Dear MAGNETOM User,

"Although Tim stands for Total imaging matrix, after experiencing the MAGNETOM Symphony with Tim myself I have decided that Tim stands for Total improvement." Dr. Johan Dehem from VZW Regionaal Ziekenhuis Jan Yperman, Belgium is excited about his upgrade to Tim Technology. From head to toe, from routine imaging to advanced applications, from gradient system to RF coils, MAGNETOM Symphony with Tim feels like a totally new system. Adding Tim and top applications such as GRAPPA, SWI (susceptibility-weighted imaging) and REVEAL (diffusion-weighted imaging) MAGNETOM users are able to see more and work faster with their systems.

When thinking about an upgrade the main concerns most likely are:

- Total cost compared to a new system
- New technologies that come with the upgrade
- Clinical advantages of the upgrade
- Total downtime to install the upgrade
- Throughput after the upgrade

This issue of MAGNETOM Flash magazine addresses these topics. We show examples with different business models and different needs from the more than 70 Tim upgrade installations worldwide in less than 1 year. Let us take you to France, Belgium, Australia, and the US for a glimpse into the routine work with MAGNETOM Symphony, A Tim System.

Enjoy this MAGNETOM Flash and the advantages of the Tim Upgrade,

Gustavo Ribeiro
The Editorial Team

We appreciate your comments.
Please contact us at magnetomworld.med@siemens.com
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MAGNETOM Symphony, A Tim System

The upgrade from MAGNETOM Symphony to Tim will bring remarkable clinical advantages to your routine MR applications. The technical comparison shows the improvements ranging from coil technology to gradients and parallel imaging techniques.

### Gradient Strength

- **MAGNETOM Symphony, A Tim System**
- **MAGNETOM Symphony with Ultra gradient system**
- **MAGNETOM Symphony with Sprint gradient system**

### Slew Rate

- **MAGNETOM Symphony, A Tim System**
- **MAGNETOM Symphony with Ultra gradient system**
- **MAGNETOM Symphony with Sprint gradient system**

### Independent Digital RF Channels

- **MAGNETOM Symphony, A Tim System**
- **MAGNETOM Symphony (standard)**

### Coil Elements

- **MAGNETOM Symphony, A Tim System**
- **MAGNETOM Symphony (standard)**

### Max. PAT Factor

- **MAGNETOM Symphony, A Tim System (standard)**
- **MAGNETOM Symphony (standard)**

### New Integrated CP Body Coil

- No-tune transmit / receive coil
- Circularly polarized (CP)
- Optimized RF efficiency and signal-to-noise ratio (SNR)
**MAGNETOM Symphony, A Tim System: Tim Matrix coils**

- **Head Matrix coil**
  - 12 elements, iPAT-compatible (integrated Parallel Acquisition Technique)

- **Head+Neck**
  - 16 elements, iPAT-compatible

- **Whole CNS (head + whole spine)**
  - 40 elements, iPAT-compatible

- **Shoulder**
  - Shoulder array 2 sizes, 4 elements, iPAT-compatible

- **Breast Matrix coil**
  - 4 elements, iPAT-compatible

- **Neck Matrix coil**
  - 4 elements, 16 elements in combination with Head Matrix coil, iPAT-compatible

- **Whole abdomen**
  - 2 x Body Matrix coil + Spine Matrix coil, 24 elements, iPAT-compatible

- **Prostate**
  - Endorectal coil + Body Matrix coil + Spine Matrix coil, 13 elements, iPAT-compatible

- **Knee, ankle**
  - CP extremity coil or 8 channel knee coil, iPAT-compatible

- **Peripheral Angiography**
  - PAA Matrix, 16 elements, can be combined with Body Matrix coil + Spine Matrix coil, iPAT-compatible

- **Spine CTL**
  - Spine Matrix coil, 24 elements, iPAT-compatible

- **Chest, heart, abdomen, pelvis**
  - Body Matrix coil + Spine Matrix coil, 12 elements, iPAT-compatible

- **Wrist**
  - CP Flex small/large or high-res wrist coil, 4 elements, iPAT-compatible

- **Spine CTL**
  - Spine Matrix coil, 24 elements, iPAT-compatible

**MAGNETOM Symphony: Integrated Panoramic Array (IPA) CP array coils**

**Image Processor**

<table>
<thead>
<tr>
<th>MAGNETOM Symphony, A Tim System</th>
<th>MAGNETOM Symphony</th>
</tr>
</thead>
<tbody>
<tr>
<td>Processor</td>
<td>AMD Opteron</td>
</tr>
<tr>
<td></td>
<td>2 CPUs, 2.6 GHz; 8 GB RAM</td>
</tr>
<tr>
<td>Reconstruction speed</td>
<td>1002 recons/sec for 256 x 256 matrix</td>
</tr>
<tr>
<td>Reconstruction speed with RecFoV</td>
<td>8694 recons/sec for 256 x 256 matrix with 25% RecFoV</td>
</tr>
<tr>
<td>Parallel scan and reconstruction</td>
<td>possible with up to 12 data sets</td>
</tr>
<tr>
<td></td>
<td>355 recons/sec for 256 x 256 matrix</td>
</tr>
<tr>
<td></td>
<td>3226 recons/sec for 256 x 256 matrix with 25% RecFoV</td>
</tr>
<tr>
<td></td>
<td>possible with up to 4 data sets in parallel</td>
</tr>
</tbody>
</table>

**Applications**

**MAGNETOM Symphony, A Tim System**

- All advanced applications including diffusion tensor imaging with up to 256 diffusion directions
- Prostate and breast 3D spectroscopy
- mSENSE and GRAPPA
- up to 200 cm whole-body scan
- real-time 3D BOLD imaging
- Coronary Artery Imaging, SWI, REVEAL, SPACE, BEAT, BLADE
- All future applications developed for Tim systems
Cardiac

[Aortic Valve 01]  T1 FLASH cine retro / TR/TE 52.2/2.4
TA 45s / SL 5 mm / slices 25
FoV 320 mm / matrix 384

[Aortic Valve 02]  TrueFISP cine retro, GRAPPA 2 / TR/E 42.4/1.4
TA 8 s / SL 6 mm / slices 20
FoV 280 mm / matrix 192

[Coronary]  T1 3D FLASH FatSat / TR/TE 233.2/2.5 / TA 3:31min
eff. SL 1.5 mm / partitions 18
FoV 340 mm / matrix 768

[DarkBlood]  T2 TSE FatSat Dark Blood
TR/TE 1542.5/66 / TA 7.83 s / SL 5 mm / slice 1
FoV 200 mm / matrix 256

[Late Enhancement]  FLASH IR OS recon image post contrast TR/TE 700/3.5 / TI 310, TA 16 s / SL 8 mm
slice 1 / FoV 370 mm / matrix 256
Courtesy of Klinikum St. Marien, Amberg, Germany
[Grid] T1 FLASH cine
TR/TE 42.7/4.2 / TA 13.6 s
SL 6 mm / slices 15 / FoV 350 mm
matrix 256

[Radial] TrueFISP cine retro radial, GRAPPA 2
TR/TE 39.6/1.3
TA 6.62 s / SL 6 mm / slices 25
FoV 340 mm / matrix 192

[WholeHeart] 3D TrueFISP segmented FatSat, GRAPPA 2
TR/TE 260/1.6 / TA 15:02 min
eff. SL 1 mm / partitions 120
FoV 260 mm / matrix 256

[Flow 01 + Flow 02] T1 FLASH cine retro
TR/TE 46/3.8 / TA 3:18 min / SL 5 mm / slices 30 / FoV 320 mm / matrix 256
Abdomen and Pelvis

[ Abdomen FLASH outphase ]  
T1 FLASH out of phase, GRAPPA 2 / TR/TE 100/2.4  
TA 26.7 s / SL 6 mm / slices 2 x 20 (in and out of phase)  
FoV 350 mm

[ Liver VIBE ]  
T1 3D FLASH FatSat, GRAPPA 2  
TR/TE 5.8/2.8 / TA 22.5 s  
eff. SL 2.5 mm / partitions 60  
FoV 350 mm / matrix 256

[ T1 cor ]  
T1 TSE, GRAPPA 2  
TR/TE 350/16.6 / TA 1:34 min  
SL 6 mm / slices 9 / FoV 380 mm / matrix 320

[ Cor T1 ]  
T1 TurboFLASH, n GRAPPA 2 / TR/TE 5948/7.5/3  
TI 900 / TA 1:14 min / SL 7 mm slices 20 / FoV 400 mm / matrix 512

[ Abdomen T2 cor ]  
T2 TSE, GRAPPA 2 / TR/TE 4288/59  
TA 1:40 min / SL 6 mm / slices 25  
FoV 450 mm / matrix 384

[ Abdomen TIRM ]  
T2 TIRM, GRAPPA 2 / TR/TE 9000/91  
TI 130 / TA 4:40 min / SL 5 mm slices 25 / FoV 480 mm / matrix 448
**Pelvis**

- **T2 TSE, GRAPPA 2**
  - TR/TE: 4640/92 / TA: 3:57 min
  - SL: 4 mm / slices: 25 / FoV: 230 mm
  - Matrix: 384

  Courtesy of H. Lee Moffitt Cancer Institute, Tampa, USA

**Pelvis T1**

- **T1 TSE, GRAPPA 2**
  - TR/TE: 582/14 / TA: 3:07 min
  - SL: 4 mm / slices: 21 / FoV: 400 mm
  - Matrix: 512

**Pelvis cor**

- **T2 TSE**
  - TR/TE: 5800/107 / TA: 2:92 min
  - SL: 3 mm / slices: 21 / FoV: 200 mm
  - Matrix: 512

**Pelvis T2 sag**

- **T2 TSE**
  - TR/TE: 582/14 / TA: 3:07 min
  - SL: 4 mm / slices: 25 / FoV: 380 mm
  - Matrix: 448

  Courtesy of Klinikum St. Marien, Amberg, Germany

**Pelvis**

- **T2 TSE**
  - TR/TE: 5900/107 / TA: 2:04 min
  - SL: 3 mm / slices: 21 / FoV: 200 mm
  - Matrix: 512

**Pelvis T2 sag**

- **T2 TSE**
  - TR/TE: 582/14 / TA: 3:07 min
  - SL: 4 mm / slices: 21 / FoV: 380 mm
  - Matrix: 448

  Courtesy of H. Lee Moffitt Cancer Institute, Tampa, USA

**Pelvis TIRM**

- **T2 TIRM, GRAPPA 2**
  - TR/TE: 6000/118 / Ti: 150 / TA: 4:07 min / SL: 3 mm
  - Slices: 15 / FoV: 240 mm
  - Matrix: 320
Abdomen and Pelvis

**[Pelvis transversal]**  
**T2 TSE**  
TR/TE 6360/104 / TA 5:18 min  
SL 3 mm / slices 30 / FoV 160 mm  
matrix 512

**[Myelography]**  
**T2 3D HASTE**  
TR/TE 8000/273 / TA 5:48 min  
eff. SL 0.9 mm / partitions 44  
FoV 230 mm / matrix 256

Courtesy of H. Lee Moffitt Cancer Institute, Tampa, USA

**[MRCP]**  
**T2 3D TSE Restore, GRAPPA 2**  
TR/TE 3773.6/678  
TA 3:43 min / eff. SL 1.5 mm  
partitions 40 / FoV 380 mm  
matrix 384

Courtesy of Klinikum St. Marien, Amberg, Germany

**[T2 TSE]**  
**T2 TSE, GRAPPA 2**  
TR/TE 6734.8/81 / TA 3 x 3:07 min  
SL 4 mm / slices 30 / FoV 360 mm  
matrix 512

**[T2 TSE FatSat]**  
**T2 TSE FatSat, GRAPPA 2**  
TR/TE 5898.9/72 / TA 3 x 2:49 min  
SL 4 mm / slices 30 / FoV 360 mm  
matrix 512

**[TrueFISP]**  
**T2 TrueFISP, GRAPPA 2**  
TR/TE 4500/43 / TA 13 s / SL 4 mm / slices 9  
FoV 370 mm / matrix 384
WholeBody T2
T2 TSE, GRAPPA 2, 6 steps
TR/TE 5430/78 / TA 6 x 1:43 min
SL 5 mm / slices 21 / FoV 1826 mm
matrix 512

WholeBody TIRM
T2 TIRM, GRAPPA 2, 6 steps
TR/TE 9000/85 / TI 130
TA 2 x 2:15 min / SL 5 mm
slices 15 / FoV 1881 mm
matrix 384

WholeBody TIRM cor
T2 TIRM, GRAPPA 2, 6 steps
TR/TE 2670/101 / TI 150
TA 6 x 2:58 min / SL 7 mm
slices 18 / FoV 1876 mm
matrix 512

Courtesy of Klinikum St. Marien, Amberg, Germany
First Experiences with Tim Upgrades in France

Eric de Kerviler, M.D.¹, Bruno Boyer, M.D.²

¹Imaging Department, APHP Saint Louis Hospital, Paris, France
²Radiology Department, Begin Military Hospital, Paris, France

Introduction
Since the introduction of the MAGNETOM Avanto, the first Tim system, we have frequently been told: “Tim is extremely interesting; is it possible to have this functionality with my Symphony?”
The answer to the question is “yes”. In France, the Saint Louis Hospital’s and the Begin Military Hospital’s MAGNETOM Symphonies were both upgraded to Tim (Total imaging matrix) in November 2005.
The Saint Louis Hospital has been working with a MAGNETOM Symphony since 1999. The activities of the department are diverse, with a strong orientation toward oncology and abdominal imaging. Caring for fragile patients requires working fast and being able to rapidly examine various regions of the anatomy. In 2002, the system had already benefited from the Maestro Class Quantum iPAT gold upgrade to make use of multiple element coils and Parallel Acquisition Techniques, and as such improve system performances for key applications of the department.
Since 1999, Begin Military Hospital has been working with a MAGNETOM Symphony. Activities are general in nature. Despite being a military hospital, the department is open to the public. The system has been upgraded on a regular basis.
After 6 months’ experience with the MAGNETOM Symphony Tim upgrades, we would like to present a report of the experiences of these two hospitals.

The French context
Whilst MRI activity has greatly advanced these last few years, the number of MRI systems per inhabitant in France remains small. Moreover, the renewal of systems in a hospital setting is taking longer and longer. Consequently, it is important for these teams to keep their systems maintained and to operate them at peak levels. The upgrade is a very appropriate response.

Why did the hospitals decide to upgrade their MAGNETOM Symphony?
For Saint Louis Hospital, Tim technology seemed to be the tool to enable them to make huge progress by breaking through the present restrictions in gastrointestinal and functional applications and to develop extensive imaging, thanks to signal gain and Parallel Acquisition Techniques. The upgrade
Patient follow-up in a liver pathology case

**[Figure 2]** Free-breathing TSE T2-weighted sequence with fat saturation using PACE and the Body Matrix coil.

<table>
<thead>
<tr>
<th>parameter</th>
<th>value</th>
</tr>
</thead>
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<tr>
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<tr>
<td>SL</td>
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<tr>
<td>slices</td>
<td>20</td>
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<tr>
<td>TA 2*</td>
<td>53.93</td>
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**[Figure 3]** FLASH 2D IN OPP sequence.

<table>
<thead>
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<th>value</th>
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</thead>
<tbody>
<tr>
<td>PAT</td>
<td>2</td>
</tr>
<tr>
<td>matrix</td>
<td>320</td>
</tr>
<tr>
<td>SL</td>
<td>7</td>
</tr>
<tr>
<td>slices</td>
<td>40</td>
</tr>
<tr>
<td>TA 2*</td>
<td>12.55</td>
</tr>
</tbody>
</table>

**[Figure 4]** 3D VIBE

<table>
<thead>
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<th>parameter</th>
<th>value</th>
</tr>
</thead>
<tbody>
<tr>
<td>PAT</td>
<td>2</td>
</tr>
<tr>
<td>matrix</td>
<td>384</td>
</tr>
<tr>
<td>SL</td>
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<tr>
<td>slices</td>
<td>60</td>
</tr>
<tr>
<td>TA</td>
<td>16.48</td>
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</tbody>
</table>
would also reduce lead times thanks to increased throughput. Begin Hospital considered Tim technology particularly suited to develop specific female pelvic exams because of the developments in 3D imaging. This upgrade also allows the hospitals to incorporate new 3D applications such as syngo SPACE and syngo SWI (Susceptibility-Weighted Imaging).

**Experience after 6 months**

**Practical aspects:**
The departments were already familiar with MAGNETOM Symphony’s IPA (Integrated Panoramic Array) system that allows the combination of several coils in a phased array. With Tim, however, extended coil combinations are available with the 10 sockets of the examination table and thereby make the use of coils even easier. There are now new possibilities to combine more coils with the new Matrix technology that offers a higher number of channels. As a further benefit, the Tim “Matrix” coils are lighter. Total integration of the coils has really changed lives. Now, it is sufficient to add the Body Matrix for abdominal-pelvic exams. There is no compromise between quality and coverage. The sites gained in utilization flexibility as well. For instance, the use of 2 Body Matrix coils side-by-side (left-right) enables straight-forward handling of heavy patients with a large waist size.

In cardiology, the new Bluetooth system (wireless ECG leads) is very easy to use: no wires, great reliability, optimized reproducibility of the exams. The interface is even clearer, thanks to Tim. The position of the anterior coils, for example, is precisely identified and viewed on the console. The Phoenix tool – which by simple drag and drop reproduces all the parameters of a previously acquired image – enables very quick implementation of new sequences used on the MAGNETOM Avanto.

**Clinical aspects:**
Tim promised signal gain. Users were able to observe this regardless of the type of application, thereby increasing resolution while reducing acquisition times.

On the one hand, the combination of the Tim Head and Neck Matrix Coils, by offering more elements (16) and thus, signal, enables coverage up to the cervical vertebra, or even an analysis extended to the complete spine when required by the pathology.

On the other hand, the 3D sequences such as the SPACE sequences which are very useful for diagnosis – are much easier to use. “We cannot do without them any longer” says

**Renal transplant evaluation at day one:**

**Figure 5** 3D VIBE isotropic

<table>
<thead>
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<th>PAT</th>
<th>2</th>
</tr>
</thead>
<tbody>
<tr>
<td>matrix</td>
<td>320</td>
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<tr>
<td>FoV</td>
<td>247*380</td>
</tr>
<tr>
<td>SL</td>
<td>1.6</td>
</tr>
<tr>
<td>slices</td>
<td>120</td>
</tr>
<tr>
<td>TA</td>
<td>26.03</td>
</tr>
</tbody>
</table>

*MPR isotropic VIBE allows an optimal assessment of vessels and renal transplants.*
Professor Boyer. It seems therefore to be a win-win situation. For abdominal exploration, Saint Louis Hospital has been using 12 element anterior and posterior body coils; Tim has ensured equivalent element density and improved signal. This has permitted development of the use of isotropic sequences such as the 3D VIBE sequence (Volume interpolated breath-hold examination). Thanks to a better fat saturation and to PACE (for free-breathing abdominal examinations), the T2-weighted fat saturated sequences are of excellent quality.

**Conclusion**

These first results of MAGNETOM Symphony Tim upgrades have been very positive and the users are very satisfied. Quoting Professor Eric de Kerviler: “We truly had the impression that we were working with a completely new system. It would be difficult to go back now.”

The MAGNETOM Symphony, A Tim System holds true to its promises in terms of ergonomics and image quality. The users of the Symphony Tim are therefore eagerly anticipating the arrival of new applications in the future.

### Feedback from Saint Louis Hospital

“A few months have passed since we upgraded our MAGNETOM Symphony to Tim. The first step after the upgrade consisted of using the same imaging protocols as before. We noticed an immediate and dramatic improvement in image quality due to the new Matrix coils, using a larger number of elements. The second step was to speed up the sequences or increase the spatial resolution without having to trade in image quality. This was easy and most of our examination protocols have now been redesigned. As a result, we obtain better images faster. The last step was to take advantage of the coil matrix. This has been a major change in the way we work. Adding a coil in the field of view is now performed with a single mouse click. The system, which allowed us to plug in a large number of coils at the beginning of the examination, provides us now with the possibility of combining coils. This prevents signal loss at the margins of the field of view and offers us great flexibility for studies requiring table offset/movement. After 6 months, our feedback of the MAGNETOM Symphony Tim upgrade is very positive.”

*Prof. Eric de Kerviler*

Scanner model used:
- Saint Louis Hospital: MAGNETOM Symphony, A Tim System
- Begin Military Hospital: MAGNETOM Symphony, A Tim System
Image Quality Improvements after MAGNETOM Symphony Tim Upgrade

J. Dehem, M.D.
Jan Yperman Ziekenhuis, Ieper, Belgium

Our MAGNETOM Symphony, installed since 2000, was upgraded to a MAGNETOM Symphony, A Tim System in February 2006. With the Tim (Total imaging matrix) platform implementation, we have raised our 6-year-old system to the level of the recent generation of MR-systems, offering up-to-date technology and service to our patients. Immediately, three major improvements become clear:

1. Image quality improved: comparing data of the same patients, taken before and after the upgrade with the same parameter setting, shows a more homogeneous image and an improved fat saturation (fatsat) due to the renewed body RF coil and completely new RF system.

2. Patient positioning is easier and patient comfort has improved: head, neck, spine and body array coils were exchanged for Matrix coils, containing multiple “clusters” of 3 coil elements. Several plug-ins on the table allow multiple coil combinations and allow to position the patient in head-first or feet-first direction. These coil combinations are indeed ‘seamless’ and have major advantages in orthopedic and angiographic imaging as illustrated in the following chapters. The Body Matrix coil is especially appreciated by the patient, because of its light weight. It is also easier to fix the coil around the patient, improving patient comfort.

3. Multiple coil combinations allow extensive use of iPAT (integrated Parallel Acquisition Technique): As the patient is covered by multiple coil elements, iPAT is used in all 3 directions. In 3D imaging, combination of iPAT in phase and slice direction is possible (iPAT3). We routinely use higher PAT factors (PAT3) and combined iPAT imaging (iPAT2 x PAT2 = PAT4). As we do not have the Tim Whole Body Suite, we only comment on ‘local’ exams in different clinical applications.

1. Head and Head/Neck imaging

In our experience, overall image quality has improved compared to the former CP head coil. Introduction of Tim meant introduction of iPAT in all 3 directions in neuro head applications. The major improvement we see in the use of iPAT in
head imaging is in the use of EPI (echo planar imaging) sequences (diffusion, perfusion, EPI_hemo sequence) which decreases distortion. iPAT is extensively used in our 3D TOF (Time of Flight) to obtain high resolution and high coverage in a reasonable time.

Tim implementation will allow us to follow the most recent MR techniques (BLADE, SWI (susceptibility-weighted imaging), SPACE) available on the last generation machines, as our software platform is common.

The high Signal-to-Noise 12-element Head Matrix coil gives good quality images for TMJ (Temporo-Mandibular Joint) imaging.

The seamless combination with the Neck Matrix coil and the good fatsat in this region are welcomed for head and neck, spine and plexus imaging.
2. Spine imaging

We routinely used iPAT on MAGNETOM Symphony for all spine imaging. Tim allows additional use of iPAT in the coronal direction with Left-Right (LR) phase encoding, thanks to the new 24-element Spine Matrix coil with LR orientation of the clusters. An iPAT factor of 3 can be obtained.

We routinely use a combination of Spine Matrix with the Body Matrix coil (thoraco-lumbar spine) which allows us to use iPAT in transverse orientation, too. In C-spine using the Neck Matrix coil, iPAT is always activated in all orientations. High S/N allows us to use iPAT routinely in all spine applications.

3. Musculoskeletal applications

In this section we want to focus on the clinical applications with the standard Tim upgrade coils and the improvements we see. Since we had acquired the 8-channel Invivo shoulder and wrist coil, we continue to use these coils on MAGNETOM Symphony, A Tim System, but these are not further discussed in this report.

The former extremity coil is re-used on MAGNETOM Symphony, A Tim System, but needs an additional connector to plug into the new table. We routinely use the Invivo knee coil for knee imaging as it offers iPAT capabilities. The quality enhancements pre-post Tim that we comment on here with the Invivo knee coil are similar to what we see with the CP extremity coil or the other Invivo coils. These are:
1. Larger range of RF and improved RF-homogeneity.
2. Better anatomical delineation.
3. Improved fat saturation.

The CP extremity coil is used for very large, fatty knees and for ankle examinations. We admit that we also often use the Head Matrix coil for upper and lower extremities e.g. foot, toe, hand and finger, which can successfully be imaged in the Head Matrix coil. No rebuilding of the patient table is needed, patient positioning is quick and easy, and the coil allows use of iPAT, increasing patient throughput.

For hip imaging we upgraded from a 6 element spine array coil to a 3 x 8 element Spine Matrix coil. This allows:
1. A larger RF range in the hip and upper leg region for tall patients, registered head first.
2. Combination with the Body Matrix coil in all directions, especially in the coronal plane, often used in the hip. We use routinely a PAT factor 3 in LR encoding. For higher PAT factors in coronal orientations (often large field of view (FoV), consequently high S/N), gradients can be swapped to HF phase encoding and PAT4 can be used.
An inventive way to use the Tim platform is to use the Spine and Body Matrix for elbow imaging. There are several benefits:

1. Fast patient positioning.
2. Patient comfort.
3. Use of iPAT in all directions, reducing scan time. Total scan time is reduced from 20 to 13 minutes.
4. Offering better image quality than with the use of flex coils, where iPAT can not be used.

[Figure 6] Consistent image quality in lumbar spine with extensive use of iPAT in all directions – even in coronal plane L-R encoding, PAT3 – with clear, sharp images.

[Figure 7] Comparison of the same patient before and after the Tim upgrade with the Invivo knee coil. We observe here an extension of the RF range, improved fat saturation and high anatomical detail.

[Figure 8A] Comparison of the same patient before and after the Tim upgrade. Note the greater signal in the peripheral zone due to more spine elements.

[Figure 8B] Higher signal, same high resolution and less time, thanks to iPAT in coronal direction.
4. Breast imaging

In MR-mammography, Tim allows more combinations of coils. We always removed the spine coil in breast examinations: using our MAGNETOM Symphony it was impossible to combine the body phased array with the breast coil, as the plug for the body part was on the spine array coil. On MAGNETOM Symphony, A Tim System, however, both Breast and Body Matrix coil can be plugged in simultaneously which is a great advantage. This improves the S/N in both breasts and even more important, in the axillae. With this large S/N, we optimized our dynamic protocol to a spatial resolution of (0.9 x 0.9 x 1.1 mm\(^3\)) within the same time resolution (1 min). Fatsat has improved. We look deep into the pectoral and axillary region.

5. Abdominal and pelvic applications

We see the following improvements in abdominal and pelvic imaging:

1. The Body Matrix coil offers more comfort due to its light weight and it fits well around the patient.
2. S/N increase allows us to use thinner slices (5 mm in 2D, 2 mm in 3D VIBE (Volume Interpolated Breathhold Examination)) and/or higher resolution scans (use of 320 resolution instead of 256).
3. Multiple plugs allow the imaging of body or pelvis in head first or feet first position.
4. PAT2 is routinely used for almost all exams, mostly to shorten breathholds.
5. Very homogeneous fat saturation is obtained in abdomen and pelvis.

[Figure 9] Patient with tendinosis. Use of combination of Body and Spine Matrix coils. Easy patient positioning. Total imaging time 13 minutes. This protocol includes a 3D DESS (Dual Echo Steady State) high resolution scan (0.7 x 0.6 x 0.8 mm\(^3\)) of 3 minutes. iPAT is used in all 3 dimensions.

[Figure 10] Comparison of the same patient before (upper left) and after (upper right) the Tim upgrade in a fatsat T2-weighted high-resolution scan. Better delineation of the lesion with MAGNETOM Symphony, A Tim System. Lower segment depicts thin MIP reconstructions of the subtraction dataset on the axillary lymph nodes on MAGNETOM Symphony, A Tim System. Note the visualization of the mediastinum and axillae when both Breast and Body Matrix coil are combined.
[Figure 11] Pre- and post-gadolinium 3D VIBE on MAGNETOM Symphony, A Tim System. Our standard protocol in transverse orientation is a 96 partition scan of 25 s with a resolution of (1.4 x 1.3 x 2.0 mm³). Late enhanced coronal scan with 72 partitions in 22 s with a resolution of (2.0 x 1.4 x 2.0 mm³). PAT2 is used in transverse and coronal orientation.

[Figure 12] T2_tse_FS in the prostate in 2 different patients. Upper segments (pre-Tim): coronal plane with resolution of 1.0 x 1.0 x 4.0 mm³ in 2 minutes (left), transverse plane in 0.8 x 0.8 x 4.0 mm³ in 3 minutes (right) (no. of slices = 14, TR = 3530 ms). Lower segments show the quality of MAGNETOM Symphony, A Tim System: coronal plane with resolution of 1.0 x 1.0 x 4.0 mm³ in 2 minutes (left), transverse plane in resolution 0.8 x 0.8 x 4.0 mm³ in 4 minutes (no. of slices = 25, TR = 6230 ms).

[Figure 13] Homogeneous signal from aorta to ICV (intracranial vessels) through use of a seamless combination of Head, Neck and Spine Matrix coil. We use 176 partitions of 1.1 mm, resulting in coverage of the complete anatomy in A-P direction. Resolution of (0.8 x 0.7 x 0.8 mm³) in 38 s. High PAT factor (PAT2 x PAT2) in phase encoding and slice direction are used to keep acquisition time short.

[Figure 14] Large coverage from the subclavian to iliac vessels through a seamless combination of 3 spine clusters, Neck and Body Matrix coil. We use 176 partitions of 1.1 mm (1.1 x 0.9 x 1.1 mm³) in 24 s. High PAT factor (PAT2 x PAT2) in phase encoding and slice direction.
[Figure 15] Coverage from above the carotid bifurcation, the subclavia and thoracic aorta until renals through combined use of Head, Neck, 2 spine clusters and Body Matrix coil. We use 192 partitions with an isotropic resolution of 1 mm (1.0 x 1.0 x 1.0 mm³) in 24 s. High PAT factor (PAT2 x PAT2) in phase encoding and slice direction. Note the better visualization of the peripheral regions, which is improved by the new body send coil.

[Figure 16] In renals, a large coverage is less important. Small pixel size is important here, to reduce dephasing as much as possible. We use a combination of Body Matrix coil with 3 spine clusters. We use 96 partitions with a resolution of 1.1 x 1.0 x 1.1 mm³ in 27 s, PAT3.

[Figure 17] Aorto-iliacal 64 partitions of 1.2 x 1.1 x 1.8 mm³ in 23 s, PAT3 Iliaco-femoral 56 partitions of 1.2 x 1.1 x 1.8 mm³ in 19 s, PAT3. Lower legs 56 partitions of 1.2 x 1.1 x 1.3 mm³ in 16 s, PAT2.
6. Contrast enhanced angiography

In angiographic applications, the major Tim improvements are:

1. Extensive use of iPAT (PAT3, PAT2) which allows higher resolution and/or larger coverage for the same acquisition time.
2. Use of maximum FoV with high S/N with seamless combination of coils.

Use of both allows us to obtain a very high resolution of our angiographies in an acceptable breathhold time. High concentration Gadolinium compensates for the loss of S/N in these high resolution scans with high PAT factors.

In peripheral angiography, we use a robust technique covering the extremities from renals to toe. The inguinal region is clearly depicted through the use of the Body Matrix coil, in combination with the spine elements and the Peripheral Angiography Matrix coil. Following the Tim upgrade, which extended our receiver circuit with 18 independent receiving channels, we are routinely applying iPAT in all ranges. We obtain high resolution and sufficient coverage in a short time and, consequently, a better delineation of the vessel.

The renal arteries are always included in the picture and can give additional information to the clinician.

In a creative way, the Head, Neck, Spine and Body Matrix coil can be combined to depict the vessels of both left and right lower arm and hand. Seamless combination of all coils elements gives a homogeneous overview.

This image quality and resolution makes magnification possible, with still sufficient diagnostic quality.

Conclusion

Tim (Total imaging matrix) has proven to be a Total improvement. The new RF body send coil and completely new RF system in combination with the new matrix coils offer a general improvement.

Tim allows higher quality (higher resolution, more S/N) and more patient comfort (lightweight coils, easy positioning and shorter breathholds). The overall scan time is diminished, resulting in a higher patient throughput and shorter waiting lists. Tim brings your system in line with the latest MR technology.

„Although Siemens launched Tim as the Total imaging matrix, Tim means to me Total improvement.“

J. Dehem, M.D., Jan Yperman Ziekenhuis, Ieper, Belgium
In May 2006, Westmead private Hospital became the first radiology facility in Australia to undertake a MAGNETOM Symphony Tim Upgrade. We asked Dr. David Ho, lead radiologist, Mrs. Francis Gray, lead MR technologist, and Ms. Kumaransi Silva, MR technologist, to comment on the difference this technology has made.

Ergonomics
The technologists are very excited about the coil concept provided with Tim. The ability to have multiple coil elements plugged in at once and to be able to use various flexible combinations of these coils simultaneously has had a major impact on both workflow and patient comfort. No longer is it necessary to re-enter the scan room and reposition the patient or coils when scanning multiple body regions. The coil set up is easy and quickly adaptable for patients with various different body habitus. The patients love the light weight nature of the matrix coils and the unique modular nature means that anatomical and pathological coverage is no longer limited by individual coil design. The "Intelligent Coil Control" provides a quick and easy way to visualize the "active coil elements" on the monitor, thereby helping to optimize image quality and saving sequence planning time. As the practice gets busy these factors become more important and assist in reducing the pressure on staff.

Image quality
Having used a MAGNETOM Avanto previously, Dr. Ho has been very impressed with the image quality on his MAGNETOM Symphony, A Tim System. He has found the fat suppression to be very uniform, even in hard to fat suppress areas such as the neck. The referrers have also been impressed by the carotid angiographies, the higher resolution now possible with the Tim Matrix coils playing a major part in this. The Tim open architecture has created the opportunity to use a wide range of third party coils: e.g. specific musculoskeletal coils, resulting in beautiful orthopedic images. Signal to noise has noticeably increased with the Tim Matrix coils, with examinations such as the head and angiographies being noticeably better. Larger matrixes and more slices, are now able to be used without extending the examination time, by using iPAT (integrated Parallel Acquisition Technique). The Body Matrix coils give greater clarity and coverage. If we see something peripherally it is easier to zoom in and examine the pathology in more detail.

Referring physicians and patients
As a private provider this site is in competition with other practices. The referring doctors may not understand technology, but they want their patients to have the best leading edge technology.

Of course our competitors used the fact that we had a 5 year old MR system against us. With the Tim Upgrade, referring doctors now know that we do have leading edge technology.
Not only have they noticed an improvement in image quality, but have fallen in love with other aspects of the new technology, such as the composed spine images, which provide an additional visualization tool to standard image assessment. The surgeons also love these composite image representations, and use them in conjunction with the individual source images.

Trigeminal studies are also improved; 3D volumes have better flow compensation. The referrers know that for their patients there is greater comfort and faster examinations thanks to the better workflow attributable to Tim. Not having to change coils greatly affects patient comfort and some referrers have told us, that they were waiting until we had the Tim Upgrade before they would refer some cases.

We have found the Tim technology to be very useful in other ways, for example when there is a lesion at the Cervico-thoracic junction, I do not have to ask for the patient to be repositioned, I can just zoom in to get the detail. It is also a fact that the speed of the examination as a whole makes patient movement less likely and consequently we see less movement in the images.

Financial outcomes and productivity
After the Tim Upgrade, we have seen a 30% increase in the number of examinations performed compared to before. It should also be noted that the upgrade has reduced waiting times. The financial impact is easy for us to see, this is due to an increase in patient volume which also translates to an improvement in cash flow.

Staff retention and development
There is a significant cost (both financial and clinical) associated with the loss and re-training of professional staff. It has been easy to attract staff and train them with this technology. There are fewer mistakes made and if there are any, they are much easier to recover from. In this environment, high levels of staff retention within radiology is very important.

Future developments
We believe that we will see other MR applications increasing in our practice now, for example an increase in the use of MR Angiography.

Conclusion
The initial business case developed in support of the Tim Upgrade for MAGNETOM Symphony has been proved so far, looking at the patient activity and financial results. Over the next year, we plan to introduce new techniques and expand the potential use of this technology. The Tim Upgrade has met and in many cases exceeded our high clinical and financial expectations.
The MR story at Methodist Hospital
In 2001 we performed 7,270 MR procedures on an antiquated Siemens MAGNETOM SP 63 and a GE 5X. It was the consensus of the Radiologists and Administration that both systems be replaced with new technology to provide superior MR imaging. The MAGNETOM Symphony and MAGNETOM Sonata systems were chosen due to superior body, vascular, neuro and new cardiac capabilities.

The growth we experienced was unprecedented in our imaging history. In 2002 we performed 11,307 procedures, a 55.5 % increase, 51.9 % over projected growth. The majority of the growth was seen in body, vascular, neuro and cardiac. In 2003 we preformed 11,745 procedures and began functional pre/post surgical planning. In 2004 we preformed 12,811 procedures upgrading both MR systems to Maestro Class. In 2005 we performed 13,167 procedures (Fig. 1).

A system utilization report was generated and showed that a 99 % and 94 % capacity level had been achieved. It was felt that we had reached saturation, and that we could not provide the level of service that was expected by the clinicians and community. Siemens MR technology had advanced, and for Methodist Hospital to continue providing cutting edge imaging, the MAGNETOM Symphony Tim Upgrade was chosen.

Why was the MAGNETOM Symphony upgrade chosen?
The MRI department is two floors below ground level. It is very expensive to install or move a large piece of equipment. We did not want a long interruption in services. Therefore we felt that the upgrade would bring us to the next level of imaging without replacing an existing piece of equipment. The Tim platform provided the advanced template that could be upgraded in the future as technology advances. Additional efficiency is what the MAGNETOM Symphony Tim upgrade added to our daily operations. The ease of moving from one body location to the next without repositioning patients or coils increased our patient throughput. We met our projected goal of 2 additional procedures a day in April 2006. In June 2006 we increased it to 3 procedures a day. We are on track to meet our projected goal of 4 procedures a day by August 2006. The projected 2006 procedure increase due to Tim is 618 –738 exams.

Benefits of the MAGNETOM Symphony Tim upgrade
The new Tim Spine Matrix coil is larger and provides additional coverage, this gives technicians flexibility in positioning patients. The Tim Spine Matrix coil (Fig. 2A) also has 24 coil elements compared to the 6 coil elements on the MAGNETOM Symphony Maestro Class systems. This allows for easier positioning of the patients head or feet first.
The Tim Body Matrix coil is one of the advances the technicians commented about right away. The coil covers a larger area, has greater flexibility, built-in straps and a longer cable for easier positioning (Fig. 2B).
The 12-channel Head Matrix coil can reduce imaging time by 20 – 30 % with minimal decrease in quality. A 10 % increase in signal to noise across the board when switching from CP mode to 12 channels with no noticeable shading.
**Coil set up**

The coil locations are displayed anatomically on the display monitor, including the anterior coil location; this is not available on Maestro Class systems. The automatic coil selection indicates visually which coils are turned on. This reduces repeated scans by technicians who forgot to turn the anterior coils on. This also allows the technicians to see anterior coil placement and make adjustments if needed (Fig. 3A, 3B).

**Maestro Class vs. Tim**

Wireless Gating (Fig. 2C) reduces the patient positioning time on the table. Less set up time equals greater efficiency. Specific electrodes are needed to get proper contact. We found that the package that came with the upgrade worked the best. We purchased both types of electrodes from Siemens accessories.

**Imaging benefits**

**Runoffs**

Pre-upgrade Runoffs with coil setup and patient positioning were 60 minutes +/- . With MAGNETOM Symphony, A Tim System imaging time dropped to 35 minutes +/- with higher resolution on VIBE (Volume Interpolated Breathhold Examination) sequences and angiographies. We were able to increase the matrix on the VIBEs to 320 and to 384 on the angiographies.

**Body**

All Body work, including MRCP (MR cholangiopancratography), renals and livers have a new HASTE (Half-Fourier Acquisition Single-Shot Turbo Spin Echo) sequence that is much sharper, with no time increase.

**Spine, C, T & L**

We were able to decrease our imaging time by 20 % on sagittal spine sequences and increase the resolution. We saw no change with axial spine imaging.

**Neck**

By utilizing iPAT (integrated Parallel Acquisition Technique) we were able to increase the number of slices by 30 % while maintaining the same time.

**Heart**

The Dark Blood HASTE sequence provides sharper images.

**MPRAGE**

We were able to reduce the time by 50 % when using iPAT. We choose to do this on difficult patients.
Gadolinium-Bolus Carotids, Renals and Aorta
We had increased signal with coronal images while still maintaining the time using iPAT. We increased the aorta resolution from a 256 matrix to 384.

Radiologists’ Comments

Neuro
Increased S/N utilizing the 12-channel Head Matrix coil in the 12-channel mode versus the CPU mode.
Image quality of MR Angiography (MRA) have increased with the 12-channel Head Matrix coil and the 8-channel Neck Matrix coil.

Body & Cardiac
MRCPs are much sharper, greater coverage with the Tim Body Matrix coil.
Cardiac quality of the images on the MAGNETOM Symphony, A Tim System is equal to the 8-channel MAGNETOM Sonata Maestro Class scanner.

Upgrade time frame & Applications
The upgrade started on November 28th and was available for applications on December 15th, i.e. within 17 days. We were able to continue with only one scanner by reducing some of our outpatient scheduling slots during the week and increasing weekend availability.
Application training began on Thursday, December 15th and ended on Tuesday, December 20th.

The time it took for staff, 6 technicians, to feel comfortable and take advantage of the new capabilities consistently was on January 9, i.e. within 19 calendar days.

MAGNETOM Symphony, A Tim System, July, 2006
To date we are very happy with our decision to upgrade the MAGNETOM Symphony to Tim. We are still able to utilize the MRI Devices coils that we have invested in over the years. The technicians favor the MAGNETOM Symphony, A Tim System over the MAGNETOM Sonata, especially when multiple areas need to be imaged. This upgrade has extended the life of the MAGNETOM Symphony system (Fig. 4, 5).
Purchasing and installing the upgrade prior to the installation of our third scanner in June 2006 allowed for accelerated training on the new 3T MAGNETOM Trio, A Tim System.

Looking to the future
We would consider a similar upgrade for the MAGNETOM Sonata if Siemens moves in that direction.

Editor’s remark:
An upgrade of MAGNETOM Sonata to MAGNETOM Symphony, A Tim System is now available.

*Results may vary. Data on file.
Tim Trio Upgrade Benefits

Jeff Zimmers RT, (R, MR) ARRT¹, Karen Ziadie, RT (R) ARRT²

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In the recent months there has truly been a revolution for Siemens 3T users. 3T has moved from a “research only” environment to mainstream clinical MR imaging. This is possible due to the improvements and developments with the Siemens unique Tim technology, now available on MAGNETOM Trio, A Tim System and available for existing MAGNETOM Trio systems. Many Siemens customers have already upgraded to the Tim technology and their testimonies offer proof to the benefits of Tim.

Karen Ziadie is an MR Clinical Specialist from Baptist Hospital in South Florida and she offers the following insight:

Our upgrade experience
“In February 2006, we received the Tim upgrade on our MAGNETOM Trio” Karen states.” We were one of the first Trio systems to receive the Tim upgrade so a team of engineers (some from Germany along with our local service personnel) descended on our site and set to work. We had been informed that the upgrade would take three full weeks. We wondered why an upgrade would take that amount of time, we soon found out why. The engineers literally stripped the system of everything except the magnet. Every component was replaced, including the gradients, the rf transmitter, table, surface coils, even the computer.”

With Tim capabilities are expanded
The gradients and rf systems bring dramatic changes to the performance available at 3T. The TQ Engine gradients have similar peak and slewrate, however the gradient linearity improves to the best in the industry at 1.7% across 50 cm, and the audio comfort of the Tim system reduces noise by up to 90%. Previously the body coil limited the field of view to 40 cm, now with a full 50 cm field of view many users are able to scan the same protocols between their 1.5T Siemens scanners and the MAGNETOM Trio, A Tim System.

Tim brings Matrix coils, as the MAGNETOM Trio before Tim relied solely on third party coils, and as a result was not able to be integrated to a level that allowed syngo GRAPPA to be implemented outside of the brain and 8-channel body coil.

New look
Oh what a difference! Our “new” 3.0T Trio looks awesome! Now it looks like the high tech imaging machine that it is. In addition to the new high tech look, the magnet bore is also much shorter and therefore much more patient friendly.

Coils
With the Tim Matrix coils we can quickly and efficiently do multiple studies without taking the patient off the table and...
without changing the coil configuration. Many of our patients have orders for brain, cervical, thoracic and lumbar spines, and sometimes hips. With the Matrix coils we can do all five studies without moving the patient and changing the coils. In addition, we can now use iPAT (integrated Parallel Acquisition Technique) with every coil.

The coils are also flexible. These high resolution foot images were acquired using the Body Matrix coil. Imaging times are very short and iPAT is used exclusively.

**The images**

We immediately noticed how much nicer the images looked. The images now have more signal to noise than before Tim. We are able to take advantage of this increased signal by utilizing iPAT, thinner slices, and higher resolution factors. We now routinely use 2 mm and 3 mm slice thicknesses for our brain and orthopedic studies. Many abdomen studies now employ a 3 mm or 4 mm slice thickness. This knee scan was acquired using DESS (Dual Echo Steady State) and an isotropic voxel size of 0.6 mm. iPAT and high signal to noise allowed this scan to be acquired in just over four minutes. High resolution allows multiplanar reconstruction with no loss of resolution.

**Special Absorption Rate (SAR)**

One of the big challenges with 3.0T imaging was SAR. Almost every protocol required the technologist to adjust the parameters as not to exceed the SAR limitations. The Tim upgrade has reduced the challenge of SAR at 3T. We are able to achieve lower TR values without exceeding the SAR limitations, resulting in reduced scan times and protocols that run smoothly without technologist adjustments required. With the new operating system also comes an increase to the number of SAR reduction methods. In addition to Hyper Echo, GRAPPA, and SPACE there are now low SAR pulses for ceMRA, VERSE is available, which through a different gradient pulse shape dramatically lowers rf power.

**T1 contrast**

The upgrade has provided greatly improved T1 contrast abilities. As a result of reduced SAR we are able to utilize TSE-T1 pulse sequences for spine imaging and we consistently obtain good T1 contrast in the spine. The Spine Matrix coil provides a major improvement in signal to noise and iPAT imaging. The previous coil did not offer iPAT capabilities.
T1 DarkFluid imaging as well as VIBE sequences (Volume Interpolated Breathhold Examination) demonstrate excellent T1 contrast and very high signal to noise. High resolution spine imaging has now become reliable and very fast for clinical imaging.

**Abdomen examinations**

“Abdominal studies have been difficult on the 3.0T. We were very pleasantly surprised by the quality of the abdominal images after the upgrade, they were excellent. Breathholds are shorter and we are now able to take advantage of the extra signal 3T provides by using higher resolution factors. Additionally, the T1 contrast improved, and we have seen dielectric shading substantially reduced.

Our radiologists liked the images so much we now routinely do abdomen studies on our 3.0T. In fact, there are some body studies that our radiologists definitely prefer to be done on the 3.0T such as adrenal, urography and prostate studies.” says Karen Ziadie.

“Sequence development and Tim have improved signal to noise, and reduced artifacts since the upgrade was completed. “

This upgrade is, all in all, the best upgrade that we have ever received. Image quality, scan times and throughput on the MAGNETOM Trio, A Tim System are greatly improved and we routinely obtain spectacular images. The 3.0T is now a fully functional clinical system.”

Karen and I had a great experience with the Tim Trio upgrade. Many other sites have as well. Similar results have been experienced on the Symphony Tim upgrades, too. The very reliable MAGNETOM Symphony is now a cutting edge system with the Tim upgrade.
Variations in microvascular structure and pathophysiology give rise to temporospatial variation in enhancement pattern, which provides valuable information on tissue characteristics. Dynamic contrast enhanced MRI (DCE-MRI) is widely used nowadays for lesion characterization. It is also used for monitoring therapy and for detection of residual or recurrent tumor.

**Pathophysiology of the contrast enhancement**

Malignant tumors enhance faster than normal tissue. The enhancement of a lesion depends on the microvascular density, endothelial permeability and extravascular space size. In tumors, development of neovascularization results from angiogenesis. The multifunctional cytokine vascular endothelial growth factor (VEGF), also known as vascular permeability factor (VPF), induces angiogenesis and strongly increases the microvascular permeability.

Release of the promotor substance stimulates sprouting of endothelial cells from the walls of pre-existing small vessels, till the sprouts encounter another vessel with which they connect, allowing blood to circulate. The vesiculo-vascular organelles (VVO) are grape-like clusters that span the entire thickness of the vascular endothelium connecting between the vascular lumen and the extravascular space [1]. The number of VVOs and leakage rate are regulated by VEGF. The VVOs might be an important pathway for Gadolinium-chelate MRI contrast agent leakage [2]. The morphology of the malignant tumors is bizarre and the capillaries are coarse, irregularly constricted/dilated and distorted. The capillary walls have numerous openings, widened inter-endothelial junction and discontinuous or absent basement membranes. These defects make tumor capillaries leaky. Slow growing benign tumors on the other hand, show regular vascular morphology.

In many tumor types including breast, lung, prostate and head and neck cancers, measurements of microvasculature density (MVD) on histopathological samples correlate with clinical stage and act as prognostic value. On this basis, it has been suggested that DCE-MRI may also be able to provide independent indices of angiogenic activity and therefore act as a prognostic indicator.

**Technique**

Contrast is administered through a peripheral vein – preferably the antecubital vein on the right side. A single dose of contrast (0.1 mmol/kg) of a standard Gd-chelate is administered at a rate of 4 ml/sec, using a pressure injector, followed by a chaser injection of 20–30 ml of saline at the same flow rate to empty the draining vein.

**Data acquisition**

While analyzing the lesion, it is important to include the entire volume of the tumor in the region of interest. An artery should also be included in the region of interest for comparison of the signal intensity in the plasma to the signal intensity in the lesion, which in turn depends on the density of the microvasculature, capillary permeability and size of the extravascular space.

**There are two generic approaches for data acquisition**

1. Susceptibility based techniques use T2/T2* sequences. When the bolus of exogenous contrast is injected, its paramagnetic property introduces a decrease in the T2/T2* relaxation properties by dephasing. Associated with susceptibility induced gradients surrounding the paramagnetic contrast agents, the effect is significant where the contrast is compartmentalized. It has important applications in brain perfusion due to the blood brain barrier.

![Mean curves. Type I, II, III, IV and V curves are well-seen.](image)
[Figure 2 A, B] **Glioblastoma multiforme.** The contrast enhanced T1-weighed axial image (A) shows a focal lesion (arrow) in the left corona radiata, which does not enhance. However the RCBV map (B) revealed elevated CBV (arrow), suggesting malignancy. A targeted biopsy revealed a glioblastoma multiforme.

[Figure 3 A, B] **Low-grade glioma.** A left frontal lesion is seen (A), which shows a maximum normalised CBV ratio (B) of 0.19, compared with the contralateral white matter, suggesting a low-grade tumor.
2. Dynamic relaxivity contrast enhanced MRI uses T1-weighted images to detect the relaxivity effects of the contrast agent. Thus the intra and extravascular contrast will raise signal intensity.

Sequences used:
For brain perfusion: 60 measurements over a period of 1.35 min at TR/TE: 1490/40, slice thickness: 5 mm, field of view (FoV): 230, matrix: 128 x 128, EPI (echo planar imaging) factor: 128, bandwidth: 1502.
Dynamic contrast enhanced tumor protocol for the rest of the tissues: 20 measurements over 5 min, at TR/TE: 3.13/1.13, slice thickness of 1.2–2 mm, FoV: 280, slab thickness 64, matrix: 256 x 140, bandwidth: 420.

Evaluation of the data
Postprocessing is done using dedicated software for image subtraction, multiplanar reconstruction (MPR), maximum intensity projection (MIP), signal intensity-time curve analysis from relaxivity based T1-weighted sequences and mapping rCBV (relative cerebral blood volume), rCBF (relative cerebral blood flow), MTT (mean transit time) from susceptibility-weighted T2/T2* sequences. Whichever the approach selected, careful inspection of the original dynamic contrast series images is important. The signal intensity-time curve contains information in terms of amplitude and time of arrival of the contrast and curve slope. It is a semiquantitative method of analyzing the vascular density, permeability of vessels and size of the extravascular space. The various patterns have been classified into 5 basic types (Fig. 1).

Indications
Brain tumors
In gliomas, tumor capillary blood volumes measured by DCE-MRI have shown to correlate with and predict tumor grade [3]. The rCBV maps identify the areas of malignant transformation or tumor dedifferentiation. This also helps to more accurately target stereotaxic biopsy (Fig. 2).
The rCBV measurements help in differentiating high grade from low grade glioma. The ratio of rCBV (tumor / rCBV in normal white matter) in high grade gliomas is 3.33 while for low grade gliomas is 1.51 (Fig. 3).

Osteosarcoma
A mid-sagittal T1-weighted contrast-enhanced image (A) was chosen for analysis. The heterogeneity is well seen. The red curve is the arterial curve obtained through the popliteal artery for reference. The yellow and blue curves are through solid tissue, showing type V and type III patterns of enhancement. Areas of necrosis are also seen, as shown by the green curve, which is flat.
The rCBV measurements also help to differentiate between primary gliomas and a solitary cerebral metastasis depending on the difference in peritumoral rCBV measurements. In metastases, the peritumoral vasogenic edema is due to extravasation of the contrast while in high grade gliomas, there is peritumoral edema and tumor cell infiltration.

DCE-MRI helps to differentiate between extra-axial tumors and intra-axial tumors as extra-axial tumors have increased rCBV. Absence of neovascularization in malignant lymphoma leads to decreased rCBV thus helping in differentiating it from glioma.

The rCBV values for tumefactive demyelinating disease are low, ranging from 0.22–1.79 (n=12), with a mean of 0.88+/−0.46, compared to intracranial glioma with rCBV values of 1.55–19.22 (n=11) with a mean of 6.47–6.56 [4].

**Bone and soft tissue tumors**

The diagnosis of a bone tumor is made on the plain radiographs. On MRI, the radiologist is required to report on the staging, response to therapy and in the detection of recurrent/residual tumor.

Osteosarcoma is the most common childhood tumor. Pre-operative chemotherapy has produced dramatic improvement in the prognosis of the disease. Response to chemotherapy is considered good if there is at least 90% tumor cell necrosis.

Mean curve analysis from T1-weighted dynamic sequences in the region of interest are helpful in characterization of primary soft tissue tumors by defining the enhancement pattern of the tumor and in approximately defining the percentage of necrotic tissue for bone tumors [6]. Mean curve analysis helps to evaluate the response to initial chemotherapy after 2 cycles, as with good response, the size of the tumor will decrease and the area of necrosis will increase. The rapidly proliferating areas show a type V curve, viable soft tissue with an increase in the extravascular space, type III, viable marrow with stable micro-circulation type IV and necrosis, a type I curve (Fig. 4). The mean curve analysis also helps to evaluate residual/recurrent tumor tissue which will show a type VII/V curve.

The percentage of tumor necrosis following pre-operative chemotherapy has predictive value of disease free survival. This is especially important with Ewing’s sarcoma/PNET and osteosarcoma.

**Other Tumors**

Various studies have been performed to evaluate the application of mean curve analysis in the liver, bladder, prostate and bone and joint pathologies.

A study was conducted by Workie [7] to quantify dynamic contrast enhanced MR imaging of the knee in children with juvenile arthritis based on pharmacokinetic modeling. This helped to monitor the degree of inflammation and therapeutic response during the early phase. This method can also be applied to adults with rheumatoid arthritis. In bladder tumors, DCE-MRI helps in staging and predicting tumor response [8]. Using DCE-MRI, analysis of microvascular density and evaluation of the vascular characteristics can be done pre-operatively in cases of carcinoma of the prostate [9].

DCE-MRI studies were also performed to analyze perfusion changes in advanced hepatocellular carcinoma treated with antiangiogenic agents [10].

*The information about this product is preliminary. The product is under development and not commercially available in the US, and its future availability cannot be ensured.*

References


A Practical Approach to Lung MRI at 1.5T

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Introduction
The value of MRI in imaging cardiac and large vessel disease is widely accepted, but the largest organ of the thorax, the lung, is usually investigated with X-ray and CT. MRI suffers from well-known limitations such as constant artifacts from heart pulsation or respiration and the low signal intensity of the aerated lung. This results from low overall proton density in combination with high susceptibility artifacts at air-tissue interfaces. The broadly accepted routine indications for MRI of the chest are instead focused on soft tissue processes such as chest wall masses or tumors of the mediastinum. The excellent contrast and the capacity of MR to produce cross sectional images or three-dimensional data sets in any orientation designate it as the ideal tool for this purpose. However, X-ray and CT are constantly criticized for the associated radiation exposure, in particular where applied for frequent follow-up examinations of children, or during pregnancy. A non-invasive method for follow-up of lung diseases during clinical trials or for physiologic research is therefore highly desirable. Since the recent technical developments in the field have contributed to overcome the well-known limitations of lung MRI, we can now recommend a comprehensive routine imaging protocol for the whole chest including lung parenchyma diseases.

Clinical method description
To match the criteria of a comprehensive imaging protocol for clinical routine, the recommended protocol needs to keep within a reasonable time frame of 15 to 30 minutes. It uses fast sequences for single- or multiple breathhold imaging and does not require cardiac or respiratory triggering. This facilitates patient positioning and shortens the room time. Limited information on cardiac pathology is provided with the single shot T2-HASTE and the free-breathing TrueFISP. If further cardiac imaging is planned, ECG-triggering would be needed. The key to high image quality without respiratory motion artifacts is appropriate instruction of the patient. Nevertheless, the T2-HASTE as well as the free-breathing TrueFISP are robust against breathing motion and can therefore be applied even with completely uncooperative patients. The vessel imaging capacity of the TrueFISP will even allow detection of large central pulmonary emboli in dyspneic patients. Only one sequence – the optional high-resolution – T2-TSE uses a navigator.

We present below a basic protocol for non contrast-enhanced MRI of the lung, followed by recommendations for additional contrast-enhanced and functional imaging (table 1). This would be applicable to children from 8–10 years and adults. For younger children, spatial resolution and Field of View

[Figure 1] 45-year-old male with cystic fibrosis of the lung. Bronchiectases in both upper lung lobes with mucous plugging being represented by bright signal on the T2-weighted HASTE (left image) and intermediate signal of the mucus within the bright signal of the contrast-enhanced bronchus walls in the transverse 3D-GRE (VIBE) sequence. For recommended sequence parameters see table 1.
(FoV) would need to be adjusted. All components are based on common MR sequence components of current standard installations. In detail, they refer to 1.5T MAGNETOM Avanto protocols and a Body Matrix coil for thoracic imaging, but they would also be applicable to MAGNETOM Sonata and MAGNETOM Symphony installations with minimal adjustments. Where parallel acquisition techniques are not available, multi-breathhold acquisitions can be used instead. If a navigator is not available, a respiration belt could be used instead. For all coronal sequences (except for perfusion and angiography to save reconstruction time), distortion correction (“large FoV”) is activated and excitation order is interleaved, where appropriate.

**Basic protocol for non-contrast enhanced MRI of the lung**

The imaging protocol starts with a gradient echo localizer in inspiration. The first sequences are acquired in breathhold, usually starting with the coronal T2-HASTE followed by the transverse T1-weighted 3D-GRE (VIBE). After this the first set of coronal SS-GRE sequences is acquired in free breathing, giving the patient some time to recover from the breathhold maneuvers. This is followed by the T2-TIRM image series, which is acquired with multiple breathholds. Anatomic coverage should include the upper abdomen with liver and adrenal glands. This basic protocol may be completed with the single slice dynamic SS-GRE series for diaphragmatic function. For this series, the patient is instructed to breathe deeply. Total room time up to here will be approximately 15 minutes.

This basic protocol covers a range of routine indications, e.g. follow-up examinations in children with cystic fibrosis (Fig. 1) [1]. The sensitivity for lung nodules larger than 4 mm ranges...
between 80 and 90% and reaches 100% for nodules larger than 8 mm [2, 3]. Depending on the water content, nodules can be detected either on the VIBE or on the HASTE and T2-TIRM images. Atelectasis and tumor can be well distinguished on T2-weighted sequences, especially if the high resolution T2-TSE is added. If MRI is applied for staging of lung malignancies with non-contrast enhanced images, T2-TIRM is an obligatory protocol component for the detection of bone metastases [4] (Fig. 2). The dynamic coronal TrueFISP acquisitions allow estimating diaphragmatic function and the mobility of intrathoracic masses (Fig. 3). This particular part provides additional functional information to the morphologic images and is one of the essential differences between lung MRI and any other common imaging modalities [5].

**Basic protocol including contrast enhanced MRI of the lung**

For routine purposes it might appear sufficient to conclude the study, if the non-enhanced scans show completely normal findings. Nevertheless, application of i.v. contrast material markedly improves the diagnostic yield of 3D-GRE imaging of the lung by the clearer depiction of vessels, hilar structures and pleural enhancement. Parenchymal disease and solid pathologies are also enhanced. Thus, a study to exclude pulmonary malignancies e.g. for staging purposes should usually comprise a contrast enhanced series, preferably with a fat-saturated 3D-GRE sequence. Contrast enhancement is also necessary for pleural processes (empyema, abscess, metastatic spread of carcinoma, mesothelioma) or for the further evaluation of solid masses, as well as for functional imaging or angiography. If it is intended to include a contrast-enhanced series, the 3D-GRE (VIBE) sequence in the pre-contrast series should be applied with fat-saturation to allow for a direct comparison of contrast uptake. Contrast material: 0.2 mmol/kg i.v. per hand or power injector. The contrast enhanced VIBE extends the range of indications for the protocol towards lung cancer and malignant and infectious pleural processes [6]. It is important to know that calcified nodules or masses may be invisible either on the non-contrast enhanced as well as on the contrast enhanced images. In most cases, this reflects benign findings but might be crucial e.g. for the staging of osteosarcoma.

**Lung perfusion imaging**

The basic principle of contrast-enhanced perfusion MRI is a dynamic MR image acquisition following an intravenous bolus injection of a paramagnetic contrast agent. Perfusion MRI of the lung requires a high temporal resolution in order to visualize the peak enhancement of the lung parenchyma. The recommended fast acquisition technique is based on iPAT (Integrated Parallel Acquisition Techniques) and data sharing. It allows for a 3D data acquisition with a temporal resolution of 1.5 seconds per image [7]. The resulting 4D-data set can be displayed with the "Mean Curve" application, which allows one to scroll through the series in a single image position or to scroll through the images of a 3D data set obtained at a single time point. A quick, semi-quantita-
Indications for perfusion MRI

- emphysema (indirectly)
- pulmonary hypertension
- acute and chronic pulmonary embolism

tive analysis of contrast-enhanced perfusion MRI data consists of the calculation of signal time curves, SNR and contrast-to-noise ratios (CNR) using region-of-interest (ROI) analysis of the signal of the lung tissue. For documentation, contrast-enhanced 3D perfusion MRI is usually processed by subtraction of mask image data acquired before contrast bolus arrival (Fig. 5). Since the acquisition time and diagnostic yield are at least the same as for a test bolus, the dynamic perfusion series may be used to prepare the angiogram. A disadvantage compared to the test bolus method is the additional time needed for image post-processing and the slightly higher amount of contrast medium (0.07 mmol/kg patient weight). To allow for an exact calculation of the time points for the angiograms, injection speed and the volume of the bolus plus sodium chloride chaser should be the same as for the following angiogram.

Pulmonary angiography

Contrast-enhanced MRA uses a T1-weighted 3D-FLASH acquisition after intravenous injection of a paramagnetic MR contrast agent. A short TR allows for breathhold acquisitions and a short TE minimizes background signal and susceptibility artifacts. The flip angle of 25 degree produces a high contrast between lung tissue and the vessels. Adequate breathhold and exact timing of the contrast agent (0.1 mmol/kg at 5 ml/s followed by a 20 ml sodium chloride chaser, time to center 8.7 s) with an automatic power injector are essential prerequisites. Three acquisitions (non-contrast enhanced, centered on the peak signal of the pulmonary artery and centered on the peak signal of the aorta) are appreciated. For comprehensive viewing, we recommend use of the 3D-tool for multi planar reformation (MPR) or maximum intensity projections (MIP). The combination of 4D MRA perfusion and CE-MR angiography provides a useful alternative to perfusion scintigraphy and CT angiography, thus extending the range of indications for lung MRI by a considerable number of clinical conditions, e.g. suspected pulmonary embolism in pregnancy [8]:

Indications covered by lung CE-MRA

- pulmonary hypertension
- acute and chronic pulmonary embolism
- AV-malformation (e.g. M. Osler)
- pulmonary sequestration
- Swyer-James-Syndrom
- pulmonary artery aneurysms
- pulmonary vein anomalies
- masses of the hila
- tumour invasion a. pulmonalis /aorta

Respiration-triggered high resolution T2-TSE

Further options to extend the standard protocol are T1- and T2-weighted Spin Echo (SE) or Turbo Spin Echo (TSE) sequences with respiratory triggering (or gating). T1-weighted images are usually recommended for the detection of lymph nodes and tumor infiltration into the chest wall, but only the T2-weighted sequences contribute to the evaluation of lung
Indications to add HR-TSE

- interstitial lung disease
- lymphangitic spread of carcinoma
- differentiation of tumour/atelectasis
- masses with chest wall infiltration
- collateral findings (liver, adrenals)

Parenchyma pathology and provide equal information about the chest wall and mediastinum. Since these triggered sequences are time-consuming, we recommend the inclusion of only one additional T2-TSE series into the protocol, since all aspects are covered by the previous sequences.

Conclusion

The proposed imaging protocol covers a wide range of clinical indications for lung imaging and may be used for routine purposes whenever it is mandatory to avoid radiation exposure as far as possible (e.g. in childhood and pregnancy). Moreover, its potential capacity to add functional information such as analysis of diaphragm motion and lung perfusion studies make it more that just a surrogate for chest x-ray and CT. The suggested sequences for lung MRI can be easily set up on state-of-the-art MR scanners by using the Phoenix sample files from www.siemens.com/MAGNETOM-World.
### Sequence parameters for MRI of the lung

<table>
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<tr>
<th>Basic protocol 15'</th>
<th>Non-CE lung MRI</th>
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<th>CE T2-HR</th>
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<td>x</td>
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<td>x</td>
<td>(x)</td>
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<th>free</th>
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| TA (min:s)        | 0:10             | 0:18    | 0:20    | 0:56  | 1:28       | 0:19     | 0:29    | 0:21    | 0:20       | 5:05+|
| Slices per acquisition | 7               | 30      | 72      | ≤128  | 32x4       | 1        | 32x20   | 120     | 72         | 45   |

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<th>R&gt;L</th>
<th>A&gt;P</th>
<th>R&gt;L</th>
<th></th>
</tr>
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| FoV (mm) [FoV phase %] | 500 [100] | 500 [100] | 400 [87.5] | 450 [100] | 400 [75] | 400 [100] | 500 [100] | 400 [83.3] | 500 [79.7] |

| Base resolution    | 256              | 256     | 256     | 256   | 256     | 256   | 256     | 256     | 256       | 512  |
| Phase resolution (%) | 75               | 100     | 100     | 100   | 75      | 66    | 54      | 90      | 100       | 75   |
| Slice thickness (mm) | 10              | 8       | 4       | 4     | 6       | 10    | 5       | 1.6     | 4         | 4    |
| Phase partial Fourier | 6/8             | 4/8     | off     | off   | off     | off   | 6/8     | 6/8     | 5/8       |      |

| Pixel size (mm)    | 2.6x2.0          | 1.8x1.8 | 1.6x1.6 | 1.8x1.8 | 1.7x1.3 | 2.4x1.6 | 3.6x2.0 | 1.2x1.0 | 1.6x1.6 | 1.3x1.0 |
| Distance factor    | 50%              | 0%      | 20%     | -50/0% | 10%     | n.a.   | 20%     | 20%     | 20%       | 10%  |
| TR (ms)            | 8.9              | 600     | 3.15    | 437.2  | 3500    | 317.1  | 1.64    | 2.75    | 3.15       | 1700 |
| TE (ms)            | 4.38             | 31      | 1.38    | 1.16   | 106     | 1.14   | 0.64    | 1.12    | 1.38       | 100  |
| Flip-angle (degr.) | 30°              | 180°    | 8°      | 80     | 150     | 67     | 40      | 25      | 8°         | 150  |
| Band width (Hz/pixel) | 180             | 610     | 500     | 1030   | 252     | 980    | 1220    | 384     | 500        | 195  |
| Large FoV (dist. corr.) | off             | on      | off     | on     | off     | off   | off     | off     | off        | on   |

| Comments           | cor/sag A>P/tra R>L | 2 concatenations | FatSat, if CE VIBE is planned | Distance factor-50% WIP-Package | TI150 ms, 5 concatenations | 60 measurements at 3s | 20 measurements 1.5 s each, if SAR limit exceeded | Navigator needed 5 concatenations | With FatSat activated, identical with 3 | Navigator needed 5 concatenations |

**Large FoV (dist. corr.)**
- off on off on off off off off off on
Case Report MAGNETOM Avanto
Advanced Breast Cancer: MR Monitoring* of Neoadjuvant Chemotherapy

Bruce A. Porter, M.D., FACR
First Hill Diagnostic Imaging, Seattle, Washington, USA

Patient history
51-year-old marathon runner noted swelling, redness, and tenderness of her left breast that did not respond to antibiotics. Biopsy confirmed inflammatory infiltrating ductal cancer with associated high-grade ductal carcinoma in situ (DCIS). MR was requested for staging and to monitor neo-adjuvant chemotherapy of locally advanced breast cancer in a patient with mammographically dense breasts.

Imaging findings
The initial, baseline, MR exam documented inflammatory thickening of the skin, lymphedema, and malignant axillary adenopathy on pre-contrast STIR chest (Fig. 1A) and breast coil STIR images (Fig. 1B) (STIR: Short TI Inversion Recovery). An axial maximum intensity projection (MIP) of the first post-contrast DynamicVIEWS acquisition (Fig. 1C) demonstrates the extreme vascularity of the large (5.6 cm) and partially necrotic primary tumor as well as numerous nodules of extensive DCIS and/or additional foci of invasive tumor. Three months later, a follow-up axial MIP (Fig. 2A) revealed not only lack of response to the initial chemotherapy, but also enlargement of the small foci of DCIS or invasive cancer. An axial, 60-second post-contrast subtraction image (Fig. 2C) and corresponding computer aided detection (CAD) color parametric image (Fig. 2B) graphically depict the extreme vascularity, rapid enhancement and wash-out of this still viable tumor. Sagittal thin section InterVIEWS image (Fig. 2D) confirms skin invasion. The axillary lymph nodes remained enlarged. As a result, anti-angiogenic therapy was begun in an attempt to control the growth of this aggressive tumor. At 5 months, after prolonged and intensive therapy, the tumor...
felt somewhat softer on exam; however, although the sagittal InterVIEWS (Figs. 3A and 3B) documented some regression of the smaller tumors, the main mass is larger and has extended further into the skin. The malignant nature and unresponsiveness of this tumor are unfortunately made very clear with MR monitoring.

**Discussion**

Pre-operative (neoadjuvant) chemotherapy is increasingly used for larger, locally advanced but not metastatic, breast cancers. The intent is to shrink the tumor, establish tumor chemo-sensitivity, and to possibly allow breast conservation if a good response is achieved. However, monitoring of the effectiveness of treatment is notoriously inaccurate with physical exam, mammography, and ultrasound. MR has been shown to be quite an accurate method to assess treatment-related changes, which helps make decisions on therapy or surgery more tailored to the individual patient and tumor. Although PET-CT can also be used for neoadjuvant monitoring, MR has much better resolution for subtle breast findings; it is also much less expensive, more available, and does not expose the patient to additional radiation.

This case represents a particularly aggressive tumor, in which MR demonstrated rapid progression even during the most intense chemotherapy currently available. The prognosis for the patient is predicted by the MR findings and is poor.

**Figure 1C**  Axial full maximum intensity projection (MIP) from the 60-second data set shows a dominant mass posteriorly, multiple smaller nodules of malignant enhancement elsewhere, and markedly asymmetrical venous drainage.

**Figure 2A**  Follow-up MR documented absence of significant response to therapy as well as progression and coalescence of the smaller nodules, though the adenopathy was somewhat improved.

**Figure 2B**  Corresponding CAD color overlay indicates heterogeneous, persistent malignant enhancement with a significant amount of wash-out kinetics (red voxels). Plateau type kinetics is portrayed as yellow and persistent enhancement as blue. The axillary node is not appreciably labeled. The right breast remains normal.

**Figure 2C**  Coronal full MIP reconstructed from the axial 60-second data set is useful to localize and assess the full extent of the invasive malignancy and the ductal carcinoma in situ (DCIS) of the left breast. Note the contralateral venous drainage.
**Figure 2D**  The thin (0.8 mm) slices and 0.7 x 1.0 mm in-plane resolution of the DynamicVIEWS bilateral subtraction images allow high quality oblique MIP images for direct comparison to mammography (Lt. MLO view).

**Figure 2E**  True lateral mammogram of the left breast (rotated to match Fig. 2F) illustrates the difficulty of assessing tumor size and extent in radiographically dense breasts.

**Figure 2F**  Sagittal InterVIEWS at 2 min post-contrast with 0.64 mm slice thickness and 0.5 x 0.5 mm in-plane dimensions clearly shows both the centrally necrotic main tumor mass and a satellite nodule (arrow). Note the skin thickening and enhancement due to tumor infiltration (i.e. this is a T4 lesion).

**Figure 3A**  Despite further chemotherapy and addition of anti-angiogenesis drugs, the sagittal InterVIEWS exam (when compared to Fig. 2F) readily documented progressive growth of the main tumor mass and further skin infiltration. However, the adjacent small satellite lesion (arrow) became smaller.

**Figure 3B**  At the level of the nipple, ductal enlargement and a ductal enhancement pattern indicates residual DCIS and fluid.

*Works In Progress – The information about this product is preliminary. The product is under development and not commercially available in the US, and its future availability can not be insured.*
Soft Tissue Motion Correction for MR Mammography (BRACE)*

Melanie Schmitt

Siemens R&D, Erlangen, Germany

Introduction
Dynamic imaging is an important and valuable diagnostic tool for the diagnosis of breast tissue lesions in MR mammography. Typically, fast T1-weighted CE imaging is used to monitor the concentration of a contrast agent during its passage through the tissue. The examination starts with a pre-contrast scan followed by the injection of the contrast agent and 6-8 post contrast examinations. To facilitate the detection of suspicious enhancing areas different post processing tools are used, including subtraction of post-contrast and pre-contrast images and evaluation of the signal time curves in suspicious lesions. Therefore it is essential that there are no changes in the patient’s position during the entire measurement. However, because of the time duration of 6-8 min for the CE MRI examination, this requirement of no movement between measurements is often not met and may thus affect the interpretation of the images or make an accurate diagnosis even impossible.

Regardless of the many efforts undertaken to reduce or suppress motion artifacts, including optimized choice of MR sequences and protocols, instructions to the patients and fixation of body-parts, there still exists quite a substantial number of examinations which suffer from motion. Consequently registration algorithms are increasingly used to correct for these artifacts. But, image registration is particularly difficult in MR mammography because of the complex, non linear elastic behavior of breast tissue and the possibility of strong local movements.

This article describes a new motion correction, BRACE – developed by Siemens – that overcomes the limitations of existing methods in MR Mammography. BRACE (soft tissue motion correction package) will be available with the new software syngo MR B13.

Motion correction
To compensate for patient motion during the CE MRI examination two different non-rigid registration algorithms of BRACE are now available. Both correction possibilities can compensate for patient motion in all directions in 2D and 3D MR data sets. The first correction form is a Gaussian-weighted least mean square registration algorithm, which also takes into account the signal change due to the inflow of the contrast agent. The fast algorithm can perform motion correction of a 3D data set including 6 repetitions with 80 slices...
Clinical applications

The new BRACE correction tools for dynamic breast MRI were used for different 2D and 3D patient data sets, in which motion of the patient was clearly visible. Fig. 2 shows the example of a patient data set, where the fast correction algorithm was used. The images were acquired with a 2D T1-weighted FLASH (Fast Low Angle Shot) sequence on a MAGNETOM Symphony scanner with the following parameters: TE/TR/flip angle = 4.76 ms / 113 ms / 80°, matrix 384 x 384, slice thickness 3 mm, iPAT (integrated Parallel Acquisition Technique) factor 2. In the corrected images the lesion is much better delineable compared to the uncorrected case. Fig. 3 shows another example of a patient with motion artifacts. In this case the images were acquired with a 3D T1-weighted FLASH sequence on a MAGNETOM Sonata scanner with the following image parameters: TE/TR/flip angle = 6.7 ms / 113 ms / 15°, matrix 256 x 256, slice thickness 3 mm, iPAT factor 2.

Conclusion

A new motion correction BRACE has been developed and optimized for MR Mammography. This motion correction can handle large datasets acquired in breast MR and enables a higher diagnostic confidence when suspicious areas are under evaluation.

* The information about this product is preliminary. The product is under development and not commercially available in the US, and its future availability cannot be insured.
CARDIAC

Case 1

Acute Myocardial Infarction*

Jeanette Schulz-Menger, M.D., Philipp Boye, M.D.
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Introduction
A 44-year-old man had complained of sudden chest pain 5 days previously. Acute myocardial infarction was indicated by ST-elevation in ECG and increased cardiac enzymes. The coronary angiography showed an occlusion of the RCA (right coronary artery) and also stenosis of the LCX (left circumflex artery). A PCI (percutaneous intervention) of both vessels was performed. Cardiac MRI was performed to assess the extent of myocardial damage.

Image findings
The left ventricular dimensions are normal. There is impaired systolic function (ejection fraction: 34%) with regional wall motion abnormalities: akinesia of the inferior, septal, posterolateral wall and the apex. Tissue characterization imaging using PSIR (Phase-Sensitive Inversion-Recovery) sequence shows extensive transmural scar of the inferior wall including a large area of microvascular obstruction. A pericardial effusion is also seen (Figs. 1 and 2).

Results
Tissue characterization imaging can be used early after myocardial infarction to assess the extent of myocardial scar. This extent determines the prognosis with regard to functional recovery of regions with wall motion abnormality. In this patient, although PCI of the right coronary artery was performed, there would be no expectation of functional recovery in the inferior wall due to the transmurality. Furthermore, the presence of microvascular obstruction is a risk factor for late remodeling. The single-shot PSIR sequence enables fast and accurate delayed tissue characterization imaging even under difficult patient conditions without the need of TI adjustment.

*Works In Progress – The information about this product is preliminary. The product is under development. It is not commercially available in the US and its future availability cannot be assured.
Case 2
Myocarditis*

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Patient history
The patient suffered a respiratory infection a week before examination and since then has complained of repeated chest pain. The ECG was indicative of an inferolateral ST-elevation myocardial infarction (STEMI). However, coronary heart disease was ruled out with the help of coronary angiography. Cardiac MRI (CMR) was performed due to suspicion of myocarditis.

MRI findings
The size and dimension of the left ventricle is within normal limits. There is hypokinesis in the inferolateral wall. IR Turbo FLASH 2D sequence shows multiple subepicardial foci of increased intensity after using tissue characterization imaging in the lateral and inferolateral wall (Fig. 2). TIRM (Turbo Inversion Recovery Magnitude) sequence acquired in the same orientations reveal edema in the inferolateral wall (Figs. 3 and 4).

Results and discussion
This patient shows patchy delayed enhancement in the inferolateral wall and edema demonstrated by TIRM to be in the same region: this is most probably due to myocarditis. CMR enables visualization of myocarditis in many cases by use of delayed enhancement and TIRM or TSE-T1-weighted imaging. In comparison to the pattern seen in patients with ischemic heart disease, hyperintensity after using tissue characterization imaging is distributed in a completely different way in patients with myocarditis: subepicardial or intramyocardial localization and more patchy and grayish appearance are typical findings. The pathology might be visible in one sequence only in some of the patients. Therefore, to ensure high sensitivity, the use of different sequences is recommended.

*The information about this product is preliminary. The product is under development. It is not commercially available in the US and its future availability cannot be assured.
Case 3

Apical Thrombus*

Jeanette Schulz-Menger, M.D., Philipp Boye, M.D.
Franz Volhard Clinic, Charité Campus Buch, Berlin, Germany

Patient history

This 72-year-old patient was referred to cardiology department with subacute myocardial infarction. The patient had had coronary artery bypass graft surgery in 1999 (grafts to right coronary artery and left anterior descending). One day prior to the MRI exam, a stent was placed in the coronary artery bypass graft to the LAD.

Image findings

Dilated, hypertrophic left ventricle with moderately reduced ejection fraction. Tissue characterization sequences show increased intensity in the anterior wall and basal inferior and posterior wall. The TIRM sequence shows edema in anterior wall but no edema in the basal inferior and posterior wall (Fig. 1). In LVOT orientation, an apical left ventricular thrombus with a liquid core can be seen. Follow-up investigation after 3 months of anticoagulation revealed complete resolution of the thrombus (Fig. 3).

Results and discussion

Cardiac MRI can be used for a comprehensive cardiac examination of patients with coronary artery disease. Beyond assessing global and regional wall motion abnormalities by use of cine-sequences, CMR reveals in this patient the presence of an apical thrombus, helps differentiating between acute and chronic infarction by use of a PSIR (Phase-Sensitive Inversion-Recovery) and TIRM (Turbo Inversion Recovery Magnitude) sequences. Both sequences used together improve the performance of CMR in the acute setting.

*The information about this product is preliminary. The product is under development. It is not commercially available in the US and its future availability cannot be assured.
Case 4
Aortic Stenosis

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Patient history
65-year-old patient with dyspnea on exertion (NYHA III). There is no history of syncope coronary artery disease or heart failure. Echocardiography reveals aortic stenosis with a pressure gradient of 76 (max.) and 49 mmHg (mean), respectively. Coronary MR imaging (CMR) was performed to evaluate the aortic orifice area.

Image findings
The MR images show clearly a bicuspid aortic valve. No left ventricular hypertrophy is seen and there is normal left ventricular function. Planimetry using 2D TrueFISP cine sequence reveals an aortic orifice area of 0.6 cm² (0.35 cm²/m² normalized).

Results and discussion
There was a moderate to severe aortic stenosis according to the pressure gradients only as assessed by echocardiography. CMR can be used additionally in patients with aortic stenosis to assess the aortic orifice area by planimetry. This helps to identify patients with indication for aortic valve replacement. In this patient the aortic orifice area is 0.6 cm², which corresponds to a severe stenosis. Accordingly, this patient underwent surgery.

*The information about this product is preliminary. The product is under development. It is not commercially available in the US and its future availability cannot be assured.
Case 5

Osteosarcoma Metastasis

Jeanette Schulz-Menger, M.D., Philipp Boye, M.D.
Franz Volhard Clinic, Charité Campus Buch, Berlin, Germany

Patient history
17-year-old man with osteosarcoma in the right lower leg. Pulmonary metastasis was seen. Cardiac MRI was performed due to abnormal finding in transthoracic echocardiography indicating an intracardiac mass in the right ventricle.

Image findings
Using cine sequences and T1-weighted Dark Blood images* in RVOT-orientation, two tumors in the right ventricular outflow tract with extension into the pulmonary artery trunk can be visualized. In TrueFISP cine-sequence the tumors appear mobile. In the T1-weighted sequence the masses appear isointense. The right ventricle is enlarged.

Results and discussion
This patient shows a cardiac metastasis of an osteosarcoma. In most reported cases the location of a cardiac metastasis of osteosarcoma is the right ventricle. Cardiac MRI allows the detection of cardiac tumors and intracardiac masses with high sensitivity. Using T1- and T2-weighted imaging and application of contrast agent, differentiation between thrombus and tumor and between different types of tumors can to some extent be achieved.

### Differential diagnosis between thrombus and tumors

<table>
<thead>
<tr>
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<td>High</td>
</tr>
</tbody>
</table>

* Isointensity    Hyperintensity    Hypointensity    Heterogeneous intensity

---

* Works In Progress – The information about this product is preliminary. The product is under development. It is not commercially available in the US and its future availability cannot be assured.

---

**Figure 1** Morphology, TSE Dark Blood, RVOT, GRAPPA 2

<table>
<thead>
<tr>
<th>TR/TE</th>
<th>907.7/24</th>
<th>slice</th>
<th>1</th>
<th>FoV</th>
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<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>SL</td>
<td>6 mm</td>
<td></td>
<td></td>
<td></td>
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</table>

**Figure 2** Turbo FLASH dynamic signal, RVOT, GRAPPA 2

<table>
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**Figure 3** Function, TrueFISP cine retro, RVOT, GRAPPA 2

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<td>SL</td>
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</tbody>
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Case 6
Acute Infarction

Jeanette Schulz-Menger, M.D., Philipp Boye, M.D.
Franz Volhard Clinic, Charité Campus Buch, Berlin, Germany

Patient history
67-year-old patient suffered acute coronary syndrome with mild elevation of cardiac enzymes. History of myocardial infarction in 1983. Echocardiography showed wall motion abnormalities. Cardiac MRI* was performed to differentiate between chronic and acute infarction.

Image findings
Hypertrophic left ventricle with global systolic function in normal range but regional hypokinesia of apical septal and anterior wall as well as the apex. PSIR and IR Turbo FLASH 2D tissue characterization sequences show increased intensity in the apical region and circumscribed in the midventricular lateral wall, both with transmural extent. TIRM sequence shows high signal of the myocardium in the lateral wall but no high signal in the apex. The high signal apically in the left ventricular cavity in TIRM indicates “slow-flow” phenomenon.

Results and discussion
Delayed enhancement imaging using PSIR or IR FLASH sequences allows detection of myocardial infarction. To differentiate acute from chronic infarction, TIRM or T2-weighted sequences can be used in addition to delayed enhancement imaging. These sequences depict acute infarction by visualizing edema. In this patient, CMR using TIRM sequence helped to identify the region of acute infarction (lateral wall).

* Works In Progress – The information about this product is preliminary. The product is under development. It is not commercially available in the US and its future availability cannot be assured.
We see a way to evaluate myocardial infarct and vascular disease within one exam without any patient repositioning.

Tim won’t let you miss a beat.

We see a way to determine regional ventricular function in real time allowing free-breathing using 12 matrix coil elements.

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Tim offers unmatched MRI capabilities for all cardiovascular exams without repositioning the patient. Ideal not only in diagnosing subendocardial infarct and congenital heart disease, but also systemic diseases like diabetes and atherosclerosis. With its 76 matrix coil elements and up to 32 RF channels, you enjoy revolutionary acquisition speed with virtually unlimited Parallel Imaging even in double oblique slice orientations. Tim transforms workflow from an exam limited by the dimensions of local RF-coils to one determined by the disease. Tim. Very heart smart.

www.siemens.com/Tim
Patient history

A 64-year-old man had a six months history of non-healing ulcer on the left foot and experienced intermittent claudication while walking less than 10 meters. He had been a long-standing heavy smoker, but had not used tobacco for the last 5 years.

The physical examination revealed a cachectic man, who had finger clubbing. On examination, we found no pulse on femoral arteries and a wound and gangrene of the left hallux.

MR Angiography (MRA) of aorta and lower limbs was requested for diagnosis and percutaneous intervention / surgical planning.

Imaging findings

The MRA showed diffuse and subtle atherosclerotic irregularity of the aorta, mild stenosis at the origin of the common iliacs and proximal occlusion of both external iliacs (Fig. 1). Common femoral arteries are filled through collaterals. There is a distal occlusion of the left superficial femoral artery indicating previous thrombosis. The right superficial artery is occluded in its mid portion. Deep femoral arteries are patent and serve as collateral pathways (Fig. 2).

Popliteal arteries are widely patent presenting subtle atherosclerotic irregularity. Proximal tibial anterior arteries and fibular are patent bilaterally and posterior tibial are occluded (Fig. 3).
Left anterior tibial artery has multiple severe stenosis and it is occluded distally. Left fibular is patent. Right anterior tibial and fibular arteries are widely patent (Fig. 4). A panoramic view is documented using the composing software (Fig. 5).

**Discussion**

Chronic arterial insufficiency is a multiple lesion complex disease. Usually, patients with such a condition suffer chronic pulmonary obstructive disease, strokes, myocardial infarction and nephropathy. Conventional angiography is an invasive and risk-related procedure, especially in this type of patient. A good quality diagnostic image is essential to determine location and severity of lesion. A therapeutic approach (vascular bypass or percutaneous angioplasty) is a very important decision.

Doppler ultrasound is limited and time consuming in multiple lesion patients and does not offer an angiographic image. Multislice CT may be used with the inconvenience of nephrotoxic contrast media, radiation and very time consuming reconstruction of original images. In patients with extensively calcified arteries, Multislice CT may be limited and difficult to determine severity of the lesions.

This case represents an advanced atherosclerotic disease, in which MRA offered a whole-body, good quality panoramic study.
Case Report MAGNETOM Avanto

Aorta Coarctation

Paulo R. Schwartzman, M.D., Ph.D.

Hospital Moinhos de Vento, Porto Alegre, Brazil

Patient history
A 30-year-old woman complains of shortness of breath during exercise. The patient denies smoking or any previous lung problem. The physical examination reveals a systolic heart murmur and a pulse difference between arm and legs. An echo was performed and demonstrated normal left ventricular function, normal heart valves but a turbulent flow at proximal descending aorta. MR Angiography (MRA) of the aorta was requested for diagnosis of a suspected aortic coarctation.

Imaging findings
The MRA was reconstructed with different techniques and confirmed the diagnosis of coarctation of the aorta. However, this also revealed an aberrant left subclavian artery. This diagnosis is an important point for surgical planning since reimplantation of the artery will be required. Thin maximum intensity projections (thinMIP) images (Fig. 1) illustrate the findings. In addition, Volume Rendering Technique (VRT) (Fig. 2) in different projections and colors demonstrates the vascular anomaly.

Discussion
Coarctation of the aorta is one of the most common congenital heart diseases and is often under-diagnosed. The patient has symptoms only with severe aortic obstruction. Usually the vascular anomaly is suspected during a routine physical examination due to heart murmur or a pulse difference between upper and lower limbs. Echo is the first choice and
often has difficulty in visualizing the arch and the proximal portion of the descending aorta. Conventional angiography was used in the past, but currently its use is not acceptable due to risks related to the procedure. A good quality diagnostic image is essential to determine location and severity of lesion. And magnetic resonance angiography has become the non-invasive procedure of choice due to an entire non-invasive visualization of the thoracic aorta. Multislice CT may be also used, but has the inconvenience of nephrotoxic contrast media and radiation exposure in usually young patients. This case characterizes a typical patient with long-standing aortic coarctation which is easily diagnosed by MRA.

[Figure 2] Posterior view with VRT images demonstrating the vascular anomalies.
Case Report MAGNETOM Avanto
Chest Pain and Myocarditis*

Paulo R. Schvartzman, M.D., Ph.D.
Hospital Moinhos de Vento, Porto Alegre, Brazil

Patient history
An 18-year-old athletic young man woke at 6 am with chest pain and intense sweating. After taking Tylenol and the persistence of the pain, the patient was taken to the emergency department. The ECG shows ST elevation in the anterior and lateral wall. The cardiac enzymes (CK, MB and troponin) were also elevated, suggesting acute myocardial infarction. The patient was taken to the cath lab, which shows normal coronary arteries. A cardiac MRI is requested to verify the presence and type of myocardial scar.

Imaging findings
The cardiac magnetic resonance exam consisted of cine images with TrueFISP and gradient echo sequences with grid tagging to evaluate myocardial contractility. In addition, tissue characterization images were acquired 15 minutes after CM dose to detect myocardial fibrosis*.

The cine TrueFISP images acquired at long-axis 4 chamber view (Fig. 1) and 3 chamber view (Fig. 2) demonstrate normal contractility. The gradient echo with SPAMM images at the same long-axis positions as the TrueFISP, demonstrate normal segmental contraction (Fig. 3).

The tissue characterization images with increased intensity reveal a mesocardial fibrosis in the posterior and lateral wall suggestive of myocarditis.

Discussion
Myocarditis is a difficult diagnosis. The symptoms may vary widely and not infrequently the patients are taken to the cath lab due to elevated cardiac enzymes to rule out myocardial infarction. The myocardial biopsy is negative in more than 50% of cases and imaging modalities have poor markers for inflammatory myocardial disease. Nuclear medicine may play a role in the diagnosis but the few studies in the international literature show controversial results. Recently, cardiac magnetic resonance has been used in the diagnosis of myocarditis and a typical mesocardial / subepicardial fibrosis, most common in the posterior and lateral wall, suggests the diagnosis. This case represents a typical patient with a difficult diagnosis with normal segmental contractility regardless of the presence of myocardial fibrosis. The elevated cardiac enzymes demonstrated myocardial injury, confirmed by high spatial resolution cardiac MR.

* Figure 1 *
4-chamber view at diastole (left) and at end-systole (right) demonstrating normal contractility.
**Figure 2**
3-chamber view at diastole (left) and at end-systole (right) demonstrating normal contractility.

**Figure 3**
The gradient echo with grid tagging images at end-systole in four (left) and three-chamber (right) view demonstrating normal segmental contraction.

**Figure 4**
Tissue characterization 4- and 3-chamber view demonstrating myocardial fibrosis (arrow).

*The information about this product is preliminary. The product is under development. It is not commercially available in the US and its future availability cannot be assured.*
Case Report MAGNETOM Sonata
Hypertrophic Cardiomyopathy and Stress Perfusion

Paulo R. Schwartzman, M.D., Ph.D.
Escola Paulista de Medicina, Unifesp, São Paulo, Brazil

Patient history
A 38-year-old female complained of chest pain associated with exercise, which resolved after 2–3 min of rest. Echo demonstrated an asymmetrical septal hypertrophy with no left ventricular outflow obstruction. A stress nuclear test was performed and no perfusion deficit was detected. Stress perfusion cardiac magnetic resonance was requested to define if chest pain was secondary to a perfusion deficit.

Stress MR protocol
The cardiac magnetic resonance exam protocol was as follows:
1. Cine-images on three long-axis views (two, three and four chamber view) and three short axis-views (base, mid-ventricular and apical).
2. Dipyridamole stress perfusion images in four short axis slices acquired 2 min after a 4 min dipyridamole injection and CM.
3. Short axis TrueFISP cine images were obtained from base to apex to define LV volumes.
4. Rest perfusion on the same four short axis views as the slices of stress perfusion.
5. Tissue characterization images acquired 15 min after CM on the same image plane as previous long axis and short-axis views to define the presence of myocardial fibrosis.

Imaging findings
The TrueFISP cine images in the four-chamber view (Fig. 1) and short-axis view (Fig. 2) documented the extensive (32 mm – arrowhead) and asymmetrical (arrow) hypertrophy. The stress perfusion* short-axis image (Fig. 3, left) demonstrates a septal perfusion deficit (arrow) during the gadolinium injection. The rest perfusion image demonstrates normal (arrow) perfusion (Fig. 3, right). Tissue characterization image* after CM in the 4 chamber view (Fig. 4, left) and short-axis view (Fig. 4, right) demonstrate the septal hypertrophy and absence of myocardial fibrosis.
Discussion

The characteristic finding of inappropriate myocardial hypertrophy in the absence of an obvious cause for the hypertrophy is the marker for hypertrophic cardiomyopathy. This is a genetic disease associated with eight different genes, and frequently it does not present with symptoms. Often the diagnosis is made by a routine echo. Several patients may complain of chest pain and dyspnea, despite normal coronary arteries. The perfusion deficit with normal coronary arteries is secondary to a disproportion of extensive hypertrophy and normal coronary flow.

Nuclear stress testing is usually performed to rule out coronary artery disease in patients with chest pain and hypertrophic cardiomyopathy. However due to suboptimal spatial resolution, smaller perfusion deficits may not be detected. This case represents an application of cardiac magnetic resonance in patients with chest pain and normal stress imaging test. The exercise-related chest pain in this young female patient with asymmetrical septal cardiomyopathy, normal coronary arteries, normal nuclear stress testing – but an abnormal perfusion deficit – was properly diagnosed by cardiac magnetic resonance. This deficit is explained by an imbalance of severe hypertrophy and normal myocardial perfusion.

*The information about this product is preliminary.
The product is under development. It is not commercially available in the US and its future availability cannot be assured.
Case Report MAGNETOM Sonata Left Subclavian Artery Occlusion*

Paulo R. Schvartzman, M.D., Ph.D.
Escola Paulista de Medicina, Unifesp, São Paulo, Brazil

Patient History
51-year-old woman complains of left arm pain, which lasts for 15–20 minutes. The pain is increased with arm exercise movements and decreases with rest. At examination, there is a slight decrease on radial pulse (3+/4) compared to right radial pulse (4+/4). A doppler was performed and suggested decreased flow to left arm. Magnetic resonance angiography (MRA) was requested to define the presence or absence of arterial stenosis.

Imaging findings
An MRA was performed on a Siemens MAGNETOM Sonata using care-bolus technique (Fig. 1) followed by a 3-dimensional (3-D) coronal MRA acquisition (Fig. 2). The 3-D acquisition consisted of a coronal slab positioned on the middle of the thorax with 80 slices of 1.5 mm thickness. Subtraction was performed (Fig. 3) between the images with and without contrast, documenting the proximal left subclavian artery occlusion. Also the mid and distal flow filling due to collaterals was confirmed. Thick (Fig. 4) and thin (Fig. 5) maximal intensity projections (MIP) were performed confirming the left subclavian artery occlusion.

Discussion
Magnetic resonance angiography has been used routinely for several years, but recently with the improvement in MRA hardware and software, the results have been more reliable. The goal of MR angiography is to detect vascular abnormalities non-invasively. This case represents a nice example of how MR angiography can be utilized to confirm or exclude vascular pathologies.

*The information about this product is preliminary. The product is under development. It is not commercially available in the US and its future availability cannot be assured.
[Figure 3] Raw data MR Angiography image shows the subclavian artery stenosis.

[Figure 4] MIP image shows the subclavian artery stenosis.

[Figure 5] Raw data MR Angiography images showing the subclavian artery stenosis.
Special Offer: Cardiovascular

Cardiovascular MR imaging is increasingly becoming a part of the daily routine of MR users. The exciting applications includes, for example, cardiac function, flow analysis, morphologic diagnosis, tissue characterization and angiographic information.

Siemens offers a wide range of software and hardware options for cardiovascular MR. And from now through June 2007 these are available to our existing MAGNETOM customers at special promotion conditions.

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Or send an e-mail to magnetomworld@siemens.com to learn more about this promotion.

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- Advanced Cardiac
- Flow Quantification
- Argus Flow

Vascular
- syngo Vessel View
- syngo 3D VRT (Volume Rendering Technique)

**MAGNETOM Espree**

Cardiac Options
- PMU Wireless
- Physio Control
- Advanced Cardiac
- Flow Quantification
- Argus Flow
- Advanced High Order Shim

Vascular
- syngo Vessel View
- syngo 3D VRT (Volume Rendering Technique)

**MAGNETOM Symphony, A Tim System**

Cardiac Options
- PMU Wireless
- Physio Control
- Advanced Cardiac
- Flow Quantification
- Argus Flow

Vascular
- syngo Vessel View
- syngo 3D VRT (Volume Rendering Technique)

**MAGNETOM Trio, A Tim System**

Cardiac Options
- PMU Wireless
- Physio Control
- Advanced Cardiac
- Flow Quantification
- Argus Flow

Vascular
- syngo Vessel View
- syngo 3D VRT (Volume Rendering Technique)
### MAGNETOM Harmony

**Cardiac Options**
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- syngo Advanced Cardiac
- Flow Quantification
- Argus Flow
- Advanced High Order Shim

**Vascular**
- syngo Vessel View
- syngo 3D VRT (Volume Rendering Technique)
- syngo Fly Through
- Peripheral Angiography
- Matrix coil

- Cardiac MR Pack SW
- Cardiac MR Pack HW

**Cardiac Options**
- Argus Function
- Argus Dynamic Signal
- Interactive Real Time

**Vascular**
- syngo Panoramic Table
- syngo Advanced Angio Pack
- syngo Core Bolus
- Interact. RM. Imaging

### MAGNETOM Sonata

**Cardiac Options**
- PMU Electronics
- syngo Advanced Cardiac
- Flow Quantification
- Argus Flow
- Advanced High Order Shim

**Vascular**
- syngo Vessel View
- syngo 3D VRT (Volume Rendering Technique)
- syngo Fly Through
- Peripheral Angiography
- Matrix coil

- Cardiac MR Pack SW
- Cardiac MR Pack HW

**Cardiac Options**
- Argus Function
- Argus Dynamic Signal
- Interactive Real Time

**Vascular**
- syngo Panoramic Table
- syngo Advanced Angio Pack
- syngo Core Bolus
- Large FoV Adapter
- Interact. RM. Imaging

### MAGNETOM Symphony

**Cardiac Options**
- PMU Electronics
- syngo Advanced Cardiac
- Flow Quantification
- Argus Flow
- Advanced High Order Shim

**Vascular**
- syngo Vessel View
- syngo 3D VRT (Volume Rendering Technique)
- syngo Fly Through
- Peripheral Angiography
- Matrix coil

- Cardiac MR Pack SW
- Cardiac MR Pack HW
- Active ECG Electrodes

**Cardiac Options**
- Argus Function
- Argus Dynamic Signal
- Interactive Real Time

**Vascular**
- syngo Panoramic Table
- syngo Advanced Angio Pack
- syngo Core Bolus
- Large FoV Adapter
- Interact. RM. Imaging

### MAGNETOM Trio

**Cardiac Options**
- PMU Electronics
- Advanced Cardiac
- syngo Flow Quantification
- Argus Flow
- Advanced High Order Shim

**Vascular**
- syngo Vessel View
- syngo 3D VRT (Volume Rendering Technique)

- Cardiac MR Pack SW
- Cardiac MR Pack HW
- Active ECG Electrodes

**Cardiac Options**
- Argus Function
- Argus Dynamic Signal
- Interactive Real Time

**Vascular**
- syngo Fly Through
- Peripheral Angiography
- Matrix coil

- Exam Room
- PMU Display

- CP Body Array Flex
Prior to allowing patients and other individuals into the MRI environment, careful screening is performed to determine the presence of implants or devices that may pose hazards or risks [1]. Once an implant or device is identified, the labeling for a given object is examined and a decision is made with regard to whether or not it is acceptable for the patient to undergo an MRI procedure or for an individual to enter the MRI environment.

“Old” Terminology

The terminology applied to implants and devices relative to the MRI environment has evolved over the years. In 1997, the Food and Drug Administration, Center for Devices and Radiological Health, proposed definitions for the terms “MR Safe” and “MR Compatible” [2]. These terms are defined as follows:

**MR Safe** – the device, when used in the MRI environment, has been demonstrated to present no additional risk to the patient or other individual, but may affect the quality of the diagnostic information. The MRI conditions in which the device was tested should be specified in conjunction with the term MR safe since a device which is safe under one set of conditions may not be found to be so under more extreme MRI conditions.

**MR Compatible** – a device shall be considered “MR compatible” if it is MR safe and the device, when used in the MRI environment, has been demonstrated to present no additional risk to the patient or other individual, but may affect the quality of the diagnostic information. The MRI conditions in which the device was tested should be specified in conjunction with the term MR compatible since a device which is compatible under one set of conditions may not be found to be so under more extreme MRI conditions.

Using this terminology, MR safety testing of an implant or object involved assessments of magnetic field interactions, heating, and, in some cases, induced electrical currents while MR compatibility testing required all of these as well as characterization of artifacts. In addition, it may have been necessary to evaluate the impact of various MRI conditions on the functional or operational aspects of an implant or device.

“New” Terminology

Over the years, manufacturers generally used the terms MR safe and MR compatible to label medical devices. However, in time it became apparent that these terms were confusing and were often used interchangeably or incorrectly [3]. Therefore, in an effort to clarify the terminology and, more importantly, because the misuse of these terms could result in serious accidents for patients and other individuals, the MRI task group of the American Society for Testing and Materials (ASTM) International developed a new set of terms with associated icons [4]. The new terms, MR Safe, MR Conditional, and MR Unsafe are defined by the ASTM document [4], as follows.

**MR Safe** – an item that poses no known hazards in all MRI environments. Using the new terminology, “MR safe” items include non-conducting, non-metallic, non-magnetic items such as a plastic Petri dish. An item may be determined to be MR Safe by providing a scientifically based rationale rather than test data.

**MR Conditional** – an item that has been demonstrated to pose no known hazards in a specified MRI environment with specified conditions of use. Field conditions that define the MRI environment include static magnetic field strength, spatial gradient, dB/dt (time varying magnetic fields), radio frequency (RF) fields, and specific absorption rate (SAR). Additional conditions, including specific configurations of the item (e.g., the routing of leads used for a neurostimulation system), may be required.

For MR Conditional items, the item labeling includes results of testing sufficient to characterize the behavior of the item in the MRI environment. In particular, testing for items that may be placed in the MRI environment should address magnetically induced displacement force and torque, and RF heating. Other possible safety issues include but are not limited to, thermal injury, induced currents/voltages, electromagnetic compatibility, neurostimulation, acoustic noise, interaction among devices, and the safe functioning of the item and the safe operation of the MR system. Any parameter that affects the safety of the item should be listed and any condition that is known to produce an unsafe condition must be described.
References

MR Unsafe – an item that is known to pose hazards in all MRI environments. MR Unsafe items include magnetic items such as a pair of ferromagnetic scissors.

In addition to the new terms, the ASTM document introduced corresponding icons, consistent with international standards for colors and shapes of safety signs. The icons are intended to be used on items that may be brought into or near the MRI environment as well as in product labeling. The icons may be reproduced in color or in black and white, however the use of color is encouraged because of the added visibility. The “MR unsafe” icon consists of the letters ‘MR’ in black on a white field inside a red circle with a diagonal red band.

For MR Conditional items, the item labeling must include results of testing sufficient to characterize the behavior of the item in the MRI environment.

Notably, the new terminology is not being applied retrospectively to implants and devices that have already received FDA approved labeling using the terms “MR safe” or “MR compatible”.

Hopefully, the utilization of this new terminology will serve to elucidate matters related to biomedical implants and devices and help to ensure the safe use of MRI technology.

Refurbished MRI Systems Now Available through the Proven Excellence Program

The Refurbished Systems division in the USA recently incorporated MRI into its Proven Excellence product portfolio. Proven Excellence (PE) is Siemens’ multi-modality medical equipment refurbishment program. The MAGNETOM Symphony, Harmony, and Concerto are proven products from our MRI equipment portfolio that can meet your everyday clinical and technological demands. These systems are available at an affordable price point through our Proven Excellence Program.

The program starts with an initial inspection of each system, taking into consideration age, condition, service history, performance and upgradeability of software and hardware. After a professional de-installation and transportation to our Siemens Refurbishing Systems facility, the system is thoroughly cleaned, disinfected and painted, worn parts are replaced with original spare parts and applicable software updates are performed. After passing a complete system check with original test equipment and procedures, the system is rewarded with our Proven Excellence quality seal. This seal represents the fulfillment of the strict specifications of relevant international norms and standards as well as security regulations.

After transportation to the customer site, the system is installed by our Siemens installation and service team of trained technicians.

With the Proven Excellence Program, Siemens Medical Solutions also offers its customers various flexible financing solutions and service contracts as well as a warranty typically equivalent to that of new systems. Moreover, spare parts are generally available for 5 years.
Patients with coronary artery disease are often treated by percutaneous transluminal coronary angioplasty (PTCA). Rennarrowing at the angioplasty site, or restenosis, occurs in as many as 50% of patients following PTCA. Therefore, after coronary artery intervention, either a bare metal or drug eluting stent is placed in an effort to prevent restenosis. There is considerable attention focused on the use of drug eluting stents to prevent coronary artery restenosis that tends to occur in a substantial number of patients following stenting with "bare" devices. Studies have reported that drug eluting stents reduce the incidence of target vessel failure compared to uncoated metallic stents. As such, drug eluting stents are now used on a widespread basis (upwards of 80%) in patients with coronary artery disease.

Recently, MR safety information has been obtained for several bare wire and drug eluting coronary stents, which have been reported to be safe for patients undergoing MR procedures at 3 Tesla or less (i.e., based on assessments of magnetic field interactions and MRI-related heating). These coronary artery stents include the following:

**Endeavor Drug Eluting Coronary Artery Stent (Medtronic Vascular)** – Through non-clinical testing, the Endeavor stent has been shown to be MRI safe at field strengths of 3 Tesla or less, and a maximum whole body averaged specific absorption rate (SAR) of 2.0 W/kg for 15 minutes of MRI. The Endeavor stent should not migrate in this MRI environment. MRI at 3T or less may be performed immediately following the implantation of the Endeavor stent. Non-clinical testing has not been performed to rule out the possibility of stent migration at field strengths higher than 3 Tesla.

In this testing, the stent produced a maximum temperature rise of 0.5 degrees C at a maximum whole body averaged SAR of 2.0 W/kg for 15 minutes of MRI. The temperature rise was observed to be similar for comparable bare MR imaging quality may be compromised if the area of interest is in the exact same area or relatively close to the position of the stent.

**TAXUS Express Paclitaxel-Eluting Coronary Stent (Boston Scientific Corporation)** – Through non-clinical testing, the TAXUS Express stent has been shown to be MRI safe at field strengths of 3 Tesla or less, and a maximum whole body averaged specific absorption rate (SAR) of 2.0 W/kg for 15 minutes of MRI. The TAXUS Express stent should not migrate in this MRI environment. MRI at 3T or less may be performed immediately following the implantation of the TAXUS Express stent. Non-clinical testing has not been performed to rule out the possibility of stent migration at field strengths higher than 3 Tesla.

In this testing, the stent produced a maximum temperature rise of 0.65 degrees C at a maximum whole body averaged SAR of 2.0 W/kg for 15 minutes of MRI. The effect of heating in the MRI environment was similar for overlapping bare metal stents (2 to 5-mm overlap at the ends), made of the same stainless steel material and having the same stent design. The effect of heating in the MRI environment on stents with fractured struts is not known. The temperature rise of 0.65 degrees C for 15 minutes is calculated to result in an increase in cumulative drug release of 0.001% of the total dose.

**Liberté Coronary Artery Stent (bare metal coronary artery stent, Boston Scientific Corporation)** – The Liberté Stent has been shown to be MR safe at field strengths of 3 Tesla or less, and a maximum whole body averaged specific absorption rate (SAR) of 2.0 W/kg for 15 minutes of MRI. The Liberté Stent should not migrate in this MRI environment. MR imaging at 3T or less may be performed immediately following the implantation of the Liberté Stent. Non-clinical testing has not been performed to rule out the possibility of stent migration at field strengths higher than 3 Tesla.

In this testing, the stent produced a maximum temperature rise of 0.65 degrees C at a maximum whole body averaged SAR of 2 W/kg for 15 minutes of MR imaging. The temperature rise was observed to be similar for comparable bare
metal overlapping stents (2 to 5 mm overlap at the ends). Heating has not been determined for fractured struts. MR imaging quality may be compromised if the area of interest is in the exact same area or relatively close to the position of the stent. This stent has not been evaluated to determine if it is safe in MRI systems with field strengths greater than 3T.

**TAXUS Liberté Paclitaxel-Eluting Coronary Stent (Boston Scientific Corporation)** – Through non-clinical testing, the TAXUS Liberté stent has been shown to be MRI safe at field strengths of 3 Tesla or less, and a maximum whole body averaged specific absorption rate (SAR) of 2.0 W/kg for 15 minutes of MRI. The TAXUS Liberté stent should not migrate in this MRI environment. MRI at 3T or less may be performed immediately following the implantation of the TAXUS Liberté stent. Non-clinical testing has not been performed to rule out the possibility of stent migration at field strengths higher than 3 Tesla. In this testing, the stent produced a maximum temperature rise of 0.65 degrees C at a maximum whole body averaged SAR of 2.0 W/kg for 15 minutes of MRI. The effect of heating in the MRI environment was similar for overlapping bare metal stents (2 to 5 mm overlap at the ends), made of the same stainless steel material and having the same stent design. The effect of heating in the MRI environment on stents with fractured struts is not known. The temperature rise of 0.65 degrees C for 15 minutes is calculated to result in an increase in cumulative drug release of 0.001% of the total dose. MR imaging quality may be compromised if the area of interest is in the exact same area or relatively close to the position of the stent.

**CYPHER Sirolimus-eluting Coronary Stent (Cordis Corporation/Johnson and Johnson)** – Through non-clinical testing, single and two overlapping CYPHER Stents have been shown to be MRI safe at field strengths of 3 Tesla or less, and a maximum whole body averaged specific absorption rate (SAR) of 4.0 W/kg for 15 minutes of MRI. Single and two overlapping CYPHER Stents should not migrate in this MRI environment. Non-clinical testing has not been performed to rule out the possibility of stent migration at field strengths higher than 3 Tesla. In this testing, single CYPHER Stents up to 33 mm in length produced a temperature rise of less than 1 degree C, and two overlapped 33 mm length CYPHER Stents produced a temperature rise of less than 2 degrees C at a maximum whole body averaged specific absorption rate (SAR) of 4.0 W/kg for 15 minutes of MRI. The effect of heating in the MRI environment for stents with fractured struts is not known. The effect of heating in the MRI environment on the drug or polymer coating is not known. MR imaging quality may be compromised if the area of interest is in the exact same area or relatively close to the position of the stent.

Note: “TAXUS” is the trademark name on the drug coating, and refers to the drug eluting coating addition to the bare metal stent. As such, when there is the addition of the drug eluting coating, it will be referred to as the TAXUS and then the specific name of the stent. For example, TAXUS Express stent. The bare metal stent does not contain the TAXUS prefix.

Note: This statement applies to all currently marketed CYPHER Stents in the United States.

References
The Siemens MR application development group initiated the 2nd Pediatric User Meeting.
The meeting, held in Philadelphia, Pennsylvania, USA, was attended by approximately 50 internationally-known pediatric radiologists and experienced technologists from the US, Germany, Great Britain, the Netherlands, Belgium, Austria and Australia.
The whole of the first day (Saturday) was dedicated to clinically focused presentations from many MAGNETOM users.
Contributions were grouped into the following sub-topics:

- Neuro Imaging
- Neuro Spectroscopy and Neuro Research
- Cardiac Imaging
- Body Imaging
- Orthopedic Imaging
- Fetal Imaging

The presentations gave a very distinct overview on the range of clinical questions that are answered by the high performance MR scanners of today. The participants use MAGNETOM Symphony, MAGNETOM Avanto and MAGNETOM Trio. The final presentation on fetal imaging* – a domain for ultra sound – ranged from a demonstration of the capabilities of this established technique right through to advanced 3D capabilities.

Sunday morning started with presentations on current developments at Siemens with respect to the upcoming software version syngo MR B13. The focus was on Clinical Applications, Advanced Neuro, Spectroscopy and Cardiac Imaging. In addition, Advanced Imaging Research Inc. (located in Cleveland) presented their coils and support tools for scanning of neonates.

The Sunday morning session was followed by three parallel working group meetings on Neuro, Body/Ortho and Cardiac applications. Each of the three working groups discussed in detail the current clinical routine, future use and clinical and technical requirements to address future applications. The topics ranged from clinical questions, hardware and advanced sequences to new approaches in image post-processing.

The two days spent together at the Pediatric User Meeting resulted in a renewing of old contacts and in creating new contacts between users, in valuable feedback and fruitful input to the Siemens development groups. Siemens already offers on Tim (Total imaging matrix) systems a set of pediatric optimized, age dependent protocols (Pediatric Suite) and will use this input to further facilitate the work of clinicians and MR technologists doing pediatric examinations.

We would like to thank all participants and clinical partners for their contributions, either in the presentations they gave or their input to discussions.

*The safety of imaging fetuses/infants has not been established.
Pediatric imaging

MR’s safety due to lack of x-ray is one of the major factors making MR the examination of choice for pediatric patients. In recent years MR imaging with Tim (Total imaging matrix) has created major developments in this area: BLADE has helped decrease motion artefacts, GRAPPA has dramatically improved imaging speed and sequences like TrueFISP fatsat have been extremely useful for state-of-the-art skeletal imaging.

[Figure 1]  MAGNETOM Trio, A Tim System T2 TSE coronal, GRAPPA 2, 5-year-old.

| TR/TE     | 5780/103 |
| TA        | 2:48 min |
| SL        | 3 mm     |
| slices    | 27       |
| FoV       | 200 mm   |
| matrix    | 512      |

Courtesy of University of North Carolina, Chapel Hill, USA

[Figure 2A]  MAGNETOM Avanto T1 SE sagittal, GRAPPA 2, without motion correction, 12-year-old.

| TR/TE     | 700/11  |
| TA        | 2:31 min|
| SL        | 5 mm    |
| slices    | 24      |
| FoV       | 200 mm  |
| matrix    | 256     |

[Figure 2B]  MAGNETOM Avanto, T1 TIRM sagittal, with BLADE, 12-year-old.

| TR/TE     | 1970/55 |
| TI        | 860     |
| TA        | 5:06 min|
| SL        | 5 mm    |
| slices    | 21      |
| FoV       | 200 mm  |
| matrix    | 256     |

Courtesy of Children’s Hospital, Denver, CO, USA
[Figure 3] MAGNETOM Symphony
T1 SE sagittal, post contrast, 4-year-old, ependymoma

- TR/TE: 600/17
- TA: 3:30 min
- SL: 3 mm
- Slices: 11
- FoV: 230 mm
- Matrix: 256

Courtesy of UZ Gent MR 1, Gent, Belgium

[Figure 4] MAGNETOM Avanto
T2 HASTE transversal, GRAPPA 2, 9-year-old, poly cystic kidneys

- TR/TE: 199950/102
- TA: 3:21 min
- SL: 5 mm
- Slices: 38
- FoV: 260 mm
- Matrix: 256

Courtesy of Children's Hospital, Denver, USA

[Figure 5] MAGNETOM Espree
T2 3D TrueFISP sagittal water excitation, 7-year-old, osteochondrosis dissecans

- TR/TE: 9.5/3.5
- TA: 4:17 min
- eff. SL: 1.5 mm
- partitions: 48
- FoV: 160 mm
- Matrix: 384

Courtesy of Klinikum Bremen-Mitte, Bremen, Germany