth plate, this last one being the most common location of the disease in the tubular
bones according to the literature [4], and prominent edema in the sacrum near the sacroiliac joints, a finding less common (Fig. 13). In rheumatic diseases, $b_0$ images had a similar performance in comparison with STIR images to detect muscle and bone marrow edema, as well as in the detection of enthesopathy. The $b_{800}$ images were not helpful in these cases, because the lesions had no real restricted diffusion.

**Conclusion**

WBDWI may play an important role in the near future for the detection of visceral and mostly bone metastases from many types of cancer. The method is also promising in the demonstration of a visual notion of the disease as a whole in some auto-immune and inflammatory illnesses, probably helping in the assessment of their severity. One of our colleagues is studying a larger number of patients with chronic recurrent multifocal osteomyelitis (CRMO) using WBDWI, so that we soon may have more answers about the advantages and disadvantages of WBDWI in these cases. We have not studied other possible indications of WBDWI, for example the assessment of bone marrow features in patients with other hematological non-neoplastic conditions, such as sickle cell disease and thalassemia, which is certainly a large field to be explored. Our center is not an academic institution and we didn’t enroll patients in a randomized and double-blind fashion. So far, the exact role of WBDWI in the oncological and non-oncological setting and its sensitivity, specificity and accuracy in comparison with other exams (e.g. bone scintigraphy, PET-CT, PET-MR) must be accessed by large clinical trials.

**References**

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Parallel Transmission and its Clinical Implementation: Enabling new Clinical Imaging Paradigms

Fernando Boada¹; Tim Shepherd¹; Andrew Rosenkrantz¹; Eric E. Sigmund¹; Jurgen Fütterer²; Hersh Chandarana¹; Mari Hagiwara¹; Henry Rusinek¹; Artem Mikheev¹; Mary Bruno¹; Christian Geppert¹; Christopher Glielmi¹; Josef Pfeuffer³

¹Department of Radiology NYU Langone Medical Center, New York, NY, USA
²Department of Radiology, Radboud University Nijmegen Medical Centre, Nijmegen, The Netherlands
³Siemens Healthcare

Introduction

The use of multiple and independent receive channels to accelerate MR data acquisition has enabled numerous clinical applications that are now part of the standard clinical workflow at state-of-the-art imaging centers. Parallel transmission (pTX) was, likewise, introduced as a means to enable and improve the use of extended RF excitations schemes such as multidimensional spatially-selective excitation, which, although conceptually powerful, were previously rendered impractical due to their long RF pulse durations.

The 3T MAGNETOM Skyra is the first clinical system to feature a parallel transmission architecture capable of enabling applications beyond B₁ shimming. This revolutionary capability allows enhancing the performance and reliability of several imaging sequences that are essential for advanced clinical applications. These enhancements are particularly significant at 3T, where B₁ field inhomogeneity can be problematic over large fields-of-view (FOV). We present several examples where the role of this technology demonstrates clinical benefits by enabling previously impractical imaging paradigms and/or improving the performance of existing ones.

1 A Standard b1000 image (1A) shows wrap artifact (*) not present on zoomed EPI image (1B). There is greater clarity of prostate capsule on zoomed EPI (arrow, 1A and 1B).

2 A There is greater clarity of prostate capsule (solid arrow) and of transition zone boundary (dotted arrow) on zoomed EPI image (2B) than on standard EPI (2A).
Head-to-Toe Imaging

Clinical

Skyra pTX architecture

The Skyra pTX system builds upon the existing clinical MAGNETOM Skyra platform, namely, a 3T magnet with a 70 cm patient bore diameter and parallel RF excitation via two fully independent, phase-coherent RF channels (TimTX TrueShape). The two pTX RF channels allow extended dynamic RF excitation schemes using a two-channel RF body coil. State-of-the-art imaging gradients (45 mT/m peak amplitude, 200 mT/m/s slew rate) and a full complement of Tim 4G multi-channel receive coils round out the hardware configuration of the system. The scanner operates under software version syngo MR D13C providing full access to further new technologies such as Dot, syngo REVEAL etc. While there are pTX features that can be applied during the adjustments (patient-specific B1-shimming, volume-selective B1-shimming), we have focused on the improved sequences and applications which provide new features with pTX. This includes 2D-selective RF pulses with an echo-planar TX trajectory that excite a selective volume (so-called ‘inner-volume’) and allows to reduce the FOV in the phase direction [1, 2] for zoomed echo planar imaging (EPI) (syngo ZOOMit).

Prostate imaging

Diffusion-weighted imaging (DWI) is a critical sequence for prostate cancer detection and localization, but often suffers from spatial-distortion EPI artifacts related to the field inhomogeneity induced by the neighboring air spaces; these challenges can be more problematic at 3T. In this challenging imaging scenario, 2D-selective zoomed EPI in conjunction with a more homogeneous localized (B1-shimmed) RF excitation can be particularly helpful. The impact of zoomed EPI on image quality of prostate DWI at 3T was assessed in six volunteers who underwent prostate MRI using an 18-channel body matrix receive coil. Scans included a single-shot EPI DWI sequence (b-values 50, 500, and 1000 s/mm²) performed with a regular sinc pulse (‘Standard’) and with the advanced 2D spatially selective RF pulse (‘ZOOMit’) combined with a reduced FOV approach (zoomed EPI). The b1000 images and ADC maps were assessed for various image quality measures on a scale from 1 to 5 (5 = highest image quality). Also, peripheral zone (PZ) ADC and estimated signal-to-noise (eSNR: determined as mean/SD of PZ) on b1000 images were measured. These measures were compared between standard and zoomed EPI.

Compared with standard EPI, zoomed EPI b1000 images showed improvements in ghosting, wraparound artifacts, clarity of prostate capsule, and clarity of peri-urethral region (Fig. 1, Table 1). By contrast, zoomed EPI ADC maps showed improvements in clarity of prostate capsule and overall image quality (Fig. 2). eSNR was nearly identical between standard and zoomed EPI b1000 images. The prostate showed a small increase in mean ADC on zoomed EPI images (mean increase 0.07 x 10⁻³ mm²/s). However, ADC reproducibility between standard and zoomed EPI DWI remained high (mean coefficient of variability of ADC (4.4 ± 4.0)%, range 0.3 to 11.0%).

This preliminary assessment showed improvements in numerous measures relating to artifacts and anatomic clarity

Table 1: Comparison of standard and zoomed DWI.

<table>
<thead>
<tr>
<th>Feature</th>
<th>Standard EPI</th>
<th>Zoomed EPI</th>
</tr>
</thead>
<tbody>
<tr>
<td>b1000 s/mm² images</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Absence of ghosting artifact</td>
<td>3.8 ± 0.4</td>
<td>4.3 ± 0.5</td>
</tr>
<tr>
<td>Absence of wrap artifact</td>
<td>3.7 ± 0.8</td>
<td>4.7 ± 0.5</td>
</tr>
<tr>
<td>Clarity of prostate capsule</td>
<td>4.0 ± 0.0</td>
<td>4.8 ± 0.4</td>
</tr>
<tr>
<td>Clarity of peri-urethral region</td>
<td>3.5 ± 0.5</td>
<td>4.2 ± 1.0</td>
</tr>
<tr>
<td>Overall image quality</td>
<td>3.7 ± 0.5</td>
<td>4.0 ± 0.6</td>
</tr>
<tr>
<td>ADC maps</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Reduced distortion of prostate</td>
<td>3.5 ± 0.5</td>
<td>3.8 ± 0.4</td>
</tr>
<tr>
<td>Clarity of transition zone boundary</td>
<td>3.2 ± 0.8</td>
<td>3.5 ± 0.4</td>
</tr>
<tr>
<td>Clarity of prostate capsule</td>
<td>3.2 ± 9.4</td>
<td>3.8 ± 0.4</td>
</tr>
<tr>
<td>Overall image quality</td>
<td>3.2 ± 0.4</td>
<td>3.7 ± 0.5</td>
</tr>
<tr>
<td>ADC (x 10⁻³ mm²/s)</td>
<td>1.34 ± 0.3</td>
<td>1.41 ± 0.8</td>
</tr>
</tbody>
</table>

¹Firevoxel and Igor Pro are not Siemens Healthcare products. Siemens bears no responsibility for these products including, but not limited to, their regulatory status.
when applying 2D-selective zoomed EPI with for prostate DWI at 3T. While the FOV was held fixed to ensure comparability between the sequences, the use of a smaller FOV is easily achieved with zoomed EPI and currently under investigation. Nonetheless, even without this adjustment, zoomed DW-EPI using 2-channel pTX has potential to improve image quality for DWI of the prostate at 3T.

Renal imaging
Diffusion-tensor imaging (DTI) uses multiple diffusion sensitizing directions to evaluate anisotropic microstructure and is a promising imaging technique for the functional assessment of kidneys. Microstructure is a key factor in renal physiology, where cortex contains randomly oriented structures, while medulla holds more aligned vessel and tubular networks. Kidney DTI has shown that renal medulla has inherently higher fractional anisotropy (FA) in comparison with the isotropic cortex [2-4]. However, kidney DTI using single-shot spin-echo EPI typically suffers from imaging artifacts such as field-inhomogeneity-related distortions, and low spatial resolution due to the matrix size and long echo trains required for abdominal imaging. In this context, shortening the echo train length could reduce the aforementioned distortions. To achieve the same level of spatial resolution, while keeping a shorter echo train length, the imaging FOV must be reduced along the phase direction by the same factor used to shorten the echo train length. Zoomed EPI achieves such a goal through the use of 2D-selective RF excitation.

Four consecutive healthy volunteers without any history of renal disease (three males, one female; mean age 28 (24-30 years); non-fasting conditions) were imaged on the Skyra pTX system. Free-breathing single-shot 2D-selective zoomed EPI DW images were acquired with reduced FOV in the left-right direction with the following parameters: 10-14 coronal slices, slice thickness 4 mm, no gap; FOV of 83 x 400 mm; 64 x 308 acquisition matrix, 1.3 mm resolution, TR 4000 ms; TE 65 ms; six diffusion directions; monopolar diffusion gradients; two b-values (b0 and b500 s/mm²); parallel imaging (GRAPPA) factor of two and scan time 5:28 min. An additional saturation band was applied to suppress signal from the left side of the body extending slightly beyond the stop-band. Right kidney images were coregistered by using a two-dimensional rigid body transform algorithm, and MR images at the same b-value and direction were then averaged by using locally developed software (Firevoxel®). DT processing was performed with custom software written in Igor Pro® (Wavemetrics, Portland, OR, USA). Parametric maps were generated of DTI eigenvalues λ, and eigenvectors, mean diffusivity (MD) and fractional anisotropy (FA). The ZOOMit DTI technique allowed for high quality diffusion images of the kidney with reduced blurring and distortion (Fig. 3) compared to full FOV EPI DTI. Medullary anisotropy and radial orientation pattern was evident in all volunteers (e.g. Fig. 3). The medullary / cortical MD and FA values were 2.04 ± 0.14 / 2.21 ± 0.12 μm²/ms, and 0.34 ± 0.08 / 0.13 ± 0.04, respectively, consistent with literature [3-5]. MD showed significantly higher, and FA significantly lower, cortical than medullary values. ZOOMit DTI may be valuable for clinical assessment of kidney pathology, particularly for applications closely scrutinizing cortico-medullary differentiation.

Breast imaging
Diffusion-weighted imaging (DWI) of the breast is increasingly used for the differential diagnosis and treatment response monitoring of breast cancer. Conventional EPI techniques are prone to characteristic artifacts, such as susceptibility artifacts, image blurring and spatial distortion resulting from gradient non-linearity and eddy currents. Thus, there can be a significant mismatch in lesion appearance and position between morphologic sequences and EPI which can potentially affect the diagnostic accuracy. In addition, the strong demand
on the gradient system limits the spatial resolution to only moderate in-plane resolution around 1.8 x 1.8 mm² or even coarser. Thus spatially-selective excitation with EPI ZOOMit was applied to a) mitigate the spatial distortion by shortening the phase encode burden and the echo train length and b) to allow the user to restrict the acquisition volume to the breast alone resulting in a higher spatial resolution.

3 healthy subjects were scanned using a 4-channel combined (biopsy and diagnostic) breast coil. For comparison, in both protocols 24 axial slices were acquired with three b-values in 3-scan trace mode with GRAPPA factor of 2. The nominal in-plane resolution was 1.8 x 1.8 mm² vs 1.3 x 1.3 mm² for zoomed EPI where the FOV could be reduced from 332 to 260 mm as well as the
matrix slightly increased (192 x 78 to 200 x 78). Both protocols were set up for comparable acquisition times, 3:53 min (‘standard’) and 3:39 min (ZOOMit).

Figure 4 shows an overlay of a T1-weighted VIBE with a diffusion-weighted (b400) dataset in multi-planar reconstruction to illustrate the excellent co-registration and resolution. In Figure 5 the comparison between standard DW-EPI (top) with the described ZOOMit DWI protocol (bottom) is shown. There is a clear improvement regarding resolution and distortion, in this example the reduced SNR however is obvious as well. This is due to two reasons, first due to increased echo time with a longer RF pulse; but secondly also a higher resolution protocol was acquired. In signal-starved situations, such as fatty breasts higher averaging might still be advised. Clinically, the higher spatial resolution allows the more accurate DWI evaluation of enhancing foci and small non-mass enhancement on MRI. ZOOMit DWI shows potential to mitigate spatial distortions commonly observed in standard EPI. Further developments and investigations are anticipated to balance SNR and spatial resolution, as well as a quantitative comparison including measured ADC and reduction of artifacts.

The preliminary data suggest that using ZOOMit DWI can enable higher spatial resolution in breast applications at reduced distortions when supported by sufficient SNR.

**Medial Temporal lobe imaging**

The hippocampus and entorhinal cortex form a critical neuronal circuit for declarative memory formation that is often altered by different pathologies, including Alzheimer’s disease. Unfortunately, standard single-shot EPI DTI fails to properly characterize the medial temporal lobe structures and their functional connectivity due to geometric distortions from subjacent temporal bone airspaces. In this setting, the 2D-selective RF excitation approach using a Skyra pTX system can be used to overcome these distortions at 3T and, therefore, quantify...
diffusion tensor properties for specific medial temporal lobe structures. To evaluate the role of ZOOMit EPI in this setting, five healthy human volunteers were imaged using the 20-channel head/neck receive coil. Single-shot EPI diffusion-weighted images with fat saturation were obtained (TR 2200 ms, TE 83 ms, NEX 15, acquisition time 8 min, 10 gradient directions, b800 s/mm²) using ZOOMit EPI. The protocol used a 13.3 x 4.4 cm FOV (read x-phase axes) with 1.5 mm in-plane resolution (90 x 30 image matrix). 18 contiguous 3 mm thick oblique coronal slices were obtained orthogonal to the long axis of the temporal lobe with the most posterior slice prescribed tangential to the vertical portion of the hippocampal tail. ZOOMit DW images had acceptable signal-to-noise (6.6 ± 1.1 @ b800 s/mm²) and significantly reduced geometric distortions from subjacent temporal bone airspaces compared to full FOV acquisitions. Diffusion-weighted images resolved specific components of the medial temporal lobes such as the entorhinal cortex (EC), perforant pathway white matter (PP), hippocampal head, subiculum, dentate gyrus (DG), molecular and neuronal layers of the hippocampus (CA1). Quantitative data (Table 2) was consistent with prior DTI parameter values from human hippocampus autopsy samples [5]. Color fiber orientation maps demonstrated coherence from CA1 neuron apical dendrites as previously shown [5] and data may allow tractography of the perforant pathway between the entorhinal cortex and hippocampus. The central portion of the color fiber orientation map (Fig. 6) also demonstrates the midline optic chiasm (OC, red) and paired, anterior coursing optic nerves (ON, green, 4.5 mm diameter). This is a nice illustration of the capability for ZOOMit diffusion to also achieve sufficient resolution to accurately measure diffusion parameters in cranial nerves. Similar results have been observed for the trigeminal nerves and may provide improved diagnosis/monitoring for trigeminal neuralgia.

**Table 2: Quantitative diffusion values using zoomed diffusion.**

<table>
<thead>
<tr>
<th>Region</th>
<th>Mean diffusivity (x 10-3 mm²/s)</th>
<th>Fractional Anisotropy</th>
</tr>
</thead>
<tbody>
<tr>
<td>CA1</td>
<td>0.82 ± 0.04</td>
<td>0.25 ± 0.06</td>
</tr>
<tr>
<td>DG</td>
<td>0.86 ± 0.06</td>
<td>0.27 ± 0.06</td>
</tr>
<tr>
<td>ERC</td>
<td>0.77 ± 0.08</td>
<td>0.32 ± 0.07</td>
</tr>
<tr>
<td>PP</td>
<td>0.81 ± 0.07</td>
<td>0.40 ± 0.08</td>
</tr>
</tbody>
</table>

Similarly to ZOOMit EPI, SPACE with reduced FOV using ZOOMit can increase acquisition efficiency enabling improved spatial resolution or T2-weighting for a given scan duration. For example, ZOOMit SPACE was acquired using the 20-channel head/neck receive coil to visualize the internal auditory canal (IAC) in a patient with Meniere’s disease (axial orientation, TR 1000 ms, TE 125 ms, 2 averages, echo train length of 54, 160 x 80mm FOV, 2 x 2 x 0.5 mm resolution, flip angle 100 degrees, BW: 255 Hz/Px, acquisition time 2:31 min). The acquisition time in this case was approx. 50% shorter then with the non-zoomed reference protocol. As shown in Figure 7, this case shows clear anatomical features including cochlear and vestibular nerves, as well as posterior and lateral semi-circu-

![Image](109)
lar canals in an axial slice (top) and thin maximum intensity projection (bottom). Another application is depicted in Figure 8 using ZOOMit SPACE in the prostate. This example was acquired using a combination of the body-18 and spine coil elements. TR 2000 ms, TE 99 ms, echo train length of 57 and 600 Hz/pixel at 320 x 160 matrix and 72 slices; using a 230 x 115 FOV this resulted in a voxel size of 0.7 x 0.8 x 1 mm at a total acquisition time of 7:15 min providing the capabilities of reformatting the data in any orientation.

Conclusions
The results shown clearly demonstrate that 2D-selective RF excitation on a 2-channel pTX system enables zoomed EPI acquisitions on 3T scanners that can significantly reduce the limitations imposed by spatial distortions due to field inhomogeneity. Together with the inherent ability to simultaneously improve excitation inhomogeneity while providing effective reduced FOV implementations, the TimTX TrueShape clinical platform is ideally suited to fully capitalize on a wide range of improved imaging approaches that require a practical and fully integrated multi-channel transmission platform. These approaches have been previously demonstrated to be of significant value in similarly challenging imaging settings, albeit at the expense of requiring a complex hardware set up due to the lack of commercially available, fully-integrated, multi-channel transmission platforms. The introduction of the clinical Skyra TimTX TrueShape platform is poised to have a significant impact on the clinical implementation of such powerful imaging tools.

For further scientific details of the above shown studies, please refer to Proc ISMRM 2013 abstract numbers (#):

<table>
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<th>Reference</th>
<th>Description</th>
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References
Introduction
Liver disease is a global burden with a growing incidence and prevalence. The World Health Organization recently estimated that there are 800,000 cirrhosis-related deaths per year worldwide [1]. Chronic liver disease has a great impact on public health care costs with therapeutic options ranging from antiviral treatment for viral hepatitis to orthotopic liver transplant for end stage cirrhosis. A variety of pathogens, which can be toxic, viral, metabolic or autoimmune in nature, can induce fibrosis which may progress to cirrhosis if the disease is not detected and treated. An estimated 150 million people worldwide are chronically infected with hepatitis C virus, approximately 350,000 people die due to hepatitis C related liver disease [2]. Liver fibrosis may be reversible at an early stage, which indicates the importance of screening and detection of liver disease. Many forms of liver fibrosis and cirrhosis especially secondary to viral hepatitis increase the risk for the development of liver cancer, namely hepatocellular carcinoma. Non-alcoholic steatohepatitis is emerging as a major pathway into chronic liver disease and is closely related to other metabolic disease entities such as diabetes and morbid obesity. The incidence and prevalence of these diseases has risen steadily over recent years. In a clinical context, liver disease is often reflected by a combination of several contributing factors, fibrosis, hepatic steatosis and iron overload, each with different forms of manifestations. Although these diseases are considered ‘diffuse’, actual hepatic parenchymal involvement by any of these can be irregular and patchy, leaving other parenchymal areas unaffected. Clinical management of patients with diffuse liver disease requires tools to accurately detect and classify the various forms of liver disease. Even with decades of experience in imaging, liver biopsy and the histological workup of the specimens have traditionally been the reference standard in the characterization of liver disease [3]. However, biopsy is prone to sampling errors if less affected parenchyma is sampled and may not reflect the true disease severity and distribution in a particular organ due to the variance in the heterogeneous pattern of histological changes on a macroscopic scale [4, 5]. Biopsy, associated with the risks of an invasive procedure, is employed for disease detection and staging, but periodically repeated biopsy is not a practical
method for disease monitoring. Additionally, given its attendant risks and costs, biopsy is only performed in patients in whom liver disease is strongly suspected, and is not suitable for evaluating those patients with only mild or questionable symptoms. Thus, there has been great interest in non-invasive methods to assess diffuse liver disease, and imaging modalities, in particular magnetic resonance imaging (MRI), have evolved as potential tools to measure certain biomarkers.

Liver MRI offers a variety of methods to detect and quantify parenchymal changes which occur in chronic liver disease [6, 7]. In contrast to liver biopsy, liver MRI allows for assessment and evaluation of the entire liver volume by means of quantitative measurements and color coded maps which reflect the geographical disease distribution. Quantification of excessive fat and iron deposition was shown to be accurate as demonstrated by many studies in the literature [8, 9]. The characterization of focal liver lesions and determination of treatment options for hepatocellular carcinoma is a well-established clinical application of liver MRI, and this use will grow as the prevalence of chronic liver disease increases [10]. The detection and accurate classification of liver fibrosis and cirrhosis remains challenging, despite the usefulness of liver MRI in the aforementioned scenarios [7].

While liver MRI may soon provide a comprehensive evaluation of liver disease it is a complex technique requiring highly trained personnel, cooperative patients and optimized scanning conditions to produce diagnostic images acceptable for clinical interpretation. MRI must compete with other imaging modalities, sonography and computed tomography (CT) in categories such as availability, cost, acquisition time, robustness, reproductibility, patient acceptance, and comfort. In particular, it has been advocated that MRI needs to close the gap in acquisition time compared to CT; simultaneously methods should be employed to standardize the image acquisition workflow, improve reproducibility of a quantitative imaging protocol, and reduce the time spent performing redundant data acquisition or preparatory steps [11]. However, in terms of diagnostic performance, lack of ionizing radiation compared with CT, and the variety of tissue contrasts available, MRI has several clear advantages. In the following article we will discuss new methods which address MRI related issues like artifacts and breathing motion while improving image quality and spatial resolution, offer fast automated screening techniques for the detection of parenchymal changes, and optimize the imaging workflow to decrease overall acquisition time.

Liver imaging

A modern liver imaging protocol must accomplish at least two main goals. The presence of diffuse liver disease, fat, iron deposition and possibly fibrosis/cirrhosis should be detected and ideally quantified, even if not expected at the time that the examination is initiated. Additionally, focal hepatic lesions, in particular in the setting of cirrhosis, must be characterized to allow for classification into benign entities, such as simple cysts, hemangiomas, focal nodular hyperplasias (FNH) or adenomas versus malignant hepatic tumors like hepatocellular carcinoma, cholangiocarcinoma or metastatic disease. These tasks must be accomplished within a reasonable amount of time without compromise in image quality or obtainable information. A number of methodologies have become recently available which provide automated diagnosis of diffuse liver disease, higher spatial resolution imaging, and automated workflows.
Abdominal Imaging Clinical

Screening Dixon

Fat and water separation can be realized by 3D In- and Opposed-Phase T1-weighted data acquisition with two-point Dixon reconstruction. This 3D gradient-echo imaging sequence produces four sets of images with In-/Opposed-Phase, Water-Only and Fat-Only depiction. The Water-Only image set can further be employed as a standard pre-contrast fat-suppressed sequence, one of the standard in a liver MRI protocol with contrast material application [12]. The two-point Dixon method offers visual, qualitative assessment of hepatic steatosis but the acquired data allow for a semi-quantitative estimation of fat deposition as well as iron overload [13, 14]. This pulse sequence provides two essential image sets (In- and Opposed-Phase and pre-contrast T1) in a single breath-hold.

The Screening Dixon method represents a two-point Dixon technique with an additional liver sampling algorithm that automatically segments large portions of the liver. Within a large volume of the segmented liver, dual signal intensity ratios from In-phase/Opposed-phase and Fat-Only/Water-Only data sets are calculated by the algorithm to produce an assessment regarding the presence of diffuse liver disease. The result of the algorithm (normal, fat, iron, or combined disease) can be coupled with a recommendation to perform a dedicated quantitative sequence for the detected abnormal metabolite (e.g. iron quantification sequence) [13, 15].

Hence, the Screening Dixon technique offers an automated approach to diffuse liver disease but simultaneously allows for a quantitative MRI protocol tailored to the individual patient. Instead of performing a time consuming all-in-one MR imaging protocol that comprises any type of quantification, the Screening Dixon methods stratifies protocol steps in a way that only essential quantitative sequences are acquired, potentially reducing overall acquisition time compared with an exam which performs quantification unnecessarily (Figs. 1, 2). Notably, although this method can provide estimates of proton density fat fraction (PDFF) and R2* (a surrogate for iron concentration), these are not corrected for a variety of potentially confounding factors, and should be confirmed by a dedicated quantification sequence when abnormal. Even so, this algorithm performs well for the task of detecting diffuse steatosis/siderosis [16].

Quantification results from the same volunteer as in Fig. 2. Top: HISTO (3A, B) voxel positioning, (3C) quantification results in a graphical norm range display. Bottom: Multi-echo Dixon (3D) fat percentage map, (3E) R2* map.
Fat and iron quantification

R2* and PDFF can be quantified in two different ways. Multi-echo, T2 corrected, single breath-hold spectroscopy (HISTO) gives highly reproducible values from a single voxel [17]. Multi-echo 3D gradient echo (VIBE) imaging, with Dixon reconstruction and correction for T2* as well as the multi-spectral nature of fat, allows quantification with good spatial resolution [18]. Figure 3 shows quantification results from the same volunteer as in Fig. 2. The results are consistent and support the initial Screening Dixon estimation.

CAIPIRHINA

There are two distinct time points within a contrast enhanced liver MRI protocol which are crucial in the acquisition of diagnostic images. Hepatic arterial phase imaging is the mainstay in the detection of hepatocellular carcinoma [18]. Here, accurate timing is important [22]. The images may be acquired as a multi-arterial phase to improve temporal resolution [17]. A ‘late phase’, whether the vascular equilibrium phase obtained with extracellular contrast agents or the hepatocyte phase obtained with hepatobiliary agents, can differentiate lesions based on their contrast retention behavior. For both elements within a liver MRI protocol, a compromise must be made between achievable image resolution and acquisition time. This conflict is even more challenging for single breath-hold/multiple arterial phase imaging. The duration of a breath-hold remains the limiting factor in contrast enhanced liver imaging, and a sequence must balance the two factors; sufficient spatial resolution within a reasonable acquisition time. The evolution of parallel imaging techniques has allowed multiple acquisitions within the arterial phase time window to reliably capture the late hepatic arterial phase, a critical image for hepatocellular carcinoma imaging. However, 20-25 second breath-holds remain challenging for some patients. Hepatobiliary phase imaging is ideally done using high spatial resolution to utilize the diagnostic information derived from hepatocyte specific contrast agent for all liver abnormalities, in particular for smaller otherwise indeterminate lesions (> 1 cm lesion diameter). Although the time window for the hepatobiliary phase is wider compared to arterial phase imaging, the achievable spatial resolution is similarly limited by the duration of a breath-hold which may be even shorter at the end of an examination due to developing fatigue of the patient. Patients with impaired or poor breath-holding capabilities may render any breath-hold sequence acquisition non-diagnostic if the acquisition time exceeds their capabilities. Further increasing standard parallel imaging acceleration, however, decreases the signal-to-noise ratio (SNR) and induces image artifacts.
CAIPIRINHA (Controlled Aliasing in Parallel Imaging Results in Higher Acceleration), a new parallel imaging technique differs in the k-space sampling pattern compared to standard acceleration techniques and is more efficient in using the coil sensitivities [21]. Undersampling is performed in both the phase and partition directions, allowing for a higher acquisition matrix and improved image resolution. This provides dramatic improvements in spatial resolution for the same breath-hold times, and can be optimized to provide a combination of high spatial resolution and reduced breath-hold duration (Figs. 4, 5).

**Workflow – Abdomen Dot Engine**

MRI data acquisition is time consuming, and the considerably longer examination times (compared to CT) must be justified by added benefits to patient care. Total examination time is composed of time spent acquiring image data and time spent performing a variety of setup tasks, including patient positioning and pulse sequence preparation. In liver MRI, preparatory tasks have been shown to consume more time than image acquisition, as such there is substantial opportunity for improvement in addressing the efficiency of performing these tasks [19]. This leads to operator dependent workflow, inconsistent image quality, and prolonged examination times. Additional patient specific factors further influence the achievable image quality and overall examination time, for example patients vary in their breath-holding capabilities and may fatigue throughout an examination. Adjustment of the imaging strategy, breath-hold versus free breathing, triggered imaging versus a shortened, fast imaging protocol may be necessary to accommodate individual differences. Additionally, many of the pulse sequence preparatory tasks are redundant and therefore offer opportunities for automation or a guided standardized setup. The current MRI acquisition workflow can be rendered more efficient thus reducing overall room time.

The Abdomen Dot Engine is an approach that incorporates various strategies to optimize and standardize a complex abdominal MRI protocol. It allows for automatic detection and positioning of an individualized field-of-view (FOV) based on localizer images, can stratify a protocol into patient specific breathing capabilities and provides tools such as automatic bolus timing for dynamic scanning [20]. A liver MRI protocol, for example can be standardized and partitioned into its typical elements (pre-contrast, multiple arterial, portal venous, and equilibrium phases), so that key settings such as delay times between each element can be configured by the user through an interface that offers an overview of all protocol steps (Fig. 6). A standardized and guided workflow for MRI examinations is needed to release the operator from redundant tasks, such as defining patient-specific sequence parameters, and allowing him or her to focus on global protocol strategies. Furthermore, the consistency of image quality across studies may be improved, and the risk of rescanning a sequence or the entire protocol may be reduced, increasing the robustness of the modal-
Additionally, multiple scan types which differ by only a few minor components (e.g., with or without MR Cholangiopancreatography (MRCP), with or without diffusion-weighted imaging) can be combined into a single, efficient protocol with a few key decision points, reducing redundancy and allowing for simpler base protocol maintenance and modification when necessary.

**Summary**

MRI examinations face serious competition compared to sonography and CT when categories such as robustness, acquisition time, patient comfort and health care costs are considered. An abundance of information may be acquired through high resolution imaging and dedicated quantitative MRI sequences, but images and measurements should be reproducible and reliable in their diagnostic value. The redundancy of preparatory steps for the operator within an MRI protocol is an opportunity for more efficient and less time consuming imaging. In addition, the image acquisition process can be improved by means of faster imaging at higher resolution with the implementation of new parallel imaging acceleration techniques, to reduce the risk of motion in patients with limited breath-hold capabilities. Intelligent imaging protocols, which self-optimize during the course of the examination or use initial pulse sequences to tailor subsequent sequence selection, can provide faster and more efficient examinations, which include quantitative data when appropriate. Combining all of the described improvements may equip liver MRI examinations with sufficient tools to remain unique in delivering disease specific quantitative data while expanding their diagnostic value.
References


Contact
Tobias Heye
University Hospital Basel
Department of Radiology
Petersgraben 4
CH-4031 Basel
Switzerland
Phone: +41 (0)61 328 6324
tobias.heyepi@usb.ch
Faster Abdominal MRI Examinations by Limiting Table Movement

Mustafa Rifaat Bashir¹; Brian Marshall Dale²; Wilhelm Horger³; Daniel Tobias Boll¹; Elmar Max Merkle¹

¹Radiology, Duke University Medical Center, Durham, NC, USA
²Siemens Healthcare, Cary, NC, USA
³Siemens Healthcare, Erlangen, Germany

1A How to create a minimized shimming protocol. Modifications will be made to pulse sequences following the initial localizers (1A). For the second sequence in the scan protocol, ‘Shim mode’ is set to ‘Standard’, and ‘Adjust with body coil’ is selected (1B). For the third and subsequent sequences, ‘Positioning mode’ is set to ‘FIX’ (1C). ‘Shim mode’ is set to ‘Standard’, ‘Adjust with body coil’ is selected, and ‘Adjustment Tolerance’ is set to ‘Maximum’ (1D). Finally, for the third sequence, a reference is created to the table position of the second sequence (1E). The same reference is created for all subsequent sequences (1F).
Hepatic magnetic resonance imaging (MRI) is widely used for a variety of indications, including characterization of focal lesions, detection of diffuse liver disease, as well as evaluation for hepatocellular carcinoma in patients with chronic hepatitis or cirrhosis [1-8]. One of the main criticisms of hepatic MRI is scan time, both in terms of length and variability. In a small prospective study at a single center, table times in contrast-enhanced MRI of the liver were shown to vary from 19 up to 58 minutes, even when examinations were performed by an experienced technologist [9].

In a detailed analysis of MRI scanner activity during various imaging examinations, we observed that the number of adjustments performed by the scanner prior to initiating a pulse sequence was higher when imaging the liver as compared to the knee [10]. These adjustments take time, with the acquisition of shim data alone taking up to 20 s per pulse sequence. The need to acquire new adjustment data is dictated, in part, by table movement, after which field homogeneity in the scan volume must be measured, and additional adjustments made as needed. This increases the time that the MRI system spends preparing to scan but not actually acquiring image data [10].

A more recent study showed that a scan protocol in which the table moves only once during the examination achieves significant total scan time savings by obviating the need to gather new adjustment data during the course of the examination [11]. For every step in which the table moves and new shim data is acquired, this protocol change reduces the time spent collecting prescan adjustment data by approximately 30 s. In that study, a reduction in total examination time of up to 20% was observed in a non-contrast liver MRI/MRCP protocol, with no observable change in image quality.
Methods
Automated algorithms to minimize table movement have already been incorporated into MAGNETOM MRI systems under syngo MR D11 and later software versions. Under earlier software versions, a few simple steps can be performed to convert a standard MRI protocol into a minimized shimming protocol, in order to realize the time savings previously described. These changes can all be made via the Exam Explorer (Fig. 1A).

Pulse Sequence #1 – localizer
The first pulse sequence of an examination is a localizer, typically utilizing either a three-plane TrueFISP or HASTE technique. At the MRI console, under the ‘Sequence’ card, the Shim is set to ‘None’ (typically the default value), and precalibrated prescan data is used with no need to acquire new prescan data. No additional modification of this sequence is required.

Pulse Sequence #2 – first and only table move
Using the image data from the localizer sequence, the image volume for Pulse Sequence #2 is prescribed. This volume should be centered on the area of interest and rather large, covering the volume of interest for the entire examination; at our institution, we typically use a coronal HASTE sequence for this purpose. The prescan data acquired in this step, including shim data, will be carried forward for the remainder of the examination. The following modifications are made to this pulse sequence:
1. In the ‘System’ card, under the ‘Adjustments’ tab, set ‘Shim mode’ to ‘Standard’. Check the ‘Adjust with body coil’ box (Fig. 1B).

Pulse Sequences #3 and higher – no further table movements
For all subsequent pulse sequences, table movement is disallowed, and prescan adjustment data from Pulse Sequence #2 is carried forward, so that as little time as possible is spent acquiring new adjustment data. The following modifications are made:
1. In the ‘System’ card, under the ‘Miscellaneous’ tab, set ‘Position mode’ to ‘FIX’ (Fig. 1C).
2. In the ‘System’ card, under the ‘Adjustments’ tab: set ‘Shim mode’ to ‘Standard’; select ‘Adjust with body coil’; and set ‘Adjustment Tolerance’ to ‘Maximum’ (Fig. 1D).
3. From the Exam Explorer, right-click on the sequence and select ‘Properties’.

Under the ‘Copy References’ tab, check the ‘Copy reference is active’ box, then select Pulse Sequence #2 in the left-hand window and ‘Table position’ in the right-hand window (Fig. 1E). In combination with step #2 above, this ensures that the MRI system table will not move when progressing to later pulse sequences in the examination, despite different prescriptions for the imaging volume.
4. Repeat steps 1–4 for all subsequent sequences (Figure 1F).

Discussion
Preparatory adjustments made by an MRI system are essential to realize excellent image quality. In particular, adequate shimming is necessary to ensure magnetic field homogeneity. Shimming is a process whereby the main magnetic field ($B_0$) is fine-tuned to compensate for field fluctuations and inhomogeneities introduced by the presence of the human body within the scanner. These adjustments are applied specifically to a volume within the bore of the magnet (based on the anticipated imaging volume), attempting to optimize magnetic field homogeneity within that volume while sacrificing field homogeneity outside of the volume.
A scanning protocol which prohibits table movement can reduce total table time by removing the need to repeatedly acquire adjustment data, particularly time-consuming shim data, throughout the course of the examination. Even though placing the liver at isocenter during an inspiratory breath-hold means that it would be located, on average, cranial to isocenter during free breathing sequences, the above study observed no differences in image quality in any type of pulse sequence. Note we do not suggest that patient- and position-specific shimming is unnecessary, but rather that if patient position can be maintained during the examination and adequate initial adjustment data is acquired, table movement during the examination and much of the additional adjustment data may be unnecessary. This holds true for both typical quantitative and qualitative abdominal MRI applications (Fig. 2). In addition, it may be possible to achieve the same time savings by any protocol which prevents table movement, for example a protocol where the image volumes of all pulse sequences are centered in the same location.

An overall workflow analysis of abdominal MRI acquisitions shows that shortening data acquisition times can reduce overall imaging time [10]. This is likely to be of greatest benefit in examinations where the large majority of table time is spent acquiring image data, e.g. musculoskeletal and brain examinations [10]. However, much less attention has been directed to other events which contribute to total imaging time, including time spent preparing the patient for imaging as well as scanner preparation, such as image prescription and prescan adjustments. In abdominal MRI, where preparatory activities take up a large proportion of total table time, reducing the prescan time represents an important opportunity to reduce total table time substantially [10]. Since this methodology is independent of the particular details of the imaging protocol, it could be applied to a variety of routine clinical and novel imaging techniques. In conclusion, any MRI protocol can be easily modified to minimize the time spent collecting prescan adjustment data. In certain scenarios, such modifications can reduce total scan time by as much as 20% with no sacrifice in image quality.

References

Contact
Mustafa RIafaat Bashir, M.D.
Duke University Medical Center 3808
Durham, NC 27710
USA
Phone: +1 919 684 7663
mustafa.bashir@duke.edu
How-I-do-it

Bandwidth in MRI?

Joachim Graessner, Dipl.-Ing.
Siemens Healthcare, Hamburg, Germany

Questions about the term ‘bandwidth’ are most frequently asked during MR trainings and discussions regarding MR parameters and image quality. This article is intended to clarify the meaning of bandwidth in MRI, to show dependencies with other MR parameters, and to give hints for daily routine work and protocol optimization. The bandwidth occurs twice in the course of an MR sequence (Fig.1): during transmission describing properties of the radiofrequency (RF) pulses and during reception of the MR signal.

Transmission phase
During transmission you have to apply RF pulses which contain a certain range of frequencies, i.e., bandwidth, to excite a discrete slice thickness or volume while a local gradient is on as well. The bandwidth typically lies in the range of a

![Sequence Timing Diagram](image)

1 Sequence timing for se15_b130.
How-I-do-it

few kilohertz (kHz). The syngo user interface offers three types of pulses for most sequences in the Sequence parameter card, part 2: Low SAR, Normal, and Fast (Fig.2).

Pros and Cons for RF pulse types with timing examples:

**Low SAR (3.84 ms)**
- Longer RF pulse with good slice profile
- Reduced SAR values (lower amplitude)
- Less crosstalk between slice; narrower gaps tolerated
- Longer minimum TEs and TRs

**Normal (2.56 ms)**
- Normal RF pulse with good slice profile
- Optimized SAR behavior

**Fast (1.28 ms)**
- Shorter RF pulse, with a compromised slice profile
- Higher SAR compared to the other modes (higher amplitude)
- Shorter echo spacing (ES)
- Opportunity for shorter TEs and TRs
- Fewer susceptibility artifacts

The 'Fast' mode is mainly applied in sequences where measurement speed and short echo spacing (ES) – the time from one echo to the next echo – is essential for good image quality: HASTE, EPI, TrueFISP, and TurboFlash. These modes can be combined with up to three gradient modes: Whisper, Normal and Fast. The gradient modes influence the slew rate of the gradient pulses, or the steepness of the ramps from zero to full gradient strength and back. The trade-off for faster sequence timing with shorter TEs and TRs usually results in more noise and a higher potential for peripheral stimulation.

**Reception phase**

When the echo signal appears, a read-out gradient is switched on and the analog-digital-converter (ADC) samples this signal. The gradient encodes different rows of frequencies into signal; the time-amplitude integral defines the measured field-of-view (FOV). The ADC is virtually asking the echo-signal (e.g., every 30 microsecond (µs)) how high its amplitude is and writes these values in a digital fashion into the memory of the image computer. This time is called the dwell time D. For a 256 base matrix this process has to be repeated 256 times. The whole sample period Tₛ will be in this example (Fig. 3):

\[ Tₛ = 256 \times 30 \mu s = 7.680 \text{ ms} \]

\[ Tₛ = N \times D \text{ with } N = \text{matrix in read} \]

The frequency content of this sampled signal is the reciprocal of the dwell time:

\[ f = 1/D \]

In the above example it is roughly 33 kHz.

The magic term bandwidth (BW) is the reciprocal of the total sampling time Tₛ and has the unit of Hz/pixel:

\[ \text{Bandwidth: } BW = 1/Tₛ, \text{ [Hz/pixel]} \]

The parameter 'Bandwidth' is found on the Sequence parameter card, part 1 (Fig. 4). The bandwidth describes in a simplified manner which frequency range from our analyzed echo signal is transferred into one pixel:

1. lower left pixel: 0-130 Hz
2. lower left pixel 130-260 Hz
3. and so forth. If you continue this fill-up task to the last pixel of a 256 matrix you end up with 33 kHz in the lower right pixel.
How I do it

Increasing the bandwidth reduces the sampling time $T_s$ and shortens the sequence timing, allowing shorter TE and TR values and vice versa (Fig. 5). The disadvantage of a higher bandwidth is the larger amount of noise which is sampled due to larger frequency range.

![Image of bandwidth bands]

On the positive side we have the shorter echo spacing (ES) which delivers less blurring in the images. The lipid signal in turbo spin echo protocols is also influenced by the echo spacing and the turbo factor (TF) [1]; the shorter the ES and the higher the TF, the brighter the fat in TSE protocols. The larger readout (RE) gradient amplitude needed for a high BW protocol increases the minimally adjustable FOV (Fig. 6). Furthermore, high bandwidth protocols play an important role in metal artifact reduced scanning. Shorter optimized excitation pulses combined with short readout intervals and short ES enables fewer distortions due to conditional implants. This feature is enhanced with optimized protocols which use WARP and VAT (view angle tilting) [2–4].

Last but not least, the chemical shift between water and fat gets smaller with high BW and larger with lower BW. This feature is of less importance for fatsat protocols. Fat resonates at an approximately 3.3 ppm lower frequency than water. Although commonly excited in the first part of a sequence, fat and water from the same physical place answer with different frequencies. In the above mentioned sorting process of frequency ranges into pixels we have to face a missorting. Fat will always appear shifted to lower frequencies by a certain amount of pixel in the readout direction. The only exception is the EPI sequence where we have a very strong chemical shift artifact in the phase encoding direction (PH) due to the long sampling period producing echoes. This very long sampling period in the order of $>40$ ms or more translates into a quite low bandwidth ($<25$ Hz/pixel). Fatsat is therefore mandatory for EPI measurements.

The chemical shift in readout direction is indicated as a mouse-over tooltip on the bandwidth field in the Sequence parameter card, part 1 (Fig. 7).

As one can see, chemical shift is not a big issue at low field strengths (Table 1). It is getting important at 1.5T and above for the range of practical bandwidth of a few hundreds Hz/pixel. But with the rise of the magnetic field the resolution typically goes up as well, keeping the abso-
ute chemical shift in millimeters reasonably low. A protocol at 3T with a bandwidth of 195 Hz/pixel and an inplane resolution of 0.3 mm creates an absolute chemical shift of less than 0.7 mm. Figure 8 demonstrates in an exaggerated way the artificial movement of the intervertebral discs (orange) towards the upper vertebrae or the spinal cord depending on the chosen readout direction and bandwidth option. The low bandwidth examples should be avoided in daily routine, although one may gain SNR and shorter scan time with lower bandwidth protocols.

Table 2 shows the effects of optimizing a TSE protocol with respect to change of bandwidth and resolution for a 1.5T scanner. Identical SNR is the border condition for these four examples. TE was kept in the mid thirty ms range. The penalty for better resolution is usually longer measurement time. Personal taste and application dictates the parameters chosen. Starting with protocol B, one can speed up the scan by halving the bandwidth to

### Table 1: Absolute pixel shift for different field strengths.

<table>
<thead>
<tr>
<th>Bandwidth* [Hz/pixel]</th>
<th>Readout time [ms]</th>
<th>Chemical Shift* [pixel] 0.35T</th>
<th>Chemical Shift* [pixel] 1T</th>
<th>Chemical Shift* [pixel] 1.5T</th>
<th>Chemical Shift* [pixel] 3T</th>
<th>Chemical Shift* [pixel] 7T</th>
<th>Relative SNR change* [%]</th>
</tr>
</thead>
<tbody>
<tr>
<td>3000 0.330</td>
<td>0.017</td>
<td>0.05</td>
<td>0.075</td>
<td>0.15</td>
<td>0.3</td>
<td>-79</td>
<td></td>
</tr>
<tr>
<td>2000 0.500</td>
<td>0.025</td>
<td>0.07</td>
<td>0.1</td>
<td>0.2</td>
<td>0.5</td>
<td>-74</td>
<td></td>
</tr>
<tr>
<td>1000 1.000</td>
<td>0.05</td>
<td>0.1</td>
<td>0.2</td>
<td>0.4</td>
<td>1.0</td>
<td>-64</td>
<td></td>
</tr>
<tr>
<td>520 1.920</td>
<td>0.09</td>
<td>0.3</td>
<td>0.4</td>
<td>0.8</td>
<td>1.9</td>
<td>-50</td>
<td></td>
</tr>
<tr>
<td>480 2.080</td>
<td>0.1</td>
<td>0.3</td>
<td>0.4</td>
<td>0.9</td>
<td>2.1</td>
<td>-48</td>
<td></td>
</tr>
<tr>
<td>390 2.560</td>
<td>0.13</td>
<td>0.35</td>
<td>0.5</td>
<td>1.1</td>
<td>2.5</td>
<td>-42</td>
<td></td>
</tr>
<tr>
<td>260 3.840</td>
<td>0.2</td>
<td>0.5</td>
<td>0.8</td>
<td>1.6</td>
<td>3.8</td>
<td>-29</td>
<td></td>
</tr>
<tr>
<td>195 5.120</td>
<td>0.25</td>
<td>0.7</td>
<td>1.1</td>
<td>2.2</td>
<td>5.1</td>
<td>-18</td>
<td></td>
</tr>
<tr>
<td><strong>130 7.680</strong></td>
<td><strong>0.4</strong></td>
<td><strong>1.1</strong></td>
<td><strong>1.6</strong></td>
<td><strong>3.2</strong></td>
<td><strong>7.6</strong></td>
<td><strong>0</strong></td>
<td></td>
</tr>
<tr>
<td>100 1.000</td>
<td>0.5</td>
<td>1.4</td>
<td>2.1</td>
<td>4.2</td>
<td>9.9</td>
<td>14</td>
<td></td>
</tr>
<tr>
<td>78 12.280</td>
<td>0.6</td>
<td>1.8</td>
<td>2.7</td>
<td>5.4</td>
<td>12.7</td>
<td>29</td>
<td></td>
</tr>
<tr>
<td>65 15.380</td>
<td>0.8</td>
<td>2.1</td>
<td>3.2</td>
<td>6.4</td>
<td>15.2</td>
<td>41</td>
<td></td>
</tr>
<tr>
<td>50 20.000</td>
<td>1.0</td>
<td>2.8</td>
<td>4.2</td>
<td>8.4</td>
<td>19.8</td>
<td>61</td>
<td></td>
</tr>
</tbody>
</table>

*Rounded values for convenience.
Chemical shift (CS) on an example sagittal spine.

How I do it:

compensate for the SNR loss coming from the reduction of acquisitions. An increase of the inplane resolution in protocol C and D results in an obvious prolongation of the scan time which could be kept tolerable when reducing the bandwidth. In example D partial Fourier, i.e., reducing the measured turbofactor, allows a TR minimization for an acceptable scan time. One could save further time in this example by a decrease of phase oversampling, which was chosen 100% for a phase encoding in head-feet direction to minimize flow artifacts from the arteries; a slight increase of the FOV, e.g., to 200 mm, could compensate for this SNR reducing step though sacrificing some hundreds of a mm inplane resolution.

On a 3T system one should at least increase the bandwidth by 50% and save time by adjusting the other parameters. The higher signal of 3T supports this measure.
Table 2: Protocol optimization.

<table>
<thead>
<tr>
<th>#</th>
<th>Parameter</th>
<th>Bandwidth</th>
<th>Matrix</th>
<th>Resolution</th>
<th>Scantime</th>
<th>TR</th>
<th>TE</th>
<th>Echospacing</th>
<th>Avagages</th>
<th>partial Fourier</th>
<th>rel. SNR</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>low BW normal Res</td>
<td>75</td>
<td>256 x 512</td>
<td>0.7 x 0.35 = 0.245</td>
<td>2:46</td>
<td>4020</td>
<td>36</td>
<td>17.8</td>
<td>1</td>
<td>off</td>
<td>1</td>
</tr>
<tr>
<td>B</td>
<td>high BW normal Res</td>
<td>150</td>
<td>256 x 512</td>
<td>0.7 x 0.35 = 0.245</td>
<td>3:32</td>
<td>2770</td>
<td>34</td>
<td>11.2</td>
<td>2</td>
<td>off</td>
<td>1</td>
</tr>
<tr>
<td>C</td>
<td>high BW higher Res</td>
<td>150</td>
<td>336 x 448</td>
<td>0.54 x 0.4 = 0.216</td>
<td>4:43</td>
<td>2870</td>
<td>34</td>
<td>11.7</td>
<td>2</td>
<td>off</td>
<td>1</td>
</tr>
<tr>
<td>D</td>
<td>low BW higher Res</td>
<td>75</td>
<td>384 x 512</td>
<td>0.47 x 0.35 = 0.165</td>
<td>6:04</td>
<td>3240</td>
<td>37</td>
<td>18.7</td>
<td>2</td>
<td>on</td>
<td>1</td>
</tr>
</tbody>
</table>

* In Accordance to the German MR Guidelines (BÄ)

Further reading

1. Magnets, Spins and Resonances; Siemens 2003.
4. MRI the Basics: Hashemi, Bradley, Lisanti; LWW 2010.
6. MRI from Picture to Proton: McRobbie, Moore, Graves, Prince; Cambridge 2007. Especially recommended for the German speaking community:
7. Praxiskurs MRT: Nitz, Runge; Thieme 2011.

Helpful links


References

Indications for 24 Hours/7 Days Emergency MRI

Marc-André Weber, M.D., M.Sc.; Jürgen Biederer, M.D.
Heidelberg University Hospital, Diagnostic and Interventional Radiology, Heidelberg, Germany

Introduction
For many years MR imaging (MRI) has been considered a second-line procedure required for further diagnostic work-up after first-line imaging with x-ray, ultrasound or even computed tomography (CT) in the emergency room. However, the increasing performance of modern MR equipment and sequence design have broadened the range of indications, now making MRI the first-line imaging modality of choice for a number of clinical conditions. This is most obvious in neurovascular emergencies, but it also applies to a number of other indications. More and more, an ‘emergency MRI’ is being requested at night or during weekends. In most cases, the decision whether to perform it is taken according to the particular circumstances, such as the availability of sufficiently skilled staff and radiological expertise. The aim of this article is to suggest stratification criteria and to provide a list of clinical situations that might justify the performance of an MRI scan during night or weekend shifts based on the clinical relevance, i.e. immediate consequences. Conditions that do not require direct therapeutic intervention should not trigger an MRI scan outside the regular schedules. The limitation to only a small number of indications improves clinical decision-making and facilitates the preparation and training of the staff for these situations. The following suggestions have been developed at University Hospital Heidelberg in cooperation between the Department of Diagnostic and Interventional Radiology (Head: Hans-Ulrich Kauczor, M.D.), the Orthopedics and Trauma Surgery Clinic, the Spinal Cord Injury Center, the Vascular Surgery Clinic, the Department of Anesthesiology, and the Center

1 Intracranial hemorrhage in the right basal ganglia with small perifocal edema (arrows). 74-year-old man presenting with left sided hemiparesis since waking up 6 hours before. (1A) Axial unenhanced T1w, (1B) axial FLAIR, (1C) axial T2*w, (1D) axial diffusion-weighted image (b-value of 1000 s/mm²).
How-I-do-it
for Pediatric and Adolescent Medicine.
Of course, the following suggestions are subject to ongoing discussion and refinement. The Department of Diagnostic and Interventional Radiology is the central service provider at the University Hospital Heidelberg in the field of diagnostic general radiological imaging and interventions. More than 91,000 examinations in out-patients and more than 71,000 in in-patients are performed annually, covering all indications and organ systems, with more than 210,000 imaging procedures every year. It should be noted that the following suggestions have been developed for a general radiological department. Dedicated neuroradiological departments may therefore develop additional suggestions regarding brain imaging.

List of indications for emergency MRI at the Department of Diagnostic and Interventional Radiology in Heidelberg

The list of indications differentiates between emergencies requiring immediate MRI (Category A, urgent care required as soon as possible day and night) and urgent cases with high priority but no need for immediate intervention (Category B, to be performed within 12 hours, e.g. next day). It was also considered important to define a third category (Category C) for situations that do not require an immediate MRI scan since equally diagnostic alternatives are available. Although such examinations may sometimes be urgently requested, it is strongly recommended to resist and to preserve the resources of the emergency MRI staff. This list represents the current stage of management and is intended to be regularly updated.

Category A
Indications for an immediate emergency MRI

1. Cerebral and neurovascular emergencies (Fig. 1; e.g. acute cerebral ischemia or herniation syndromes in children): minimal protocol: T2-weighted TSE, dark-fluid imaging, diffusion-weighted imaging (DWI), time-of-flight (TOF) angiography, NO routine intra-venous (i.v.)-contrast medium administration.

2. Acute traumatic and non-traumatic syndromes with paraplegia and apparent neurologic deficits (such as paresis, sensory disturbances, disturbances in bladder or rectum function) that raise suspicion of a lesion of the myelon or the cauda equina. Examples include: Clinically suspected spondylodiscitis with epidural and psoas muscle abscesses. 80-year-old woman with severe back pain. The spondylodiscitis in the first and second lumbar vertebra is clearly acknowledged on the sagittal contrast-enhanced fat-suppressed T1-weighted images (arrows in 2C). T1w TSE sagittal (2A) and T2w TSE sagittal images (2B). Also the epidural enhancement within the spinal canal (open arrows in 2C and 2D) and left psoas muscle abscess (asterisk in 2D) can be evidenced best on the contrast-enhanced fat-suppressed image.
Category A

Indications that require an emergency MRI within 1 hour include:

1. Spinalis anterior syndrome (Fig. 3); suspicion of epidural hematoma following spinal anesthesia or spinal surgery; suspected spinal cord contusion; clinical relevance: surgical decompression if edema of the spinal cord is detected. 

Minimal protocol: T2w TSE fat-saturated sagittal, T1w SE sagittal, T2w TSE transversal (non-fat-saturated) findings-centered. Optional: Diffusion-weighted imaging in case of suspected spinal ischemia. In case of suspected epidural abscess MRI with i.v.-contrast medium required.

3. Strong clinical suspicion of septic arthritis (Fig. 4; clinical relevance: early joint lavage to prevent chondrolysis indicated). MRI with i.v.-contrast medium required.

4. Strong clinical suspicion of osteomyelitis in children. MRI with i.v.-contrast medium required.

5. Acute pulmonary artery embolism in pregnant women or very young patients (Fig. 5; pulmonary artery embolism protocol based on free breathing TrueFISP images. i.v.-contrast-enhanced TWIST per-

fusion and high spatial resolution MR angiography (MRA) only to be used, if exclusion of small peripheral emboli would be clinically relevant).

Category B

Indications for an MRI within 12 hours include:

1. Spinal emergencies without neurological symptoms, e.g. to exclude spondylodiscitis or a ligamentous affection following a trauma of the spine, suspicion of a discoligamentous injury according to CT findings (use standard spine MR protocols).

2. In conventional radiography inconclusive findings or suspicion of occult fractures to prevent exposure to radiation in CT (especially in childhood).

Category C

Indications that do NOT justify an emergency MRI (=> e.g. CT as alternative emergency modality or MRI the next working day):

1. Run-off MRA for arteriosclerosis or acute occlusion of the lower limb (CT angiography as an alternative).

2. Suspicion or follow-up of intracranial hemorrhage (CT as an alternative) unless classified as neurovascular emergency according to Category A 1.

3. Suspicion of cerebral metastasis (CT with contrast medium as an alternative).

4. Urgent MRI requests due to organizational issues of the referring clinical partner or because of the patient’s wish.
4 Septic arthritis of the shoulder joint in a 69-year-old man following shoulder arthroscopy and supraspinatus muscle refixation. The joint effusion is appreciated on the axial T2-weighted fat-saturated images (open arrows in 4A). The strong synovialitis (arrows) is clearly evidenced on the contrast-enhanced coronal (4B) and axial (4C) MR images (4B without and 4C with fat saturation).

5 Acute pulmonary embolism in both pulmonary arteries shown on T1/T2-weighted coronal TrueFISP images (arrows; this examination was obtained in a 64-year-old patient with renal insufficiency and suspected pulmonary embolism, being referred for non-contrast-enhanced MRI).
Musculoskeletal Imaging at 3T with Simultaneous Use of Multipurpose Loop Coils

Elena Ferrer¹; Rafael Coronado Santos²

¹Radiology Department, Clínica Creu Blanca, Barcelona, Spain,
²Imaging & Therapy Division, Siemens S.A. Healthcare Sector, Madrid, Spain

Introduction
The goal of this paper is to show how the simultaneous use of multipurpose Loop Coils in magnetic resonance imaging (MRI) enables high-resolution musculoskeletal (MSK) studies with an increased level of contrast and specificity for assessing muscles, tendons, ligaments, joints, cartilage, etc. and how this imaging procedure helps to obtain accurate clinical diagnoses. In the following sections, we present six different daily patient routine examinations carried out at Clínica Creu Blanca, a leading Spanish institution in Sports Medicine MRI (Fig. 1).

Material and methods
All MRI exams shown in this article were performed at 3 Tesla open bore system with TrueForm technology (MAGNETOM Verio, Siemens Healthcare, Germany), equipped with 32-channel (Tim [102 × 32] configuration) in combination with multipurpose Loop Coils and Flex Coil interfaces. Loop Coils come in three sizes (Fig. 2), Large (11 cm diameter), Medium (7 cm diameter) and Small (4 cm diameter). They are iPAT-compatible (integrated Parallel Acquisition Technique) in combination with other coils and can be combined with any coil and the lower part of the 32-channel Head Coil (Table 1). The Flex Coil Interface is not permanently mounted and therefore allows flexible coil positioning. The imaging protocols include axial, coronal and sagittal Proton Density-weighted (PD) Turbo Spin Echo (TSE) sequences with and without Fat Saturation (Fat Sat) and T1-weighted TSE.

Patient and coil positioning
Correct patient positioning and the selection of right loop coils for the region-of-interest have a huge influence on image quality (e.g. to avoid coil filling-factor). In addition, we have to make sure that the patient is positioned comfortably to make the scan bearable and to reduce the risk of patient movements. The following six examinations – as performed at Clínica Creu Blanca’s daily routine – explain how we do it and the results that can be achieved.
<table>
<thead>
<tr>
<th>Flex Coil Interface</th>
<th>Loop Coils</th>
<th>Loop Coil, large</th>
<th>Loop Coil, medium</th>
<th>Loop Coil, small</th>
</tr>
</thead>
<tbody>
<tr>
<td>General</td>
<td></td>
<td>No coil tuning</td>
<td>No coil tuning</td>
<td>No coil tuning</td>
</tr>
<tr>
<td></td>
<td></td>
<td>iPAT-compatible in combination with other coils</td>
<td>iPAT-compatible in combination with other coils</td>
<td>iPAT-compatible in combination with other coils</td>
</tr>
<tr>
<td>Applications</td>
<td></td>
<td>Examination of upper or lower extremities (e.g. shoulder, axilla)</td>
<td>Examination of inner ear, wrist and fingers, pediatric examinations(^*)</td>
<td>Examination of small structures near the surface (e.g. joints of fingers and toes, wrist, skin, temporomandibular joints (TMJ))</td>
</tr>
<tr>
<td>Can be combined with</td>
<td></td>
<td>Any coil and the lower part of the 32-channel Head Coil</td>
<td>Any coil and the lower part of the 32-channel Head Coil</td>
<td>Any coil and the lower part of the 32-channel Head Coil</td>
</tr>
<tr>
<td>Cannot be combined with</td>
<td></td>
<td>The complete 32-channel Head Coil and the complete 32-channel Body Coil</td>
<td>The complete 32-channel Head Coil and the complete 32-channel Body Coil</td>
<td>The complete 32-channel Head Coil and the complete 32-channel Body Coil</td>
</tr>
<tr>
<td>Weight</td>
<td></td>
<td>225 g</td>
<td>175 g</td>
<td>200 g</td>
</tr>
<tr>
<td>Diameter</td>
<td></td>
<td>110 mm</td>
<td>70 mm</td>
<td>4 mm</td>
</tr>
</tbody>
</table>

\(^*\) MR scanning has not been established as safe for imaging fetuses and infants under two years of age. The responsible physician must evaluate the benefit of the MRI examination in comparison to other imaging procedures.
Carpals examination

This examination was performed using three Flex Coil Interfaces and three 7 cm Loop Coils (Fig. 3). The obtained images are depicted in Figure 4.

Cables should not be crossed otherwise there will be a signal error when a three plane localizer is launched.
(4A) Coronal PD Turbo Spin Echo (TSE) Fat Sat, TR 3500 ms, TE 44 ms, Slice thickness (SL) 1.7 mm, in-plane resolution 0.4 x 0.4 mm², matrix 192 x 192 px².
(4B) Coronal T₁w TSE, TR 1290 ms, TE 12 ms, SL 1.7 mm, in-plane resolution 0.3 x 0.3 mm², matrix 320 x 320.
(4C) Axial PD TSE Fat Sat, TR 4400 ms, TE 47 ms, SL 2 mm, in-plane resolution 0.3 x 0.3 mm², matrix 256 x 256 px².
(4D) Axial PD TSE, TR 2400 ms, TE 34 ms, SL 2 mm, in-plane resolution 0.3 x 0.3 mm², matrix 256 x 256 px².
(4E) Sagittal PD TSE Fat Sat, TR 4100 ms, TE 44 ms, SL 1.8 mm in-plane resolution 0.4 x 0.4 mm², matrix 192 x 192 px².
(4F) Sagittal T₁w TSE, TR 1000 ms, TE 12 ms, SL 1.8 mm, in-plane resolution 0.3 x 0.3 mm², matrix 320 x 320 px².
Thumb metacarpals examination

This examination was performed using two Flex Coil Interfaces and two 4 cm Loop Coils (Fig. 5). The obtained images are depicted in Figure 6.
Axial PD TSE Fat Sat, TR 5200 ms, TE 47 ms, SL 2 mm, in-plane resolution 0.3 x 0.3 mm², matrix 256 x 256 px².

PD TSE, TR 3000 ms, TE 35 ms, SL 2 mm, in-plane resolution 0.3 x 0.3 mm², matrix 256 x 256 px².

Sagittal PD TSE Fat Sat, TR 4100 ms, TE 44 ms, SL 1 mm, in-plane resolution 0.4 x 0.4 mm², matrix 192 x 192 px².

Sagittal T1w TSE, TR 810 ms, TE 12 ms, SL 1 mm, in-plane resolution 0.3 x 0.3 mm², matrix 256 x 256 px².

Coronal PD TSE Fat Sat, TR 3500 ms, TE 44 ms, SL 1.2 mm, in-plane resolution 0.4 x 0.4 mm², matrix 192 x 192 px².

Coronal T1w TSE, TR 650 ms, TE 12 ms, SL 1.2 mm, in-plane resolution 0.3 x 0.3 mm², matrix 256 x 256 px².
Ankle examination
This examination was performed using three Flex Coil Interfaces and three 7 cm Loop Coils (Fig. 7). The obtained images are depicted in Figure 8.

U-shape cushion to place the feet and fix the ankle with a strap.

Flex Interface Coils are separated by flat pads and a couple of straps are used to immobilize the ankle.

The triangular leg pad should be placed to increase patient comfort and to avoid lumbar lordosis.

Patient and coil positioning for ankle examinations.
Orthopedic Imaging

Clinical

(8A) Sagittal PD TSE, Fat Sat, TR 5200 ms, TE 52 ms, SL 2.5 mm, in-plane resolution 0.4 × 0.4 mm², matrix 320 × 320 px².

(8B) Sagittal T1w TSE, TR 719 ms, TE 10 ms, SL 2.5 mm, in-plane resolution 0.4 × 0.4 mm², matrix 320 × 320 px².

(8C) Axial PD TSE, TR 4140 ms, TE 44 ms, SL 1.8 mm, in-plane resolution 0.3 × 0.3 mm², matrix 384 × 384 px².

(8D) Axial PD TSE, Fat Sat, TR 8710 ms, TE 40 ms, SL 1.8 mm, in-plane resolution 0.4 × 0.4 mm², matrix 320 × 320 px².

(8E) Coronal T1w TSE, TR 719 ms, TE 12 ms, SL 2 mm, in-plane resolution 0.3 × 0.3 mm², matrix 320 × 320 px².

(8F) Coronal PD TSE, Fat Sat, TR 5400 ms, TE 50 ms, SL 2 mm, in-plane resolution 0.3 × 0.3 mm², matrix 320 × 320 px².
Metacarpals or fingers examination

This examination was performed using two Flex Coil Interfaces and two 7 cm Loop Coils (Fig. 9). The obtained images are depicted in Figure 10.

---

Patient and coil positioning for metacarpals or fingers examination.

9A 9B 9C

Rectangular pad

9D

9E

9F

---

2 Patient and coil positioning for metacarpals or fingers examination.
(10A) Sagittal PD TSE Fat Sat, TR 4100 ms, TE 44 ms, SL 1.3 mm, in-plane resolution 0.5 × 0.5 mm², matrix 192 × 192 px².

(10B) Sagittal T1w TSE, TR 1000 ms, TE 12 ms, SL 1.3 mm, in-plane resolution 0.3 × 0.3 mm², matrix 320 × 320 px².

(10C) Axial PD TSE Fat Sat, TR 5600 ms, TE 52 ms, SL 2.5 mm, in-plane resolution 0.3 × 0.3 mm², matrix 256 × 256 px².

(10D) Axial PD TSE, TR 3000 ms, TE 34 ms, SL 2.5 mm, in-plane resolution 0.3 × 0.3 mm², matrix 256 × 256 px².
Distal inter-phalangeal examination

This examination was performed using two Flex Coil Interfaces and two 4 cm Loop Coils (Fig. 11). The obtained images are depicted in Figure 12.
(12A) Axial PD TSE Fat Sat, TR 5200 ms, TE 47 ms, SL 2 mm, in-plane resolution 0.3 × 0.3 mm², matrix 256 × 256 px².

(12B) Axial PD TSE, TR 3000 ms, TE 35 ms, SL 2 mm, in-plane resolution 0.3 × 0.3 mm², matrix 256 × 256 px².

(12C) Sagittal PD TSE Fat Sat, TR 4100 ms, TE 44 ms, SL 1 mm, in-plane resolution 0.4 × 0.4 mm², matrix 192 × 192 px².

(12D) Sagittal T1w TSE, TR 810 ms, TE 12 ms, SL 1 mm, in-plane resolution 0.3 × 0.3 mm², matrix 256 × 256 px².

(12E) Coronal PD TSE Fat Sat, TR 3500 ms, TE 44 ms, SL 1.2 mm, in-plane resolution 0.4 × 0.4 mm², matrix 192 × 192 px².

(12F) Coronal T1w TSE, TR 650 ms, TE 12 ms, SL 1.2 mm, in-plane resolution 0.3 × 0.3 mm², matrix 192 × 192 px².

(12G) This figure shows an angiography of a finger whose third phalange was amputated: coronal FLASH 3D post contrast, TR 1.5 ms, TE 3.82 ms, SL 0.7 mm, in-plane resolution 0.7 × 0.7 mm, matrix 192 × 174 px².
Elbow examination

This examination was performed using three Flex Coil Interfaces and three 7 cm Loop Coils (Fig. 13). The obtained images are depicted in Figure 14.

Conclusion

High-resolution MSK imaging can be acquired using a 3T magnetic field (MAGNETOM Verio) and a combination of multipurpose Loop Coils. It does not claim to replace dedicated MSK coils (e.g. knee coil, hand coil) but it might represent an alternative method for institutions whose number of MSK studies is scarce and/or that are not concerned about reducing MSK examination times and/or increasing patient throughput but that want to achieve similar levels of diagnostic accuracy as dedicated coils [1–6].

References

**14A (14A)** Axial PD TSE Fat Sat, TR 4400 ms, TE 47 ms, SL 2 mm, in-plane resolution 0.5 × 0.5 mm², matrix 256 × 256 px².

**14B (14B)** Axial PD TSE, TR 2700 ms, TE 41 ms, SL 2 mm, in-plane resolution 0.4 × 0.4 mm², matrix 320 × 320 px².

**14C (14C)** Coronal PD TSE Fat Sat, TR 5000 ms, TE 44 ms, SL 1.8 mm, in-plane resolution 0.5 × 0.5 mm², matrix 192 × 192 px².

**14D (14D)** Coronal T1w TSE, TR 1000 ms, TE 12 ms, SL 1.8 mm, in-plane resolution 0.3 × 0.3 mm², matrix 320 × 320 px².

**14E (14E)** Sagittal PD TSE Fat Sat, TR 4800 ms, TE 41 ms, SL 2 mm, in-plane resolution 0.5 × 0.5 mm², matrix 192 × 192 px².

**14F (14F)** Sagittal T1w TSE, TR 1000 ms, TE 12 ms, SL 2 mm, in-plane resolution 0.3 × 0.3 mm², matrix 320 × 320 px².
Automated Morphological Knee Cartilage Analysis of 3D MRI at 3T


1 The Australian E-Health Research Centre, CSIRO ICT Centre, Brisbane, Australia
2 Steadman Philippon Research Institute, Vail, Colorado, USA
3 School of Human Movement Studies, University of Queensland, Brisbane, Australia
4 Siemens Healthcare, Erlangen, Germany
5 Leeds Musculoskeletal Biomedical Research Unit, Leeds, University of Leeds, Leeds, UK
6 School of Information Technology & Electrical Engineering, University of Queensland, Brisbane, Australia.

Introduction
Quantitative magnetic resonance (MR) imaging including morphological and biochemical measurements have shown promise for detecting early cartilage changes [1, 2]. Morphological measures (volume, thickness, surface area and curvature) can be derived from MR images and have been shown to vary with osteoarthritis (OA) [2, 3]. As biochemical changes occur before gross tissue loss, improved sensitivity for early OA detection requires using quantitative cartilage mapping techniques such as T2, T2*, T1rho, dGEMERIC*, Sodium and CEST [4]. Although quantitative MR has been used extensively in research, it is still not commonly used clinically for multiple reasons, mostly related to limited validation of reproducibility and the time required to perform analyses. Extracting accurate and reproducible quantitative cartilage measures from MR images has been the focus of significant research but remains an open problem [5]. In recent studies, it has been found that different research groups can obtain notably different morphometric results, reflecting subtle variations in regional definitions of anatomy, methods and calculation techniques [6]. A desired direction (although difficult to achieve), is to standardize and automate processes for quantitative cartilage analyses which would improve reproducibility and post-processing time necessary for clinical application. An important aspect of this process would include automated segmentation and standardizing the sequences used for acquisition.

1 Visual comparison of all sequences acquired for representative subject; (1A) Sagittal T2w DESS (TR 16.3 ms; TE 4.72 ms; flip angle 25°), (1B) Sagittal PDw SPACE (1200 ms; 32 ms; 120°), (1C) Sagittal T1w VIBE (20 ms; 5.74 ms; 12°).
In recent years several different approaches have been developed to perform automated segmentation of articular cartilage from MR images [7–11]. A primary limitation of previous work has been limited validation of accuracy especially on pathological cohorts and different scanning sequences. Gradient echo sequences such as DESS and FLASH have most commonly been used for morphologic measurements because they provide relatively homogeneous cartilage signal and excellent delineation of the cartilage-bone interface [12–16]. Spin echo sequences have more clinical value due to their larger dynamic range and signal contrast resolution within the cartilage, other soft tissue and bone, although this makes segmentation more challenging. Scan times for the sequence chosen for segmentation must also be considered. Ideally, the scans used for quantitative cartilage segmentation and analyses would also be readily used for clinical diagnosis since patient scan time is limited.

In this article we will review the performance of a new automated segmentation algorithm for the knee cartilages. An aim of this work is to evaluate the accuracy and reproducibility of this automated approach when the scheme is applied to scans from asymptomatic volunteers acquired at 3T with different 3D MR sequences (T2w DESS, T1w VIBE and PDw SPACE), different coils (8 and 15-channel knee coils), and test re-test (3 scans of the same subject/s repositioned between scans). Additionally, initial results will be presented with scans from pathological subjects imaged using PDw SPACE.

**Method**

**Data sets:**

**Asymptomatic subjects:**

3D sequence comparison

MR imaging of the knee joint was performed on 10 healthy volunteers (3 males and 7 females; aged 23 to 45 years, mass 50-90 kg). Informed consent was obtained and the medical research ethics committee of the University of Queensland approved the current study. All images were acquired at the University of Queensland on a 3T MR scanner (MAGNETOM Trio, Siemens Healthcare, Erlangen, Germany) using an 8-channel transmit-receive knee coil. The knee for each subject was imaged using three 3D MR sequences; T2w DESS, T1w VIBE and PDw SPACE. A slice from each scan sequence of one subject can be seen in figure 1, illustrating the difference in image contrast, especially at the bone-cartilage and cartilage-synovial interfaces. The scanning parameters are given in table 1.

The articular cartilage in each DESS image was manually segmented on every sagittal slice in the volume by a musculoskeletal expert using the ROI plugin in ImageJ [17] taking on average 2 hours. Segmentations for other MR sequences were obtained by independently co-registering the sub images within a bounding box around the manual DESS segmentation (dilated by 7 mm) for each cartilage plate (implemented using ITK [18]). The segmentation masks were not adjusted to match the sequence.

**Test re-test**

MR imaging was performed on 16 healthy volunteers (5 males and 11 females; aged 23 to 52). Informed consent was obtained and the local research ethics committee approved the current study. All images were acquired and anonymized at the Leeds Musculoskeletal Biomedical Research Unit on a 3T MR scanner (MAGNETOM Verio, Siemens Healthcare, Erlangen, Germany) using 8 and 15-channel transmit-receive knee coils. The knee for each subject

![Visual comparison of automated segmentation results (colormap represents thickness streamlines). Arrows point to several focal errors. These slices correspond to those presented in figure 1.](image)

*A licensed physician may choose to use FDA-approved contrast agents in conjunction with an MRI examination, based on his/her medical opinion and discretion and in accordance with the instructions for use and indications for use supplied by the pharmaceutical manufacturer for the contrast agents.*
### Table 1: MR sequence parameters used in this study, from the University of Queensland (UQ), University of Leeds (UL) and the Steadman Philippon Research Institute (SPRI).

<table>
<thead>
<tr>
<th>Sequence</th>
<th>DESS</th>
<th>DESS</th>
<th>VIBE</th>
<th>SPACE</th>
<th>SPACE</th>
</tr>
</thead>
<tbody>
<tr>
<td>SITE</td>
<td>UL</td>
<td>UQ</td>
<td>UQ</td>
<td>UQ</td>
<td>SPRI</td>
</tr>
<tr>
<td>N</td>
<td>16</td>
<td>10</td>
<td>10</td>
<td>10</td>
<td>3</td>
</tr>
<tr>
<td>Acquisition time</td>
<td>11:10</td>
<td>9:44</td>
<td>10:14</td>
<td>08:40</td>
<td>04:46</td>
</tr>
<tr>
<td>Phase encoding direction</td>
<td>AP</td>
<td>AP</td>
<td>AP (RL, N=3)</td>
<td>AP</td>
<td>AP</td>
</tr>
<tr>
<td>Number of slices</td>
<td>160</td>
<td>160</td>
<td>80</td>
<td>176</td>
<td>176</td>
</tr>
<tr>
<td>Slice thickness (mm)</td>
<td>0.70</td>
<td>0.70</td>
<td>1.50</td>
<td>0.60</td>
<td>0.70</td>
</tr>
<tr>
<td>Slice spacing (mm)</td>
<td>0.36 x 0.36</td>
<td>0.36 x 0.46</td>
<td>0.31 x 0.31</td>
<td>0.49 x 0.49</td>
<td>0.60 x 0.60</td>
</tr>
<tr>
<td>Image matrix</td>
<td>384 x 384</td>
<td>384 x 312</td>
<td>512 x 512</td>
<td>320 x 320</td>
<td>256 x 256</td>
</tr>
<tr>
<td>Phase encoding steps</td>
<td>308</td>
<td>250</td>
<td>512</td>
<td>294</td>
<td>151</td>
</tr>
<tr>
<td>Fat suppression</td>
<td>WE</td>
<td>WE</td>
<td>WE</td>
<td>SPAIR</td>
<td>SPAIR</td>
</tr>
<tr>
<td>GRAPPA factor/ref lines</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>2/24</td>
<td>2/24</td>
</tr>
<tr>
<td>Number of averages</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1.4</td>
<td>1</td>
</tr>
<tr>
<td>Slice resolution</td>
<td>80%</td>
<td>80%</td>
<td>100%</td>
<td>91.88%</td>
<td>100%</td>
</tr>
<tr>
<td>Slice oversampling</td>
<td>0%</td>
<td>10%</td>
<td>0%</td>
<td>0%</td>
<td>0%</td>
</tr>
<tr>
<td>Phase resolution</td>
<td>100%</td>
<td>81%</td>
<td>100%</td>
<td>100%</td>
<td>100%</td>
</tr>
<tr>
<td>Phase oversampling</td>
<td>0%</td>
<td>10%</td>
<td>0%</td>
<td>0%</td>
<td>0%</td>
</tr>
<tr>
<td>Turbo factor/echo spacing/echo train</td>
<td>1/0/0</td>
<td>1/0/0</td>
<td>1/0/0</td>
<td>53/-/-</td>
<td>84/4.98/284</td>
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<tr>
<td>Repetition time (TR) ms</td>
<td>16.3</td>
<td>16.3</td>
<td>20</td>
<td>1200</td>
<td>1200</td>
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<tr>
<td>Echo time (TE) ms</td>
<td>4.72</td>
<td>4.72</td>
<td>5.65</td>
<td>32</td>
<td>45</td>
</tr>
<tr>
<td>Flip angle</td>
<td>25</td>
<td>25</td>
<td>12</td>
<td>PdVar</td>
<td>PdVar</td>
</tr>
<tr>
<td>Bandwidth (Hz/pixel)</td>
<td>352</td>
<td>352</td>
<td>130</td>
<td>539</td>
<td>425</td>
</tr>
</tbody>
</table>

WE = water excitation
was imaged using a DESS sequence with parameters outlined in table 1, and involved, scanning twice with the 8-channel coil and once with a 15-channel coil. The subject was taken off the table and repositioned between each scan. The articular cartilage in the first 8-channel coil DESS images was manually segmented on every fourth sagittal slice from the original MR by a trained radiographer then reviewed and adjusted where necessary by a musculoskeletal radiologist with experience of cartilage segmentation using Analyze 10.

Pathological case studies
For qualitative evaluation of the automated segmentation performance with clinical scans, automated segmentation from three patient scans obtained preoperatively were qualitatively assessed. All images were acquired and anonymized at the Steadman Clinic on a 3T MR scanner (MAGNETOM Verio, Siemens Healthcare, Erlangen, Germany) for this study by the Steadman Philippon Research Institute (SPRI) with approval from the Vail Valley Medical Center Internal Review Board. The knee for each patient was imaged using a 15-channel knee coil with the SPACE sequence with parameters outlined in table 1. Subject 1 was a patient with excessive synovial fluid and synovitis in the joint space. Subject 2 was a patient with extensive cartilage damage. Cartilage damage was graded arthroscopically using the International Cartilage Repair Society (ICRS) grading system. The femoral cartilage lesions included grade 4 on the medial femoral condyle, grade 2 on the lateral femoral condyle, and grade 2 in the central trochlea. The tibial cartilage lesions included grade 3 on the posterior lateral tibial plateau. Grades 2 and 3 patellar lesions were also present. Subject 3 was a patient with moderate cartilage damage. The femoral cartilage damage included grade 2 damage to the lateral femoral condyle. The damage to the tibial cartilage included diffuse grade 1 damage to the lateral plateau, and grade 3 damage to the anterior lateral plateau. The patella cartilage had grades 2 and 3 damage.

Analysis:
MRI automatic segmentation
The MR images from each subject were automatically segmented using an improved version of a recently published 3D shape model based scheme [11]. The improvements primarily consist of optimization and tradeoffs for speed (total computation time 4 to 5 minutes), multi-sequence performance and some changes to better handle MR artifacts (particularly bias-fields) and pathologies such as synovitis.

The basic scheme consists of four stages 1) preprocessing, 2) atlas alignment 3) bone segmentation and 4) cartilage segmentation.

1) Preprocessing: Noise on raw MR images was reduced with a median filter implemented in ITK [18].
2) Atlas alignment: An average MR atlas image was affinely registered to the subject. The obtained transform was used to propagate the bone atlas surface into the preprocessed image and then estimate pose and shape parameters. The atlas surface was partitioned into 8 regions (patella, medial and lateral tibial, medial and lateral femoral condyle, medial and lateral central femoral and femoral trochlea).
3) Bone segmentation: The bone (femur, tibia and patella) is segmented

<table>
<thead>
<tr>
<th>Table 2: Computational efficiency of our approach.</th>
</tr>
</thead>
<tbody>
<tr>
<td>CPU clock speed:</td>
</tr>
<tr>
<td>Machine CPU count:</td>
</tr>
<tr>
<td>Machine memory:</td>
</tr>
<tr>
<td>OS</td>
</tr>
<tr>
<td>Compiler</td>
</tr>
<tr>
<td>Libraries</td>
</tr>
<tr>
<td>Image dimensions</td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td>Multi-threaded</td>
</tr>
<tr>
<td>Total segmentation time (min)</td>
</tr>
<tr>
<td>Peak memory used:</td>
</tr>
</tbody>
</table>
using an adaptive threshold shape based segmentation scheme described in Fripp et al. [19]. This technique was improved by including robust regional thresholds to handle signal attenuation commonly found in the femoral-patella region, as well as aliasing artefacts that can cover the patella bone.

4) Cartilage segmentation: The cartilage segmentation scheme is based on the technique described in Fripp et al. [11, 19]. This technique was improved by including the aforementioned robust regional cartilage statistics, and secondly an explicit synovial fluid segmentation was performed.

Starting from the bone segmentation, a statistical model extracts the bone-cartilage interface (BCI). The thickness based cartilage segmentation is performed iteratively driven locally by local edges and estimated tissue probabilities within each region. Initially it assumes that all bright tissue is cartilage, and uses a statistical model to constrain the estimated cartilage thickness. Possible synovial fluid patches are then detected as non cartilage tissues with high signal intensity. If found, the 'synovial fluid' tissue probabilities are then combined with the other tissue probabilities and local edge information to estimate the local thickness which is statistically constrained. At each iteration tissue and edge properties are re-estimated, local areas that definitely have no cartilage are disabled, and local thickness is estimated within a reduce search space to refine the cartilage segmentation.

Validation of automatic segmentation

The extent of spatial overlap between the manual segmentations (A) and automatic cartilage segmentations (B) were evaluated for the femoral, tibial, and patellar cartilage using the Dice similarity index (DSI) [20]. The DSI ranges from 0 (no overlap in segmenta-
Table 3: Average and standard deviation of the validation measures comparing automatic and manual segmentations for each cartilage plate and MR sequence in the University of Queensland

<table>
<thead>
<tr>
<th>Cartilage</th>
<th>Site</th>
<th>Sequence</th>
<th>N</th>
<th>DSI</th>
<th>Median DSI</th>
<th>Vol. RD (%)</th>
<th>R</th>
</tr>
</thead>
<tbody>
<tr>
<td>Patellar</td>
<td>UL</td>
<td>DESS</td>
<td>16</td>
<td>82.74±6.11</td>
<td>85.15</td>
<td>11.58±11.06</td>
<td>0.95</td>
</tr>
<tr>
<td></td>
<td>UQ</td>
<td>DESS</td>
<td>10</td>
<td>83.44±4.65</td>
<td>84.61</td>
<td>9.11±6.1</td>
<td>0.98</td>
</tr>
<tr>
<td></td>
<td>UQ</td>
<td>SPACE</td>
<td>10</td>
<td>79.77±4.46</td>
<td>80.74</td>
<td>10.49±10.75</td>
<td>0.90</td>
</tr>
<tr>
<td></td>
<td>UQ</td>
<td>VIBE</td>
<td>10</td>
<td>82.57±5.53</td>
<td>84.10</td>
<td>7.98±20.16</td>
<td>0.76</td>
</tr>
<tr>
<td>Tibial</td>
<td>UL</td>
<td>DESS</td>
<td>16</td>
<td>83.3±1.68</td>
<td>82.97</td>
<td>4±5.83</td>
<td>0.98</td>
</tr>
<tr>
<td></td>
<td>UQ</td>
<td>DESS</td>
<td>10</td>
<td>80.47±2.23</td>
<td>81.48</td>
<td>12.39±8.73</td>
<td>0.97</td>
</tr>
<tr>
<td></td>
<td>UQ</td>
<td>SPACE</td>
<td>10</td>
<td>79.08±3</td>
<td>79.46</td>
<td>-5.06±9.26</td>
<td>0.97</td>
</tr>
<tr>
<td></td>
<td>UQ</td>
<td>VIBE</td>
<td>10</td>
<td>78.14±3.05</td>
<td>78.85</td>
<td>15.26±10.85</td>
<td>0.94</td>
</tr>
<tr>
<td>Femoral</td>
<td>UL</td>
<td>DESS</td>
<td>16</td>
<td>82.64±2.85</td>
<td>82.18</td>
<td>-1.51±7.31</td>
<td>0.94</td>
</tr>
<tr>
<td></td>
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<td>DESS</td>
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<td>83.87±2.04</td>
<td>83.88</td>
<td>1.79±6.27</td>
<td>0.93</td>
</tr>
<tr>
<td></td>
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<td>SPACE</td>
<td>10</td>
<td>80.14±2.72</td>
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<td>2.71±5.28</td>
<td>0.91</td>
</tr>
<tr>
<td></td>
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<td>VIBE</td>
<td>10</td>
<td>80.76±2.73</td>
<td>81.30</td>
<td>4.46±9.21</td>
<td>0.88</td>
</tr>
</tbody>
</table>

Results

Asymptomatic subjects: 3D sequence comparison

The results of the segmentation on each sequence for a representative subject can be seen in figures 2 and 3, whose computational efficiency is outlined in table 2. Overall, the segmentation of the DESS appears qualitatively to have the highest accuracy, likely due to the good contrast-to-noise at both the cartilage-bone and cartilage-synovial fluid interfaces. The VIBE sequence was reasonably accurate, although the larger slice thickness and lower cartilage-synovial contrast, especially when imaged coro-

nally resulted in mis-segmentation in the patellar-femoral and tibial-femoral cartilage regions. Although the cartilage tissue is inhomogeneous and the bone-cartilage interface can be indistinct, the syno SPACE sequence generally worked well, and due to the good cartilage-synovial contrast overall its precision was similar to the VIBE. A summary of the quantitative validation for all datasets is provided in table 3. It can be seen that the DESS sequence obtained the highest and most consistent DSI in all cartilage regions (80–84%), while the VIBE (78–83%) and SPACE (79–80%) obtained fairly accurate
and consistent results. The manual DESS volumes and those calculated automatically for the DESS (R = 0.93–0.98 in both UQ and UL datasets), VIBE (R = 0.76–0.94) and SPACE (R = 0.91–0.97) were strongly correlated (Table 3). As the manual volumes were calculated on the DESS, there could be a bias towards the DESS, resulting in improved accuracy and stronger correlations. To fairly compare the sequences, manual segmentations of the SPACE and VIBE sequences should be performed.

In this study no gross segmentation errors were observed. The most common segmentation errors were focal errors, particularly at the cartilage interfaces (patella-femoral and femoral-tibial) as well as towards the ends of the cartilage regions (top of patellar, edges of the tibial). In a few cases, the cartilage signal drops in the inter-chondylar notch, sometimes resulted in mis-segmentation of cartilage as bone. In most scans, a non-uniform signal attenuation around the patella-femoral region was observed, which made detection of and separation of cartilage-synovial fluid more difficult (especially for VIBE) and increasing the amount of mis-segmentation in a few cases.

**Asymptomatic subjects:**

**test re-test and different coils**

Bland-Altman plots of the (patellar, tibial and femoral) cartilage volume from the three scans of all 16 subjects can be seen in figure 4, and had an average coefficient of variation of (4.7, 2.7, 2.2)%.

The automated volumetric analysis of the repeated scans with the 8-channel coil showed strong correlation (R = 0.97, 0.97, 0.99), with all but four (of 48) cartilage plates having an absolute volume difference of less than 3%, three of these were patellar (4.2–5.5%) and one tibial (5.5%). Comparing the volume estimated automatically from the 8-channel and 15-channel coils also showed strong correlation (R = 0.96, 0.98, 0.99), however, the volume estimated from the 8-channel coil appears

![Bland-Altman plot comparing automatic cartilage volume obtained for the (top) patella and (middle) tibial (bottom) femoral cartilages.](image)
systemically larger than that of the 15-channel coil in all cartilage plates (between 0.67–5.5%). Manual analysis is required to determine if this bias is due to actual MR imaging differences or simply a bias in the segmentation algorithm due to subtle changes in the image contrast or other effects.

Case studies: pathological cases
Figures 5 to 8 show the SPACE sequence with overlaid automated segmentation contour for each subject (1, 2, 2 and 3 respectively).

The segmentation results for subject 1 were excellent and the excessive joint fluid and synovitis did not appear to have a significant effect on the segmentation. There were some segmentation errors where the superficial femoral and tibia cartilage was under-segmented at the medial tibial femoral interface (Fig. 5). There was also some under-segmentation of the medial patellar cartilage. Subject 2 had more severe cartilage damage and there were some segmentation errors. Figure 6 shows an ICRS grade 4 lesion on the medial femoral condyle and a resulting segmentation error. The segmentation performed well on an ICRS grade 2 lesion to the lateral femoral condyle, shown in figure 7. Again under-segmentation errors were noticed at the tibial femoral interface. The trochlea also exhibited under-segmentation.

The segmentation for Subject 3 was excellent. The ICRS grade 2 cartilage lesion on the patella was well segmented as shown in figure 8, although there was some over-segmentation of the patella into the synovial fluid.

Discussion
The purpose of this paper was to assess the accuracy and repeatability of a novel automated cartilage segmentation routine.

The automatic segmentation of the DESS sequence had the highest correlation with the manual segmentations defined originally on the DESS. The SPACE and VIBE had lower correlation and precision, making the detection of very subtle longitudinal changes more challenging. These results were not surprising, with VIBE having poor cartilage-synovial and muscle contrast, while SPACE has a less homogeneous cartilage appearance and the bone-cartilage interface (BCI) is less well defined [21]. However, as SPACE allows more accurate detection and grading of cartilage lesions and other internal cartilage changes compared to gradient echo techniques and thus has more clinical value [22, 23]. The scan time for SPACE is significantly faster than the other sequences, even if parallel imaging was enabled.

The shape and thickness based segmentation scheme was performed directly on the 3D scans and the results did not show any critical problems due to image contrast, resolution, sectioning or planar choice. In general the areas that cause the most errors and variability are at low contrast interfaces, particularly carti-
age-cartilage. Strong signal inhomogeneity was observed to cause some focal mis-segmentation, particularly by cartilage being mis-identified as synovial fluid. Although the quantitative analysis for the coronal VIBE images was reasonable, errors due to partial voluming artifacts were observed in the patellar and femoral chondyles. For reproducible automated analysis using our technique a higher slice resolution would be recommended.

The automatic segmentation on the DESS sequence was found to have high same day reproducibility when repositioned and using two different knee coils. Manual segmentations of all scans showed promise for use of the auto-segmentation routinely with patients. Areas with segmentation errors included the interface between the tibial and femoral cartilage, the superior tibial cartilage, and the medial patella. Excessive joint fluid and synovitis did not appear to affect the segmentation results. The subject with the most severe cartilage damage had the least successful segmentation results which may indicate that current segmentation regime is limited to less severe cases.

Segmentation processing time is very important for routine clinical use, the current algorithm is fast enough (< 5 minutes) to be run while other images are being scanned, allowing results to be available for assessment by radiologists at the end of the MRI session. Further validation using different image contrasts, resolutions, field strengths and coils as well as pathologic scans is required, especially to investigate any biases that could occur.

Although, the initial case reports show promise with pathologic cases further quantitative comparison on a larger dataset between manual and automated segmentation is required. There are several clinical applications of automated cartilage segmentation. Manual segmentation is too time intensive and cannot be implemented into the routine clinical sessions. Automated cartilage segmentation allows for quantitative assessment of the morphological and biochemical state of the cartilage. Morphological cartilage data can be used for treatment planning and can be tracked over time to determine progression or healing of cartilage damage. Additionally, cartilage segmentations can be registered to biochemical sequences such as T2 and T1-mapping which have been shown to be more sensitive to early cartilage degeneration before morphological measurements.
Auto-segmentation results from Subject 2. Good segmentation results were noticed at location of an ICRS grade 2 cartilage lesion on the femoral condyle (see arrow head). The superficial tibial cartilage is under-segmented (see arrow).

Auto-segmentation results from Subject 3. Good segmentation results were noticed at location of an ICRS grade 2 cartilage lesion on the patella (see arrow). There was some over-segmentation of the patella into the synovial fluid (see arrow head).
Conclusion
The accuracy and reproducibility of automatic cartilage segmentation and subsequent volume measures obtained from T2w DESS, T1w VIBE and PDw SPACE were assessed. The validation results indicate that the automatic segmentation algorithm performed best on the T2w DESS images. The SPACE sequence has lower accuracy automated morphological cartilage assessment, although less accurate than DESS.

Acknowledgements
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References
MAGNETOM Prisma Image Gallery

MAGNETOM Prisma*, Siemens’ new and powerful 3T MRI system, is built to tackle the most demanding research challenges. It delivers maximum performance under prolonged high-strain conditions opening new possibilities for imaging functional processes and understanding the most threatening diseases. Only one of many high-performance features is the new XR 80/200 gradient system. With its higher gradient amplitude it delivers significantly higher sig-

*The product is under development and not commercially available. Its future availability cannot be ensured.
nal-to-noise ratio, enhancing for example physiological imaging or morphometric measurements. With MAGNETOM Prisma’s higher spatial and temporal resolution you can see excellent anatomical detail, for example displaying functional and structural brain connectivity or smallest lesions with zoomed MRI. Additionally, it delivers benchmark 3T magnet homogeneity – the basis for superior quantitative evaluations.
Product News

Hips, PD TSE SPAIR coronal, PAT factor 2, matrix 448, 18-channel Body/32-channel Spine coil

Wrist, PD TSE transversal, PAT factor 2, matrix 320, 16-channel Hand/Wrist coil

Hips, T1 TSE coronal, PAT factor 2, matrix 512, 18-channel Body/32-channel Spine coil

Knee, PD TSE FatSat coronal, PAT factor 2, matrix 384, 15-channel Knee coil

Ankle, PD TSE FatSat sagittal, PAT factor 2, matrix 384, 16-channel Foot/Ankle coil

Ankle, T1 3D VIBE water excitation sagittal, PAT factor 2, matrix 384, 16-channel Foot/Ankle coil

Ankle, T2 TSE transversal, matrix 320, 16-channel Foot/Ankle coil
**The New, High-Performance MR Gradient System XR 80/200. Design, Benefits and Safe Operation**

Daniel Fischer; Eva Eberlein

Siemens Healthcare, Erlangen, Germany

**Introduction**

At the RSNA 2012, Siemens introduced MAGNETOM Prisma*, a new 3 Tesla MR imaging (MRI) scanner. This system is designed to fit the needs of those involved primarily in research activities. The foundation of the system is a new 3T magnet, based on the proven MAGNETOM Trio. It provides the same robust base as Trio did – with high homogeneity, only now with zero helium boil-off. MAGNETOM Prisma combines Siemens’ latest technological advancements in MR signal transmit and receive technologies – TimTX TrueShape and Tim 4G technology, with the latter forming the backbone of other Siemens MR scanners 1.5T MAGNETOM Aera and 3T MAGNETOM Skyra. Both technologies have shown an increase in image quality and imaging speed, proving their value in a number of clinical MRI applications [1]. On top of these components, MAGNETOM Prisma includes XR 80/200, a new, high-end gradient system that delivers high gradient amplitudes and fast switching capabilities in a combination that is currently unmatched in the market.

This article is to provide insight into the evolution of the new gradient engine: a brief history of gradients in MR, the technical aspects surrounding the XR gradient engine, safety mechanisms, as well as the benefits that MR applications stand to gain from the outstanding performance.

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*The product is still under development and not commercially available yet. Its future availability cannot be ensured.*
A few words on gradients

The gradient system is one of the major sub-systems of any MR scanner. It plays a key role in the creation of the MR image by providing a means to:

1) spatially select a region that should be imaged and
2) encode the contents of this region. Both selection and encoding are done by applying electric current to the gradient coil, thus creating a field slope on any one of the physical axes x, y, or z. Through precise timing of these additional fields, or gradients, a specific section of an object can be selected and then encoded. By altering the physical characteristics of the selected image region, it can be seen that the gradients have a tremendous influence on the overall quality of the acquired image.

In order to create a gradient pulse in the context of an MR measurement, a lot has to happen, and it all has to happen very fast. The gradient sub-system can be generalized as a chain of several individual units that receive data from their previous neighbor on the chain, process it, and then pass it down the chain to their next neighbor. First in this chain is the measurement control unit, which controls the entire MR pulse sequence and image reconstruction task. This unit has the task of ‘ordering’ not only gradient pulses, but also the necessary RF pulses and performing the reconstruction algorithms on the received MR data.

The ‘ordered’ gradient pulse information is passed onto the next unit, the gradient control module, which works digitally and in sub-millisecond intervals, executing the desired gradient pulse forms in real-time and performing algorithms on these waveforms in order to compensate for effects such as eddy currents, delays, and other gradient-field-induced abnormalities. Next, the small-signal unit processes the digital signal further and converts it into an analog signal, to be fed into the power stage. Here, the signal is amplified and finally fed to the last member of the chain, the gradient coil, in which the rapidly flowing current creates a gradient field in the region of interest.

The two main performance characteristics of any gradient sub-system are the maximum attainable amplitude (Gmax), which is measured in millitesla per meter (mT/m), and the slew rate (SR), which describes how fast a gradient can attain a desired amplitude and is measured in Tesla per meter per second (T/m/s). Ideally, the gradient sub-system is designed in a way to allow the highest amount of performance (Gmax and SR) without having to compromise one measure for the other. It is not uncommon to find clinical scanners with a Gmax anywhere from 33 mT/m to 45 mT/m, while slew rates can vary from 100 T/m/s to 200 T/m/s.

To get an idea as to how gradient performance has evolved over the past 30 years: In 1983 the first super-conducting 0.35T clinical Siemens MAGNETOM system, was commissioned in St. Louis, USA and could deliver a maximum gradient amplitude of 3 mT/m and a slew rate of 3 T/m/s. Ten years later in 1993, the MAGNETOM Vision was introduced and set a new standard in gradient performance for clinical systems with a maximum amplitude of 25 mT/m and a slew rate of 42 T/m/s. Shifting to the new century, MAGNETOM Avanto, the first Tim system, was introduced in 2003 and boasted gradients with a 45 mT/m maximum amplitude and a 200 slew rate. MAGNETOM Prisma raises the gradient performance bar again, delivering maximum gradient amplitude of 80 mT/m and a slew rate of 200. In very few industries has there been such a dramatic enhancement in performance in such a short period of time.

So what are the benefits of maximum amplitude and slew rate? The gradient sub-system has a direct influence on the spatial resolution and acquisition time of the MR image [4]. By increasing the performance of the gradient sub-system, you can increase the spatial resolution and decrease the acquisition time. In terms of MR images, increased gradient performance translates into more signal-to-noise (SNR), fewer distortions, higher in-plane resolution and thinner slices. In short, better image quality. Fast imaging techniques such as echo planar imaging (EPI) and turbo spin echo (TSE) have evolved and thrived in part because of the higher performance that gradient sub-systems have been able to offer.

Diffusion-weighted imaging (DWI) benefits greatly as well because there is a direct relation of the measured ADC data to the square of the diffusion gradient amplitudes. By utilizing higher gradient amplitudes, higher levels of SNR can be achieved, improving DWI and making it even more relevant in the clinical and research environments.

The technical challenge in designing a gradient system is two-fold. On one hand, the desire exists to deliver outstanding performance, which allows the user to realize the previously mentioned imaging benefits. On the other hand, guaranteeing safe operation is paramount. Striking the right balance of performance and safety is the key guiding principle in this process. The characteristics of high gradient performance and safety are largely dependent on the design of the analog components in the gradient sub-system, namely the power amplifier and gradient coil, and play a major role in achieving the desired image quality results. However, the surrounding digital infrastructure also contributes significantly. The following sections give an overview of the XR gradient system in the MAGNETOM Prisma.

The XR gradient system

The XR gradient is a key component of the technological foundation upon which the MAGNETOM Prisma is built. This gradient is capable of driving maximum amplitudes of up to 80 mT/m with a slew rate of 200 T/m/s, on each axis, simultaneously. Let’s take a deeper look at these aspects as they relate to designing the XR gradient system.

The gradient coil can truly be considered a masterpiece of engineering as it brings a number of separate topics together: physics, electro-magnetics, thermodynamics, mechanics, and manufacturing. As the gradient coil is responsible for precisely encoding the physical characteristics within the measurement volume, it is imperative that particular attention be paid to its design. The XR gradient coil itself is constructed of numerous
individual layers of different materials, making the coil substantially thicker and, as a result, much heavier than conventional gradient coils. These layers all have a specific function – whether the conductors to create and adjust the gradient fields, the isolation layers to aid in protection from RF disturbances as well as those potentially induced by the coil itself, or the fill layers to ensure a stable and rigid coil. Additionally, a number of these layers are dedicated water-cooling.

High gradient performance delivers a strong, stable and linear gradient field. A byproduct of all of this current is induced mechanical vibration [3] and thermal noise as well as heat as a result of simple electrical resistance. Through an efficient force compensation design on all axes, these vibrations can be reduced to a minimum. The sheer size of the gradient coil plays to its advantage, whilst the mass of the coil helps reduce vibrations, acoustic noise, and eddy currents further. Heat must still be dealt with as efficiently as possible. Failure to do so can result in a decrease in contrast and an increase in susceptibility artifacts due to a thermally induced drift of the $B_0$ frequency, a common effect with fast imaging techniques such as EPI. With this in mind, the designers of the XR gradient coil utilized a special conductive material that allows thermal energy to be transferred easily to the coolant flowing through these layers in the coil. Each cooling layer is then placed next to a coil conductor, effectively giving each axis its own individual cooling source. In total, the MAGNETOM Prisma has the ability to cool up to 60 kW, thus ensuring that any thermal energy created by the system is also removed from the system, guaranteeing stability and performance at the same time.

Driving the gradient coil is a power amplifier system designed specifically to handle the needs of such a high-performance gradient system. Modular in design and small in footprint, the solid-state amplifier provides 2250 V at maximum voltage for fast switching gradients and up to 900A of peak current for high-amplitude pulses, which this amplifier can achieve simultaneously. It has the total specified power available to it at all times – there is no need to switch to a special hardware mode which would create some compromise with respect to maximum amplitude or slew rate.

It is these specific capabilities that made this amplifier so attractive for the renowned high-performance gradient system supplied by Siemens for the Human Connectome Project (HCP). For this project, two unique MR systems were built, using this amplifier specifically for neurological research. One such system is able to deliver up to 300 mT/m of peak gradient amplitude with a slew rate of 200 T/m/s! While power is important to drive performance, it is just as important to have the ability to accurately reproduce the pulses from scan to scan. Unless the unit controlling the amplifier and thus the gradient coil is deterministic and precise in its functionality, then stable performance of the entire system cannot be guaranteed. The gradient control unit of the XR gradient system consists of a real-time digital signal-processing unit working in sub-millisecond intervals to control the entire operation of the gradient system as requested by the MR sequence. Like many other control units critical to the MR system functionality, the gradient control unit contains a vast amount of intelligence. It has the ability to influence the state of the entire MR

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2 Tim 4G control architecture.

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Safe operation

Unfortunately we cannot continuously increase the amplitudes and slew rates of our gradient pulses and reap their benefits. A rapidly changing magnetic field, which is essentially what MR gradients are meant to produce, has been shown to produce physiological effects such as peripheral nerve stimulation (PNS) in humans [2]. These effects can be uncomfortable. All MR systems with a gradient system that has the potential to stimulate must provide measures in order to ensure that this does not occur. How is it possible to maximize performance of the gradient system while maintaining safe operation under these conditions?

The XR gradient system, like all other Tim 4G scanners, employs several safety mechanisms to ensure safe operation at all times. The accuracy of a constant monitoring system is more important than having lots of alerts, because the user would be notified of this. The XR 80/200 gradient system of the MAGNETOM Prisma employs three main layers of safety.

First — prediction. A so-called ‘look-ahead’, or prediction monitor examines the critical pulses in a measurement protocol and determines whether or not it will be possible to run the protocol without exceeding the stimulation limit. If it is deemed that the protocol will exceed stimulation limits, then the user is presented with a set of possibilities as to how to adjust the protocol, allowing it to run. If the MR protocol checks out OK, then it is allowed to run. Prior to the actual execution of the sequence, the look-ahead monitor sends measurement parameters to the second safety layer — the online monitor.

The online monitor has the task of calculating the current stimulation level and checking this against that calculated by the prediction model. Each calculation occurs at regular sub-millisecond intervals during a running sequence. Should any time during a running sequence the measured levels exceed the predicted levels, the measurement is stopped. Upon completion of a measurement, the prediction model obtains the final stimulation results from the online monitor and does a comparison: does the calculated prediction model stimulation correlate with the actual measured stimulation? By checking the prediction model with the online calculation, there is the added security that the hardware is performing as intended and the user is receiving the maximum performance available. If there were a discrepancy, the user would be notified of this.

The third and final safety layer involves Tim 4G technology. We are not referring to the high-density coils that Tim 4G is known for, but rather the infrastructure that makes it possible to integrate them flawlessly into a complex system.

Through the intense use of fiber optics that enable the DirectRF real-time data transfer, the scanner can digitally transmit large amounts of raw data over dedicated channels from the receivers located directly at the magnet, to the measurement control unit located in the equipment room in a fast and efficient manner. At the same time, other MR components, such as the transmitter or gradient controller, utilize the same fiber optic technology to communicate with each other, at all times, and in real-time. By doing so, the free exchange of information between MR-critical components is possible. Information such as measurement results, calibration data, status updates, and safety information is broadcast over this ‘network’ at all times to all components. Should one component in this network exhibit abnormal behavior, this is registered and the scanner is immediately brought into a safe state. Fiber optics is just part of the digital backbone that makes Tim 4G possible — not just for excellent image quality, but also for robust communication and safe operation of all components at the scanner.

Outlook

The XR 80/200 gradient system of the MAGNETOM Prisma sets a new standard in gradient performance. The design of a unique gradient coil that is easily able to handle the stresses demanded by MR researchers, coupled with a powerful amplifier and an intelligent control system make this the gradient system the perfect centerpiece for a 3T MR research platform. Ensuring safe operation is of the utmost importance, and multiple, redundant layers of safety make this possible. Many MR applications will surely benefit from the gradient power that the XR 80/200 gradient system provides. Adding the technologies TimTX True-Shape, Tim 4G, and a solid 3T magnet to the XR gradients and you get MAGNETOM Prisma — a powerful and flexible tool that will offer MR researchers innumerable possibilities.

References

Introduction

Cryoablation is a promising minimally invasive therapy used to treat malignancies in various organs, and has gained large acceptance in the treatment of prostate and liver cancer [1]. Cryoablation is less painful than high-temperature coagulation techniques such as Laser Induced Thermo Therapy (LITT) or Radio-frequency Ablation (RFA). An additional advantage of cryoablation is the possibility of 'sculpting' the ice-ball using multiple probes, in order to contour complex or large lesions. Magnetic resonance imaging (MRI) guidance of cryoablation allows the monitoring of the ice-ball boundary in three dimensions with excellent contrast between the frozen and unfrozen tissue and potentially allows noninvasive temperature mapping around the ice-ball [2-9].

Previous studies reported MR thermometry inside the ice-ball based on temperature dependent T2* changes during cryoablation. Given the very short T2* values within the ice-ball, short echo time (TE) values need to be used, in the range ≤ 1 ms [2]. Kaye et al. [10] showed exponential dependency of the T2* relaxation time in the temperature range between –20°C to –5°C. This exponential behavior is similar for different tissues (liver, heart, kidney, prostate) [11, 12].

Temperature monitoring in the tissues proximal to the ice-ball would be a valuable safety tool, particularly considering at-risk structures adjacent to the target tissue/organ (e.g. nerve bundles, wall of large vessels, urinary bladder). Currently, temperature monitoring around the ice-ball is performed using invasive temperature probes, which must be placed by the operator, a time consuming procedure prone to puncture complications. To the best of our knowledge, little investigation has been reported on using MR to measure the near-zero temperatures induced around the ice-ball. As long as the tissue contains liquid water, the Proton Resonance Frequency Shift (PRFS)
method should be applicable. PRFS MR thermometry is currently the preferred method for on-line monitoring of thermal therapy, due to its linear calibration and tissue-type independence [13] (excluding the adipose tissue).

Nevertheless, the susceptibility contrast between frozen and unfrozen tissue disturbs the local magnetic field proximal to the ice-ball. This effect is time-dependent as the volume of the ice-ball varies during the procedure and yields temperature estimation errors when using the conventional PRFS method. In particular, the measured values of temperature may be under- or overestimated around the ice-ball depending on the relative orientation from the magnetic field and on the shape of the ice-ball.

This study demonstrates a time-effective 3D correction of gradient-recalled (GRE) phase shift around an ice-ball that is due to the susceptibility contrast between frozen and non-frozen tissue. This correction is necessary in order to deliver accurate temperature maps in tissue proximal to the ice-ball using the PRFS method. The study was published first in Magnetic Resonance Materials in Physics, Biology and Medicine [14].

**Material and methods**

**Theory**

This section summarises the underlying physical model for a time-effective correction of the GRE phase shift induced around the ice-ball during cryotherapy [15]. For more details please see [14]. It must be considered that the magnetic field at one point of an object is the summation of the magnetic contributions of bulk susceptibility (χ) from all other points of the object.

For convenience, the vacuum permittivity is set to 1. According to fundamental physics, if there are no circulating currents within a region of matter, the magnetic field strength (H) in the object can be described as an irrotational field, which means that a magnetic potential (Φ) as a scalar field exists:

\[
\vec{H} = -\nabla \Phi \quad \text{with} \quad \nabla = \left( \frac{\partial}{\partial x}, \frac{\partial}{\partial y}, \frac{\partial}{\partial z} \right)
\]

As known from Maxwell’s equations, the divergence operator of the magnetic induction is always zero. Furthermore, magnetic induction and magnetic field strength are related by \( \vec{B} = (1 + \gamma) \vec{H} \).

Combining this knowledge enables a relation to be established between the scalar potential and the magnetic susceptibility:

\[
\nabla^2 \Phi + (1 + \chi) \nabla^2 \Phi = 0 \tag{2}
\]

This magnetic scalar potential can be expressed as a linear summation:

\[
\Phi = \Phi_0 + \Phi_{in} + \Phi_{obj} \tag{3}
\]

where \( \Phi_0 = -H_0^2 \) represents the ideal field of the magnet (i.e. \( H_0 = (0,0,0) \) is a uniform field), \( \Phi_{in} \) represents the intrinsic inhomogeneities of the magnetic field (including shimming), and \( \Phi_{obj} \) represents the inhomogeneities induced by the object itself.

Neglecting all terms and variables smaller than 10^{-5} and mention that \( \nabla \Phi_{in} = 0 \) (source-free field in the region of interest) equation (2) and (3) can be combined and simplified to:

\[
\nabla^2 \Phi_{obj} = H_0^2 \chi \tag{4}
\]

If the spatial dependence of \( \chi \) is known, the magnetic induction observed by one NMR nucleus which has an electronic screening coefficient \( \sigma \) can be described as [15]:

\[
B_{nuv} = (1 - d + \frac{1}{3} \chi) H_0 + h_{nuv} + h_{obj} \tag{5}
\]

Using equation (5) and the Fourier Transformation (FT) of partial-derivative operators, equation (4) can be written as:

\[
FT(\Phi_{obj}) = -iH_0z FT(\chi) \tag{6}
\]

With \( k^2 = k_x^2 + k_y^2 + k_z^2 \) the squared norm of the inverse space distance, the non-thermal phase shift in the complex GRE image due to the time-dependent susceptibility changes between the time points \( t_0 \) and \( t \) can be written as:

\[
\Delta \Phi_{nuv}(t) \equiv \gamma x H_0 \times T_z \times FT^{-1}
\]

\[
\left[ \left(\frac{1}{3} - \frac{k_z^2}{k^2}\right) FT \chi(t) - \chi(t_0) \right] \tag{7}
\]

using standard symbols (\( \gamma = 42.58 \text{ MHz/T, TE=echo time} \)). Further on the susceptibility contrast between frozen and unfrozen (or defrosted) tissue is defined as a constant value, denoted here \( \Delta \chi_{susc} \), and the shape of the ice-ball as a 3D binary function with \( S(r,t)=1 \) within the ice ball and \( S(r,t)=0 \) outside the ice ball.

Assuming the value of the susceptibility contrast \( \Delta \chi_{susc} \) is known together with the 3D shape function of the ice-ball at each time-point \( S(t) \), the master equation for the correction of the PRFS temperature maps (relative temperature changes) in tissue proximal to the ice-ball is:

\[
\Delta T_{susc}(t) = \Delta T_{PRFS}(t) + \frac{\Delta \chi_{susc} \times FT^{-1}}{[\left(\frac{1}{3} - \frac{k_z^2}{k^2}\right) FT \Delta S(t)]} \tag{8}
\]

where the PRFS temperature coefficient is \( \alpha = -0.0094 \text{ppm/°C} \) [13], the average susceptibility contrast between the ice-ball and the tissue is expressed in ppm and \( \phi(t) \) is the actual phase map acquired at time point \( t \).

**Experimental setup**

Experiments were performed at a 1.5T MR-system (MAGNETOM Avanto, Siemens Healthcare, Erlangen, Germany) on ex vivo pork liver. A home-made MR-compatible cryogenic probe was used in order to demonstrate the performance of the correction method, while using an interventional device [14]. High resolution 3D GRE T1-w images VIBE [16] were acquired as preliminary data (Fig. 1), with the following parameters: TE 1.6 ms, TR 4 ms, TA 2.55 min, FA 10°, BW 650 Hz/pixel and spatial resolution of 1.2 × 1.2 × 1.2 mm³, and were used for positioning a stack of 25 adjacent slices of MR thermometry (a similar GRE sequence was used as above, modified to a rectangular FOV = 75%, slice thickness 2.4 mm, TR 27.6 ms, TE 20 ms, FA25°, BW 230Hz/pixel, GRAPPA 1, temporal resolution 49.6 s and using a single loop coil of 11 cm diameter). The active cryogenic interval corresponded to the circulation of ethanol into the probe, for 20 minutes under MRI monitoring. The MRI acquisition was performed for a further 15 minutes after the peristaltic pump was stopped in order to monitor the initial interval of the return to thermal equilibrium in the tissue.

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A standard baseline correction for the $B_0$ drift was implemented using the GRE phase from a 36 pixel-sized region-of-interest (ROI), selected far from the state transition interface.

**Data processing**

PRFS thermometry data was corrected off line in this study. To correct for the susceptibility-induced GRE phase shift proximal to the ice-ball, the susceptibility difference between frozen and unfrozen/defrosted tissue has to be determined. The underlying physical hypothesis is that the frontier of the ice-ball has the state transition temperature, uniform at every point of the surface. First, the ice-ball is segmented using a semi-automatic region growing algorithm [17]. That is, the experimental estimation of the shape function $S(r,t)$ is obtained for each measurement. The second step is to determine the average susceptibility difference between the ice-ball and proximal tissue ($\Delta \chi_{\text{ice}}$). For this purpose, a cost function (F) was defined as the standard deviation (SD) of the series of corrected temperature values, taken from the ice-ball frontier in the central slice that is defined as the slice of maximum cross section through the ice ball.

The third and last step consisted in applying equation (8) for correcting the PRFS temperature maps using the value $\Delta \chi_{\text{ice}}$ determined above. The main steps of the algorithm are highlighted in Figure 3.

**Results**

No MR-signal was obtained within the ice-ball using a TE of 20 ms at 1.5T and therefore no temperature information was available in that region, whereas the semi-automatic segmentation algorithm always performed robustly. Total computing time for correcting one 3D measurement was less than 1s in Matlab.

The susceptibility contrast between non-frozen and frozen ex vivo tissue was found to be 0.161 ppm for a best-corrected temperature uncertainty of 0.3°C in pork liver (Fig. 2).

The home-made cryogenic probe showed excellent MR compatibility, the size of the signal void region in a plane orthogonal to the copper cylinder was practically equal to the physical diameter of the cylinder, within the available resolution of the T1-weighted MR images. The induced ice-ball at the end of the cryogenic procedure had the shape of an elongated ellipsoid of diameter 1.2 cm and height 4 cm.

Figure 4 illustrates the performance of the temperature correction method for the cryogenic probe setup. Note, after correction, the temperature values in the tissue layer at the frontier of the ice-ball are uniform and correctly measured at around 0°C. In particular, the corrected temperature curves considered at two different locations, defined at the same distance from the cryogenic probe but having different position vectors with respect to the $B_0$ direction, show similar behaviour, as theoretically predicted. The border condition at the margins of the tissue sample was set to 20°C measured from thermometer data acquired immediately before and after the cryogenic procedure. Without correction, the ice-ball frontier temperature was over- or under-estimated by the conventional PRFS method by ±10°C.

**Discussion**

Inducing an ice-ball to treat tumors requires temperature monitoring to protect healthy surrounding tissue, particularly at-risk structures adjacent to the target tissue or organ. Non-freezing cold injuries (NFCI) were documented mainly in the context of accidental casualties. The proposed mechanisms of NFCI include direct axonal damage, ischaemia and ischaemia/reperfusion [18]. Cold immersion (1–2°C) results [18] showed that large myelinated fibres were preferentially damaged, while small myelinated and unmyelinated fibres were relatively spared. The formation of oxygen derived free radicals, which are implicated in ischaemia/reperfusion injuries, have been demonstrated during re-warming of a rabbit hind limb, cooled to 0°C for 20 min [19]. Vasoconstriction, endothelial injury and thromboembolism contribute to vascular insufficiency and ischemia. Thromboembolism, from stasis and endothelial injury, may also be triggered by hemoconcentration and hyperviscosity [20]. The series of enzymatic reactions of the coagulation cascade are strongly inhibited by hypothermia below 28°C [21] that can cause or aggravate bleeding.

The placement of secondary temperature probes into ‘structures at risk’ proximal to the cryo-ablation site is performed routinely. Not only can the placement of these temperature probes be difficult and time consuming, but they can damage the structures they are trying to protect. Temperature mapping outside of the ice ball would allow for
Overview of the experimental results and correction of PRFS thermometry in ex vivo tissue when cryogenic ice-ball was induced by an MR-compatible probe.

(4A) PRFS temperature maps in a coronal mid-plane through the ice-ball: susceptibility-corrected (left) and phase-subtraction conventional PRFS (right).

(4B) Plot of the measured temperatures over time in the three positions visualized in frame a).

(4C) Illustration of the spatial profile of susceptibility-corrected temperature through the centre of the ice-ball (comprising the positions 2 and 3 from above).

3 Workflow of the correction algorithm for the PRFS thermometry in tissue proximal to the ice-ball.

1. Magnitude image of the ice ball
2. Temperatures calculated using phase difference images
   Over- and underestimation of temperature induced by perturbation of magnetic field due to the susceptibility contrast between frozen and defrosted tissue.
3. Border around ice ball.
   → iceball frontier must have a uniform state transition temperature = a uniform value
   Calculation of susceptibility contrast between frozen and defrosted tissue
4. Calculated perturbation field that minimizes the cost function in the model
5. Corrected temperature map

4 Overview of the experimental results and correction of PRFS thermometry in ex vivo tissue when cryogenic ice-ball was induced by an MR-compatible probe. (4A) PRFS temperature maps in a coronal mid-plane through the ice-ball: susceptibility-corrected (left) and phase-subtraction conventional PRFS (right). (4B) Plot of the measured temperatures over time in the three positions visualized in frame a). (4C) Illustration of the spatial profile of susceptibility-corrected temperature through the centre of the ice-ball (comprising the positions 2 and 3 from above).
non-invasive monitoring of these structures and allow the operator to moderate the ice-ball if a precipitous temperature drop is seen. However, using the phase-subtraction only PRFS method for temperature monitoring in the proximity of frozen tissue results in over- or under-estimated values in the typical error range of ± 10°C, depending on the ice-ball size/shape and also on the imaging slice orientation. This range of under- or over-estimation of tissue temperature with conventional PRFS MRT compromises the potential benefit of on-line monitoring of the temperature distribution. To recover accurate temperature maps, our study proposes and validates a numerical algorithm, performed in less than 1 second on a standard CPU, to correct in 3D for the susceptibility-induced GRE phase shift proximal to a cryotherapy ice-ball. Overall, the method described here enables the user to monitor accurately the temperature in the vicinity of an ice-ball, particularly when at-risk structures are adjacent to the target tissue. The method described for the susceptibility artifact correction is based on the fundamental property of isothermal state transition of tissue and allows the PRFS temperature correction without the placement of any probes, which is time consuming and potentially prone to complications.

In contrast to the method described in reference [11] to estimate temperatures inside the ice-ball itself, our aim was to monitor the temperature in regions external and proximal to the ice-ball and therefore to provide a safety tool to enable the physician to avoid collateral damage in surrounding organs or structures of risk.

Finally, the actual correction method does not address motion-related artifacts if any (for instance, concerning liver cryotherapy). Therefore a prerequisite condition here is a motion-robust GRE phase acquisition, prior to applying the susceptibility correction. Coupling motion-correction and susceptibility-correction methods simultaneously is beyond the purpose of the present study and requires further investigations.

Conclusion

This study demonstrates a time-effective stand-alone method for correcting susceptibility artifacts near an ice-ball induced by the susceptibility contrast between frozen and non-frozen tissue. The post processing method examined provides a fast algorithm requiring only one simple user interaction. It allows the non-invasive and accurate monitoring of near zero temperatures in at risk tissues adjacent to the target lesion using the PRFS method.

References

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