Comparison of amniotic fluid volumetry between fetal sonography and MRI - Correlation to MR diffusion parameters of the fetal kidney

Elena Moschos1 *, Daniel Güllmar2 , Anja Fiedler1 , Ulrike John4 , Diane M. Renz1 , Matthias Waginger1 , Ekkehard Schleussner1 , Dietmar Schlembach1 , Uwe Schneider3 and Hans-Joachim Mentzel1

1 Institute of Diagnostic and Interventional Radiology, Section of Pediatric Radiology, University Hospital Jena, Germany
2 Institute of Diagnostic and Interventional Radiology, Medical Physics Group, University Hospital Jena, Germany
3 Department of Obstetrics, University Hospital Jena, Germany
4 Department of Pediatrics, Section of Pediatric Nephrology, University Hospital Jena, Germany
5 Clinic of Obstetrics, Vivantes Hospital Berlin-Neukoelln, Germany

Abstract

Objective: Determine whether amniotic fluid volume (AFV) determination by magnetic resonance imaging (MRI) can be compared with ultrasound based amniotic fluid index (AFI) and whether diffusion weighted imaging (DWI) of the fetal kidneys can be used as surrogate marker for the assessment of AFV and fetal kidney function.

Methods: In the period from August 2013 to September 2014 prospective evaluation of AFV based on TRUFI MRI in 2 spatial planes (AFVMRI) and ultrasound based AFI. Inclusion criteria for the evaluation were: 1) Full image of the fetus (including AF and placenta) in an MR-study in at least two planes (sagittal and coronal), 2) Layer thickness 8 mm, 3) DWI-images of the fetal kidney in two planes (sagittal and transverse to the fetus). ROI analysis of the ADC values in normal kidneys and in pathological fetal kidneys (CAKUT) as well as the correlation of AFVMRI vs. AFI, renal volume and ADC were performed.

Results: The correlation between AFI (13.60 ± 2.46 cm) and AFVMRI (579 ± 353 mL) was 0.69 (p=0.03). There was no significant correlation between AFVMRI and kidney volume, between AFVMRI and ADC700 and ADC800. The B700-ADC values of fetuses with CAKUT (n=14) were considerably higher than the values in fetuses with healthy kidneys.

Conclusion: Compared to AFI, AFVMRI showed higher accuracy, but was associated with higher time consumption. Thus it cannot be routinely recommended for the purpose of AFV-determination alone. MRI can be used to establish a relationship between kidney diffusion parameters, renal volume, GA and AFV in fetuses with CAKUT.


Introduction

The amniotic fluid (AF) plays an important role during pregnancy [1,2]. Its determination is an essential part of prenatal diagnostics [3]. Changes in the AFV are one of the most common abnormalities diagnosed by ultrasound (US). The amount of AF is assessed during US by measurement of the Amniotic Fluid Index (AFI) or the Single Deepest Pocket (SDP). Among other pathologies congenital anomalies of the kidney and urinary tract (CAKUT) are considered to have major influence on the amniotic fluid volume (AFV). They include a wide spectrum of renal abnormalities, from mild hydrenephrosis to more severe cases, such as bilateral renal dysplasia. The etiology of the majority of cases of CAKUT remains unknown, but there is increasing evidence that genomic imbalance contributes to the pathogenesis of CAKUT [4].

For the accurate assessment of renal abnormalities, fetal MRI may be used as a further diagnostic step [5,6]. It not only determines some qualitative characteristics of the fetal kidney like the apparent diffusion coefficient (ADC) [7], but also provides a possibility for the determination of the AFV (AFVMRI). The study compared two methods of AF volumetry, and investigated the suitability of diffusion-weighted imaging (DWI) of the fetal kidney to be used as a surrogate marker of fetal glomerular density in relation to the amount of AF.

Correspondence to: Elena Moschos, Institute of Diagnostic and Interventional Radiology, Section of Pediatric Radiology, University Hospital Jena, Germany, Tel: +49 36419324925; Fax: +49 36419324832; E-mail: elena.moschos@med.uni-jena.de

Key words: magnetic resonance imaging, amniotic fluid, kidney, diffusion magnetic resonance imaging, biomarkers

Received: April 28, 2017; Accepted: May 26, 2017; Published: May 29, 2017
Materials and methods

Subjects

From August 2013 to September 2014 thirty pregnant women were investigated at a 1.5 T MRI scanner. The study protocol was approved by the local ethics committee. Informed consent was obtained from all prospectively recruited participants. All in the study included pregnant women underwent a medically indicated fetal MRI (Table 1).

Study protocols

MRI Protocol: All MR examinations were performed at 1.5 T (Magnetom Avanto, Siemens, Erlangen) using the standard body matrix coil and without application of contrast medium. All sequences listed below were included in the standard MRI protocol. They were obtained during free breathing and without sedation with the pregnant women in the supine or lateral position. For the determination of AFV a True Fast Imaging with Steady State Free Precession (TRUFI) gradient sequence was acquired in coronal and sagittal planes with the following parameters: TR 3.68 ms; TE 1.84 ms; slice thickness 8 mm; gap 12 mm in 14 cases; no gap in 16 cases; acquisition time 71; (31 slices in the sagittal plane and 21 slices in the coronal plane).

Table 1. Indications for fetal MRI.

<table>
<thead>
<tr>
<th>Number</th>
<th>Indications</th>
</tr>
</thead>
<tbody>
<tr>
<td>14</td>
<td>CAKUT</td>
</tr>
<tr>
<td>8</td>
<td>CNS – Malformation</td>
</tr>
<tr>
<td>1</td>
<td>Tuberous sclerosis complex</td>
</tr>
<tr>
<td>1</td>
<td>Planning trial of labour after cesarean section (TOLAC)</td>
</tr>
<tr>
<td>1</td>
<td>Suspected duodenal atresia</td>
</tr>
<tr>
<td>1</td>
<td>Distended fetal bowel loops</td>
</tr>
<tr>
<td>1</td>
<td>Fetal: presacral tumor mass (anterior sacral meningocele)</td>
</tr>
<tr>
<td>1</td>
<td>Left-sided diaphragmatic hernia</td>
</tr>
<tr>
<td>1</td>
<td>Fetal hydrops of unknown origin</td>
</tr>
<tr>
<td>1</td>
<td>Truncus arteriosus communis</td>
</tr>
</tbody>
</table>

### Diffusion-weighted imaging:

Diffusion parameters were evaluated with a Diffusion-Weighted-Sequence (TR 8000 ms; TE 90 ms; FOV 420 x 300 mm, matrix 192 x 192, slice thickness 8 mm; Total volume, which was counted with the ellipsoid-formula (width x depth x length x 0.523 mm) (Figure 1a, b).

Figure 1. Measurements of the normal fetal kidney (34 wk): a) width and depth in the axial plane, b) length in the sagittal plane in one of the two kidneys.

### Statistical analysis

Statistical analysis was performed using SPSS 22.0 (IBM Corporation, US). Arithmetic mean values were calculated to describe the central tendency of a frequency distribution. The dispersion of a distribution in the subject groups was measured descriptively by calculating scattering measurements. To verify the correlation of MR–tomographic volume measurements in both planes Bland-Altman plots were applied. The relationship between two characteristics in the subject groups was calculated by correlation analysis of bivariate statistics. In order to quantify the linear relationship in the subject groups, Pearson correlation coefficients were calculated.

### AF tracing:

The AFV was evaluated on TRUFI sequences using the free open source software package 3D Slicer (Version 4.2.2.1, USA). The experience of the MRI operator was less than a year. The external and internal contours of the AF were marked manually for each layer both in the sagittal and coronal planes. Where possible, the umbilical cord was not included. The contoured area was filled with color (blue for AF) (Figure 3). The tracing of each study took approximately 40–60 minutes. For the overall volume calculation, the software interpolated the boundaries in these areas.

### Fetal sonography:

After the MRI scan, 16 from overall 30 subjects underwent transabdominal fetal sonography in a supine position with the devices GE Voluson E8 Expert (GE Healthcare), GE Voluson 730 Expert (GE Healthcare) or Toshiba Apio 500 (Toshiba Medical Systems Corporation) using 3.5 MHz curved–array transducers. The experience of the US operators was more than 5 years. The AFI measurement was performed similar to the method introduced initially by Phelan et al. [8] and described later by Hinh und Ladinsky [9]. The uterus was divided into four quadrants that are defined by the umbilicus. The transducer was placed vertically in each quadrant and then the largest fluid pocket in each quadrant was measured (cm). To determine the AFI, the measurements from each quadrant were summed.
Results

In 30 cases the MRI was successfully performed. In 16/30 cases [age 30 yrs (21-39 yrs), gestational age (GA) 30 weeks (23-35wks)] the ultrasound evaluation of the AFV by AFI was performed on the same day with the MRI study (Group A). Other 14/30 patients underwent MRI alone without AFI-measurement (Group B). The GA in the Group B was 26 weeks and 2 days (19-33 wks), the age of the pregnant women 26.2 years (23-36 yrs).

The mean AFI in the Group A was 13.6 ± 2.46 cm (range 8.30 – 17.20 cm). The AFVMRI was measured in a total of 30 patients in both groups. The mean AFVMRI was 579 ± 353 mL (range 156-1732 mL) in coronal and 590 ± 335 mL (range 187–1628 mL) in sagittal plane (Figure 4).

The mean difference of AFVMRI (coronal-sagittal) was -5.09 mL (SD 36 ml), the upper and lower limits of agreement were 67 mL and -77 mL, respectively. The correlation between AFI and AFVMRI was 0.69 (p=0.03); there was a linear relationship between MR-tomographic and sonographic determined values (Figure 5). The range of renal volume obtained was 0.57–21.92 mL (Table 2).

Inclusion criteria for the evaluation were: 1) Full image of the fetus (including AF and placenta) in an MR-study in at least two planes (sagittal and coronal), 2) Layer thickness 8 mm, 3) DWI-images of the fetal kidney in two planes (sagittal and transverse to the fetus). 16 patients underwent MRI followed by the US evaluation of the AFI on the same day (group A). The other 14/30 patients underwent MRI alone without AFI-measurement (group B). Exclusion criteria for the group A were absence of AFI on the same day and for both groups inadequate image quality, absence of DWI of TRUFI sequences as well as poor orientation of the fetal kidneys in very small/rolled up fetuses.

The diffusion parameters of the fetal kidneys were measured in 30 fetuses. ADC–values of the fetal kidneys determined in 25 patients with a b-value of 700 s/mm² were on average lower (mean 1.5 x 10⁻³, min 0.67, max 2.99, SD 0.53) than in the group of 9 pregnancies measured with a b-value of 800 s/mm² (mean 1.58 x 10⁻³, min 1.22, max 2.74, SD 0.47). The average value of ROI-size was 10 mm (SD ± 6.69 mm²).

The comparison of ADC values in fetuses with and without CACTU showed that on b-value of 700 s/mm² the ADC was higher than in fetal kidneys without anomalies (Table 3).

The Pearson correlation coefficients between AFV measured both in MRI and in the sonography and diffusion parameters of the fetal kidneys for fetuses with CACTU are summarized in Table 4. There was no significant correlation between diffusivity of the fetal kidney and with both methods measured AFV in fetuses with CACTU (p>0.05).

<table>
<thead>
<tr>
<th>Kidney volume (mL)</th>
<th>N</th>
<th>Minimum</th>
<th>Maximum</th>
<th>SD</th>
</tr>
</thead>
<tbody>
<tr>
<td>KvOli_right</td>
<td>29</td>
<td>0.57</td>
<td>21.92</td>
<td>4.84</td>
</tr>
<tr>
<td>KvOli_left</td>
<td>29</td>
<td>0.57</td>
<td>20.50</td>
<td>6.13</td>
</tr>
</tbody>
</table>

Table 2. MR-tomographically measured kidney volume (n=30).

<table>
<thead>
<tr>
<th>Mean Diffusion (mm²/s)</th>
<th>CAKUT (n=14)</th>
<th>Healthy kidneys prenatal (n=16)</th>
</tr>
</thead>
<tbody>
<tr>
<td>B700ADC</td>
<td>1.03 – 2.99 x 10⁻³ (n=14)</td>
<td>Mean 1.64 x 10⁻³ SD 0.61 x 10⁻³</td>
</tr>
<tr>
<td></td>
<td>0.67 – 2.14 x 10⁻³ (n=11)</td>
<td>Mean 1.34 x 10⁻³ SD 0.37 x 10⁻³</td>
</tr>
<tr>
<td>B800ADC</td>
<td>1.30 – 1.75 x 10⁻³ (n=3)</td>
<td>Mean 1.56 x 10⁻³ SD 0.23 x 10⁻³</td>
</tr>
<tr>
<td></td>
<td>1.22 – 2.74 x 10⁻³ (n=6)</td>
<td>Mean 1.59 x 10⁻³ SD 0.58 x 10⁻³</td>
</tr>
</tbody>
</table>

Table 3. Diffusion values b=700 s/mm² ADC and b=800 s/mm²; ADC in the overall group in healthy kidneys fetuses (n=16) and fetuses with CACTU (n=14).

<table>
<thead>
<tr>
<th>Parameter</th>
<th>AFVMRI (mL)</th>
<th>AFI (cm)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>B700ADC(mm/s)</td>
<td>B700ADC(mm/s)</td>
</tr>
<tr>
<td>Pearson-correlation</td>
<td>-0.218</td>
<td>-0.594</td>
</tr>
<tr>
<td>Sig.</td>
<td>0.546</td>
<td>0.214</td>
</tr>
<tr>
<td>N</td>
<td>10</td>
<td>6</td>
</tr>
</tbody>
</table>

Table 4. Pearson correlation between the AFV measured in MRI and in the sonography and kidney – diffusivity (ADC) of different b-values in fetuses with CACTU: b=700 s/mm² and b=800 s/mm².
For fetuses with healthy kidneys the correlation data between AFV and kidney ADC was not obtained because in that group 15 fetuses presented with pathologies which can affect the AFV regardless of kidney function (Table 1).

Renal diffusion parameters in both groups were correlated to GA. With increasing GA the ADC-values in the fetal kidney decreased (Pearson correlation coefficient of -0.653 (p<0.01) when measured with b-value of 700 s/mm²). When measured with b-value of 800 s/mm² the data was not statistically significant (Pearson correlation coefficient of -0.465 (p=0.207)), which is probably due to a fewer number of patients in that group.

A correlation between the kidney diffusion parameters and total kidney volume measured in the MRI was also performed (Table 5).

The negative correlation in the Table 5 was statistically significant when measured with b-value of 700 s/mm² (p<0.05), suggesting that fetuses with reduced renal volume show higher ADC-values. This finding is in concordance with a physiological increase in kidney volume with increasing GA (Figure 6).

**Discussion**

**AFV measurements**

Fetal MRI allows measurement of AFV. AFVMRI in two planes are only very slightly different and show low SD based on the orientation. This indicates that in the future in order to determine the AFVMRI one plane will be sufficient enough.

The TRUFI sequences were selected for the AFV evaluation in the MRI. Due to the high difference in contrast, they allow a good differentiation between the bright water and other structures (fetus, placenta, uterus, umbilical cord). They can be performed with free breathing, are rapidly acquirable and have a high signal-to-noise ratio [10]. TRUFI sequences in the group B were performed with intersection gap of 12 mm, which could lead to partial volume effect [11]. In order to avoid it, achieve better image quality and thus more exact AFV evaluation TRUFI sequences in the group A were performed without intersection gap.

In the current study the manual AFV tracing has been performed. None the less, some studies on the semiautomatic MR volumetry of fetal organs, such as the fetal brain and the fetal lung have already been published [12,13]. The semitomatic method is time-saving [14]. However, AF tracing in an MR dataset, in contrast to, for example, lung

**Table 5**: Pearson correlation between the kidney volume measured in MRI and kidney – diffusivity (ADC) of different b-values: b=700 s/mm² and b=800 s/mm².

<table>
<thead>
<tr>
<th>Kidney volume (mL)</th>
<th>B700ADC (mm²/s)</th>
<th>B800ADC (mm²/s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>KVol</td>
<td>-0.416</td>
<td>-0.513</td>
</tr>
<tr>
<td>Sig.</td>
<td>0.043</td>
<td>0.158</td>
</tr>
<tr>
<td>N</td>
<td>24</td>
<td>9</td>
</tr>
</tbody>
</table>

The measured kidney volume in both groups correlated significantly with the GA with Pearson correlation coefficient of 0.673 (p=0) (Figure 6).

Two outliers were observed. They are marked in red and blue, respectively. They are accounting for one case of severe subvesical obstruction with megacystitis and enlargement of both ureters and the renal pelvis (red) and one case of bilateral pyelectasia (blue) in the fetal MRI which consequently lead to an increased kidney volume since the parenchyma could not be evaluated separately.
tracing, is a more difficult task since the AF does not possess a clearly defined shape. For this reason, no semiautomatic method was used in the present study.

The AFI-value as an equivalent of an AFV ranged from 8.30 to 17.20 cm with a mean of 13.60 cm (SD 2.46). The mean AFI of our study is slightly higher than that of Magann et al. [15], where it was 10.4 ± 4.2 cm. Nevertheless, that study also included pregnancies from 14 weeks onwards. The AFI-range in the current study is somewhat lower than that of Hebbar et al. [16], where the range was from 11.7 to 17.3 cm. This can be explained by the fact that it included women from 34 to 40 GA weeks and also multiple pregnancies. When comparing the results of our study with other data, a strong correlation between AFI and patient population or GA can be followed.

There was a statistically significant correlation between AFVMRI and AFI. The comparative study of Zaretsky et al. [17] with a similar design did not demonstrate a direct comparison between MR-tomographic and sonographic method of AF volumetry. However, it showed a correlation between AFI and AFVMRI, measured 3 hours before the Cesarean section and the AFV collected intrapartum during the Cesarean section: AFVMRI compared to AFI correlated higher with the measured during Cesarean section AFV [17]. That study also showed that AFI sensitivity declined with decreasing AFV in oligohydramnios. In the current study AFI variation between US and MRI did not change the categorization between normal AFV, reduced AFV or polyhydramnios. Another earlier study showed that AFI underestimated the AFV in oligohydramnios which consequently could lead to unnecessary labor induction [18]. The reason for this is most likely the inaccuracy of the volume estimation with the AFI, which is also confirmed by the low reproducibility of the measurement in pregnancies with low AFV [19]. That is why the application of the Single Deepest Pocket (SDP) in oligohydramnios is currently preferred [20].

Even though the use of fetal MRI to determine the amount of AF instead of the established sonographic method (AFI), is associated with high exactness of MRI, it brings a lot of methodical disadvantages of MRI over sonography. These include basic limitations of MRI such as basic limitations of MRI such as a sufficient diagnostic image quality and absence of contraindications for the MR-study. Logistic time requirement for the fetal MR-study as well its cost-effectiveness do not allow MRI to be routinely recommended for the purpose of AFV determination alone. It could be however useful in patients with sonographically diagnosed changes in the AFV. So, both in severe oligo- and polyhydramnios, it may complement to identify the cause of AFV-change and determine pregnancy outcome [21].

**Renal imaging**

In the current study the kidney volume was calculated using the formula for an ellipsoid in order to achieve a more exact comparison with the published size and volume charts [22]. However, the study of Bakker et al. [23] showed that for reliable calculation of renal size in vitro, MRI with use of the voxel-count method is preferred. Compared to the study of Michielsen et al. [2], where the MR-volumetry was performed in 240 fetuses, the range of the kidney volumes in the current study was slightly larger (Table 2). That study [2] however, included only healthy fetal kidneys, while our study involved also fetuses with CAKUT and pregnancies with earlier GA. In fetuses with CAKUT renal size is not an accurate reflect of the renal function (Beall et al. 2007). Moreover, the exact renal volume estimation in the fetal MRI using diameter measurement is only possible to a limited extent due to the small kidney size. Even though the present study allows a rough definition of the kidneys in the DWI and in the TRUFI sequences, for a volumetry a spatially higher resolution sequence (at best a 3D sequence), would have been necessary. This was not possible in the context of this study because of required longer time consumption for each MRI scan.

In most pregnancies (n = 25) the diffusivity was measured with a b-value of 700 s/mm². It was with 1.50 x 10⁻³ mm²/s on average slightly smaller than that of a b-value of 800 s/mm² (Table 3).

The other studies in which the fetal renal diffusivity was the subject of the study were performed with maximum b-values of '700 s/mm²' and minimum b-values of 0, 50, 200 and 350 s/mm², respectively [7,24-26]. With increasing b-value the signal-to-noise ratio in the image reduces and thereby the sensitivity to motion and to other artifacts increases [27,28]. Therefore, it is most likely that a signal–bias in the current study was responsible for small increase in diffusivity with b-values of 800 s/mm².

We observed a significant negative correlation between the diffusivity of the renal parenchyma and the fetal kidney volume (p<0.05) (Table 5). It was expected because it is known that with increasing GA the glomerular density increases (24). This leads to increase in the renal volume, kidney density, isotropy in the kidney and consequently to the restriction in the diffusion parameters in the parenchyma [29,30].

The study showed no statistically significant correlation between the AFVMRI/AFI and kidney diffusion when measured with b-values of 700 and 800 s/mm² in fetuses with CAKUT. Another study of 38 fetuses with hypoplastic kidneys demonstrated that in 88% of cases they developed oligohydramnios [31]. The reason for the lack of significance between fetal kidney diffusion values and AFVMRI is probably due to the relatively small number of pregnancies in the study. The diffusivity of the kidneys in fetuses with CAKUT was higher than the diffusivity of healthy fetal kidneys. The study of Chaumoitre et al. [7] showed that of total 6 fetuses (3 fetuses with nephropathy and 3 fetuses with dilatation of the urinary system) 4 fetuses had increased ADC-values with a b-value of 300 s/mm². In contrast, in another study of Savelli et al. [25], the diffusivity in 15 fetuses with CAKUT was slightly lower than in fetuses with normal renal function (b-value of 700 s/mm²). The increase in diffusion values in fetuses with CAKUT is possibly caused by changes in the glomerular density in pathological renal kidneys compared to healthy fetal kidneys. To the best of our knowledge there are currently no fetal MR-studies that have investigated these changes in fetuses with CAKUT. However, some studies on diffusivity in renal pathology in adults and children show that the diffusivity of the renal parenchyma in newborns with congenital hydrenephrosis is lower than those of fetuses with healthy kidneys [32]. ADC-values of healthy fetuses are lower than in newborns [24,32] which can be explained by the fact that the renal perfusion and the GFR are lower in fetuses and increase after birth [30], while the glomerular density is already determined at birth and thus has no influence on the diffusivity. In this study, it was shown that there is a negative correlation between the ADC-values with b-values of 700 and 800 s/mm² and the GA, which was significant for a b-value of 700 s/mm². This result is consistent with data of other studies [7,24,26], which indicates that the fetal kidney diffusivity values are generally well reproducible. We suggest that an ADC may possibly serve as an indicator of the fetal renal function. In adults, the diffusion value is already successfully used for this purpose [9,33,34].

**Limitations of the study**

The study was based on the prenatal diagnostics and it did not evaluate the fetal outcome postnatal. Further studies such as follow up of
the CAKUT fetuses are necessary in order to confirm the prenatal data and prove the concept of the glomerular density. Some of the fetuses which demonstrated no renal abnormalities on US or fetal MRI, could still have renal impairment due to their syndromes and conditions, for example those with tuberous sclerosis. Thus introduction of more homogenous healthy renal comparison group could reinforce the study outcome.

Conclusion

AFV measurement with both MRI and US demonstrated a significant correlation between the two methods. Although AFVMRI, compared to the sonographic API, showed higher accuracy, this method was associated with significantly higher time consumption. For this reason, it cannot be routinely recommended for the purpose of AFV–determination alone. One can suspect an advantage of the AFVMRI in poly- and oligohydramnios in a way that it may help to determine the underlying cause and thus pregnancy outcome. The role of AFVMRI in the fetuses without CAKUT was not explored in the current study. For this purpose, more intensive studies with larger study populations are needed.

We suggest that MRI can be used in order to establish a relationship between kidney diffusion parameters, renal volume, GA and AFV in fetuses with CAKUT. The MR-tomographically determined diffusivity can possibly serve as maturation marker, which can in accordance to GA differentiate between 'healthy' and 'diseased' fetal kidneys. More extensive research is required to secure the connection between fetal diffusivity and kidney function in order to make it prognostically relevant.

Authorship and contributorship

Byline authors: Hans-Joachim Mentzel, Daniel Güllmar, Uwe Schneider, Anja Fiedler, Dietmar Schlembach, Elena Moschos.

Contributors: Ulrike John, Diane M. Renz, Matthias Waginger, Ekkehard Schleussner.

Acknowledgements

This research was supported by the Department of Obstetrics and Institute of Diagnostic and Interventional Radiology of the University Hospital Jena. We are thankful to our colleagues PD Dr. Ulrike John, PD Dr. Diane M. Renz, Dr. Matthias Waginger who provided expertise that greatly assisted the research. We are also grateful to Prof. Dr. Ekkehard Schleussner for making substantial contribution to conception of the study.

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