Whole-Body-MR-Diffusion Weighted Imaging in Oncology
Ganzkörper-MR-Diffusionsbildgebung in der Onkologie

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Introduction

For about the last 15 years, high-resolution whole body MRI has yielded relevant information on local and systemic growth in oncological diagnostic testing when employed over the course of examination [1 – 3]. In this context, diffusion imaging initially played no significant role.

The development of rapid echoplanar sequences (EPI), more powerful and faster gradient...
systems, parallel acquisition techniques (PAT) with multichannel coils and receiver technology facilitated a more robust image acquisition in the trunk [5]. In 2004 Takahara et al. made a pioneering contribution to the development of whole-body diffusion-weighted imaging, by introducing a technology called DWIBS (diffusion-weighted whole-body imaging with background suppression), the acquisition of which can be performed during free respiration, that suppresses background signals and allows examination of the whole body within 25 minutes [6]. In the meantime, this technique has become viable for broad clinical application, since the appropriate protocols can be implemented on the majority of MRI systems [7]. Whole-body DWI supplements the conventional whole-body MRI sequence protocol for various tumor entities, where the accuracy depends on the particular tumor entity in some cases:

1. Diffusion-weighted images with high b-values have a high tumor-to-normal tissue contrast, thereby improving the detectability of tumor lesions as well as the infiltrative spread thereof.

2. Diffusion-weighted images and parameter cards of the diffusion constants provide clues about tissue architecture such as e.g. cell density and cellular membrane integrity and correlate with grading [8]. More in-depth information on the proliferation activity and micromilieu for the purpose of individual tumor characterization can be gathered from additional measurements of tumor blood supply by means of dynamic contrast-enhanced MRI (DCE) and – particularly in the case of brain tumors and prostate carcinoma – metabolism by means of MR spectroscopy.

3. DWI can be consulted as predictive and quantitative biomarker for individualized oncological therapy and constitutes an important diagnostic tool for testing new active substances (targeted drugs) [9]. ADC provides a quantitative progression parameter for monitoring therapy. Additionally, when frequent follow-up examinations are performed, it is advantageous to forego the use of ionizing rays and intravenous contrast agents, which has no effect particularly when examining young patients who require frequent follow-up and on kidney failure patients in the course of frequent therapy monitoring or for early recurrence testing of potentially tumor-free patients [10].

4. For better overview, 3D images can be generated [11] that provide similar imaging results as nuclear medicine procedures. In this way, DWI also supports the feasibility of full body MRI, since economics dictate that the time and human resources required for examination, evaluation and reporting must be kept in check without, however, compromising the quality of the examination results.

This survey article explains the mechanisms of the appearance of contrast and focuses on the current clinical uses of whole-body DWI for the systematic diagnosing of tumor spread, without addressing in detail the individual organ systems already examined in numerous original articles.

### Measuring diffusion with NMR

Owing to their thermal energy, water molecules are in constant motion (Brown’s molecular motion). The scope of movement in the body is tissue-specific and is ascertained by means of free length of travel or the diffusion constant. The principle of measuring diffusion by means of diffusion-weighted MRI involves first dephasing and then rephasing the transverse magnetization of excited water protons along diffusion gradients. Overall, the net magnetization of stationary water protons experiences no change, while that of protons in motion is decreased due to residual dephasing. The strength of signal loss in this process is function of the free length of travel of the water molecules in the particular tissue.

Precise mathematical computation of the complex diffusion processes in tissue and the measurement thereof using MR is possible only within limits and does not allow a simple relationship to be derived between, for example, cell density and diffusion constant. Ideal free diffusion in liquids can be mathematically described using the free diffusion constant $D_0$. In the case of free diffusion, the signal decreases according to $S=S_0 \exp(-bD_0)$, where $S$ represents the signal with diffusion gradients and $S_0$ the starting signal. The $b$-value is provided by the gradient amplitude $G$ and the temporal gradient profile (expressed by $\Delta$ and $\delta$). The free parameter $D_0$ can be computed from $S$ and $S_0$, by assuming a monoeponential drop in signal. If, however, there are restrictions caused by cell walls, then the signal decreases as $S=S_0 \exp(-bD_{app}(t))$ if $b$-values are low (roughly $8000$ s/$\text{mm}^2$). In this case, $D_{app}$ is what is known as the “Apparent Diffusion Coefficient” (ADC), the value of which is roughly $1\mu\text{m}^2$/ms in human tissue. It should be kept in mind, however, that $D_{app}(t)$ is different from $D(t)$, since the diffusion gradients are ultimately long. It should also be kept in mind that $D_{app}(t)$ is a function of the temporal course of the gradient profile. For example, the same $b$-value can be generated with, for example, a short $\Delta$ or a very long $\Delta$, if $\delta$ is changed. $D_{app}(t)$ is generally independent of these parameters, however. Despite this fact, $D_{app}(t)$ or ADC can usually be regarded as a good approximation for $D(t)$ [12].

### Acquisition, contrast mechanisms and image interpretation

#### Acquisition

Whole-body DWI can be performed on any MRI system with a field strength of at least $1.5\text{T}$. While diffusion imaging has proven to be more reliable at $1.5\text{T}$ than $3\text{T}$ owing to the more homogeneous fat suppression in a wider field of view (FOV) and the lower occurrence of susceptibility artifacts, there is potential for improvement in the signal-to-noise-ratio (SNR) at $3\text{T}$. Using ultra-fast echo-planar-imaging (EPI) sequences with an echo time $<80\text{ms}$ facilitates a favorable signal-to-noise ratio. Either integrated body coils or whole body surface coils can be used for signal reception. Using whole body surface coils can allow a scan area of approximately $200\text{cm}$ at higher image quality without requiring the repositioning of the patient or coil system during the examination. Examination is performed during free respiration with as many table positions as necessary to image the body from the top of the head to mid-thigh. Table 1 provides an overview of the sequence parameters for single-shot-spin-echo planar MR imaging (EPI) performed during free-respiration [13]. The acquired diffusion-weighted images with high $b$-values are usually reconstructed in...
Contrast mechanisms and image interpretation

Contrast depends on the selection of b-values during image acquisition, since the contrast intensity increases when higher b-values are selected owing to an improved contrast-to-noise-ratio (CNR). At the same time, selection of the maximum b-value is limited by the signal-to-noise ratio. In the case of oncological patients, the visual assessment of images with high b-values is crucial for detecting and, in some cases, characterizing lesions associated with tumors. What is known as the T2-shine-through-effect constitutes the problem that describes the relationship between signal intensity and transversal T2 relaxation time. When b-values are low, structures especially with long T2 time pose a potential source of errors. The simplest way of preventing misdiagnoses due to the T2-shine-through effect is acquiring multiple b-values to facilitate a quantitative evaluation. For this purpose, it is necessary to acquire at least two b-values, the corresponding signal amplitudes thereof must be adapted to the behavior of a simple mono-exponential curve [14]. This procedure allows the computation of an ADC value (apparent diffusion coefficient) that is independent of the magnetic field strength and the T2-shine-through-effect. An evaluation can be performed on the basis of either a region of interest (ROI), a volume of interest (VOI) or a pixel-for-pixel analysis [15]. There is additionally the risk of generating false-positive findings in movement-dependent anatomical regions such as the mediastinal structures including the hili, the portions of the left hepatic lobe located in close topographical relation to the heart as well the ventral thoracic chest wall and regions in immediate proximity to the air interfaces such as in the neck and shoulder region.

Normal image

To be able to use whole-body diffusion imaging diagnostically for tumor detection and therapy monitoring, familiarity with the normal images of soft tissue and bone marrow is imperative. Findings yielded by DWI should be correlated with conventional sequences. As Fig. 1 shows, several structures such as the CNS, the salivary glands, the spleen, the lymph node stations as well as the organs of the urinary tract and, depending on the female patient’s cycle, the normal female breast parenchyma as well as hepatic cysts and hemangiomas exhibit a physiologically induced T2-shine-through-effect as a result of a long T2-time and thus appear hyperintense in diffusion-weighted images [16]. If the water content of a lesion increases following a clearing reaction of dead cells or due to tumor necrosis it would appear as hyperintense owing to the T2-shine-through-effect in low b-image. Post-interventional changes to the tissue such as interstitial edema, surgical interventions and inflammatory reactions following irradiation likewise have a hyperintense appearance at low b-values and can thus result in false positive findings. The variability of the distribution of red and yellow bone marrow can be excellently visualized using this method. In diffusion weighted images, yellow bone marrow has a lower signal than red bone marrow due to its lower cellularity [17]. Red bone marrow, in contrast, emits a more hyperintense signal owing to its higher cell density [18]. The atrophy of red bone marrow above the age of 40, especially in women (apparently due to the estrogen deficit and osteoporosis), and obesity cause signal intensity to drop as age increases. From an oncological viewpoint the effect of cytotoxic chemotherapies plays an important role, since, owing to the decreasing cellularity of the bone marrow, they bring about a drop in signal as a result of the cytotoxic effect with subsequent bone marrow fattening. Conversely, administering granulocyte colony-stimulating factors (G-CSF) during chemotherapy in response to rising cellularity and the associated rise in signal can mimic tumor progress.

Tumor detection (qualitative analysis)

The sensitivity of DWI in detecting tumors depends on the tumor entity, tumor grading and the anatomical localization [8]. A hyperintense appearance was demonstrated in diffusion-weighted images for a number of entities such as prostate carcinoma (Fig. 2), breast carcinoma [19], gynecological tumors such as cervical and ovarian carcinoma, hepatocellular carcinoma (HCC), rectal carcinoma and other small cell tumors such as neuroendocrine tumors (Fig. 3), small cell tumors and many pediatric tumors such as hepatopo-, nephro- and neuroblastoma [20, 21]. The same signal behavior is also exhibited for hematological tumors such as multiple myeloma and lymphoma [13]. Similarly, the metastases also appear hyperintense in the diffusion-weighted images at high b-values [13]. In contrast, renal primary tumors as well as the metastases thereof are more difficult to see. Clear cell metastases in

<table>
<thead>
<tr>
<th>parameter</th>
<th>value</th>
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<tr>
<td>repetition time (ms)</td>
<td>9000</td>
</tr>
<tr>
<td>echo time (ms)</td>
<td>68</td>
</tr>
<tr>
<td>number of slices</td>
<td>200; four stations, 50 slices per station</td>
</tr>
<tr>
<td>field of View (FoV)</td>
<td>45 × 45 cm (variable)</td>
</tr>
<tr>
<td>matrix</td>
<td>128 × 128 interpolated to 256 × 256</td>
</tr>
<tr>
<td>number of acquired signals</td>
<td>six</td>
</tr>
<tr>
<td>fat suppression</td>
<td>short-tau inversion recovery (inversion time 180 ms)</td>
</tr>
<tr>
<td>diffusion coding</td>
<td>trapezoidal gradients, trace-weighted, bipolar gradients</td>
</tr>
<tr>
<td>parallel imaging</td>
<td>IPAT factor 2; number of reference lines: 30</td>
</tr>
<tr>
<td>slice thickness/gap (mm)</td>
<td>5/0</td>
</tr>
<tr>
<td>b-values (sec/mm²)</td>
<td>50, 800</td>
</tr>
<tr>
<td>bandwidth (Hz/pixel)</td>
<td>1628</td>
</tr>
</tbody>
</table>
particular often come across as unremarkable in diffusion imaging compared to other histological types [22]. Tumors of primarily normal histological structure as well as well differentiated or tumors of low malignancy are likewise occasionally difficult to detect [23]. Owing to their increased cellularity relative to the surrounding bone marrow tissues, osteolytic tumors are highly visible with diffusion imaging. False-positive findings are caused primarily by degenerative changes, fractures, edemas, infections, hemangiomas and dispersed red bone marrow [13]. More precise differentiability can be achieved by using the ADC value and conventional anatomical imaging. False-negative results in the detection of bone marrow tumors arise in the case of low-grade tumor infiltration of the bone marrow at the level of the base of the skull as well as the cranial vault as a result of the physiologically high signal intensity of the brain and in the case of metastases that develop in hypercellular bone marrow. Osteolytic bone metastases are more visible in diffusion imaging than primary sclerotic lesions or those induced by therapy, which can evade detection due to their low cell count and a low extracellular water content [20]. Assessing lymph nodes in cancer patients is generally difficult. Normally, lymph nodes are evaluated according to size,
which, however, provides only limited information on the tumor infiltration of lymph nodes (Fig. 4). Because of the overlapping values in the threshold range, the ADC value does not allow clear differentiation between malignant and benign lymph nodes [7], even if a few studies have established that metastatic lymph nodes exhibit a significantly lower ADC than their healthy counterparts [24 – 27]. An additional study shows that while DWI is inferior to FDG PET-CT in detecting primary colorectal carcinomas, it is superior in detecting lymph node metastases [28]. Patients with lymphoma diseases are an exception, with a study having established that nodal and extranodal manifestations of a non-Hodgkin’s lymphoma can be detected with certainty using diffusion imaging and that significant changes in ADC can be observed in patients under therapy, even at early stage, before the affected lymph nodes change in size.

Tumor characterization (quantitative analysis)
Whole-body diffusion-weighted imaging is of limited suitability for characterizing soft tissue lesions if merely the signal intensity is examined for evaluation without taking into consideration the morphological appearance and the ADC (Table 2). Only when examined together with anatomical localization and ADC can the tissue be characterized so that grading can be performed if certain tumors diseases are present. For example it was demonstrated that there is a correlation between ADC and histopathological differentiation (Gleason degree) in the case of prostate carcinoma, with moderately differentiated prostate carcinomas (Gleason 6) exhibiting higher ADC values in the peripheral zone than mildly (Gleason 7) or poorly differentiated (Gleason 8) prostate carcinomas [30, 31]. Tumor grading is also possible for additional tumor entities such as glial tumors, hepatocellular carcinoma (HCC), bladder carcinoma and rectal carcinoma [32 – 34].

Therapy monitoring

The complex changes in tumor tissue contrast over the course of therapy are a function of the particular therapy and the passage of time. In general, tumors with rising water content, e.g., due to interstitial edema, following lysis, necrosis or inflammations, appear increasingly hyperintense on diffusion-weighted images with low b-values (T2-shine-through-effect) and ADC cards (increasing diffusion). The observation of an increase in signal intensity alone on the b-images must not lead to false-positive findings. Each cytotoxic process that results in a necrosis or apoptosis on the cellular level exhibits, as early reaction owing to the failure of the sodium-potassium pump, an influx of water intracellularly and thereby a cytotoxic cellular edema that results in a drop in ADC. If, over the course of time, the cell membrane exhibits increased permeability resulting from the intracellular edema and shifting of the oncotic pressure, water flows from the intracellular space to the extracellular space [35]. This results in a drop in signal intensity in the diffusion-weighted images with high b-values and a corresponding increase in ADC value [9, 36, 37]. On the other hand, tumor progression is reflected, in addition to size progression, in the appearance of new cell-dense areas with correspondingly hyperintense signal or in the form of an increase in signal intensity in the diffusion-weighted images. Cell death resulting from response to therapy brings on a change in the diffusion-weighted image prior to the change in lesion size. Diffusion imaging can therefore be considered to be useful tool for the early measurement of response to therapy [13, 38, 39]. Increases in ADC that can be associated with the death of tumor cells have been found to be inconsistent over the course of time. A drop in ADC can be expected in the process of a clearing reaction of dead cells by macrophages and subsequent tissue reconstruction, e.g. through fibrosis. This is a potential source of error posed by diffusion-weighted imaging, since both the tissue reconstruction during the healing process and a tumor recurrence can bring about a drop in ADC. Fig. 5 provides an overview of the influence of therapy on ADC on the cellular level.

Therapy-induced changes in bone marrow
Because T1- and T2-weighted MR images often show no substantial changes with regard to the evaluation of bone marrow over the course of treatment of bone marrow diseases, these purely morphological sequences would appear to have only limited use for monitoring therapy [40].
Diffuse infiltration of bone marrow
This is where diffusion imaging provides an added clinical benefit, since it can show that in patients with leukemia, for example, there is a correlation between an increase in ADC and successful therapy [41]. Fig. 6 shows a response to therapy using a patient with multiple myeloma as an example. In contrast, patients not experiencing clinical improvement exhibit persisting hyperintensities in diffusion-weighted imaging with high b-values, which suggests an ongoing tumor-induced hypercellularity. While the long-term changes in bone marrow observed following successful treatment of the underlying disease depend on the type of tumor and treatment plan, the signal intensity and ADC slowly decline in the months following the start of therapy in most cases [42]. Like the elapsing pathophysiological process, signal alterations in diffusion imaging accompanying the curing of bone marrow disease have not yet been explained in detail. It is apparent, however, that effects such as sclerosis, clearing reactions of dead tumor cells, repopulation of (usually yellow) bone marrow, chemotherapy-induced myelofibrosis and decreased perfusion lead to a reduction in signal intensity and ADC [43].

Focal infiltration of bone marrow
Studies examining neoadjuvant response to therapy in bone tumors such as osteosarcoma likewise show significant differences in ADC between patients responding to therapy, who exhibit correspondingly higher ADC values after therapy, and patients not responding to therapy [44]. However, the signal may still remain high in diffusion-weighted imaging despite response to therapy and the associated necrosis. This is due to the T2-shine-through-effect, which appears particularly in the case of bone marrow diseases such as multiple myeloma with focal or diffuse infiltration pattern and in the case of lymphomas, but also occasionally with other solid metastatically dispersing neoplasms. Nevertheless, the T2-shine-through-effects are comparatively easy to detect through the ADC map, which in this case exhibits very high values when viewed together with T2-weighted sequences or short-tau-inversion-recovery (STIR).

Reaction in soft tissue
Rises in ADC following successful chemo and radiation therapy have been documented for several anatomical regions, including breast carcinoma, primary and metastatic liver tumors as well as for primary osteosarcoma and malignant brain tumors [13]. Just three to seven days after their first chemotherapy session, patients with hepatic metastasis of stomach and colorectal carcinomas exhibit a change in ADC correlating to a response to therapy [45]. Several clinical and pre-clinical studies have examined how radiation impacts ADC, showing that ADC increases rather rapidly in the case of radiation-sensitive tumors. In one pre-clinical study, proton MR spectroscopy detected visible changes in ADC after only 24–72 hours without any signs of the lesion changing size, but coinciding with a metabolic disturbance of the tumor tissue [46]. Rises in ADC appear incrementally with fractionated radiation therapy, where the greatest visible changes can be observed consecutively at the end of therapy [47, 48]. This effect can most likely be attributed to the death of tumor cells, the development of interstitial edema resulting from radiogenic inflammatory reactions and increased vascular permeability. Overall, it can be postulated from a number of studies that the absence of a rise in ADC following the conclusion of the early reaction with the formation of intracellular edema under chemotherapy, radiation therapy or a combination thereof suggests a poor response to therapy.

Table 2 Signal behavior in diffusion-weighted imaging.
<table>
<thead>
<tr>
<th>signal intensity at high b-values</th>
<th>ADC</th>
<th>Interpretation</th>
</tr>
</thead>
<tbody>
<tr>
<td>↑</td>
<td>↑</td>
<td>hypercellular tumor, cytotoxic edema (DD occasionally abscess, viscous fluid, hematoma)</td>
</tr>
<tr>
<td>↑</td>
<td>↑</td>
<td>T2-shine-through-effect pronounced at low b-values</td>
</tr>
<tr>
<td>↓</td>
<td>↑</td>
<td>low-protein fluids, necrosis, occasionally well differentiated, hypocellular adenocarcinoma</td>
</tr>
<tr>
<td>↓</td>
<td>↑</td>
<td>susceptibility artifacts, fibrotic tissue, muscle tissue, bones, adipose structures</td>
</tr>
</tbody>
</table>

Fig. 4 a Coronary 3D-Reconstruction of an inverted DWI of the neck, b coronary T1-w post-gadolinium of the neck, c Hybrid-image of T1-w post-gadolinium and inverted coronary DWI of the neck in a patient with lymphatic metastasis in thyroid cancer.
Comparability of whole-body DWI and FDG-PET

Fluorodeoxyglucose (FDG) PET and whole-body DWI are assuming a complementary role in oncological evaluation, since they are able to illustrate different tissue properties. While FDG PET presents metabolic information on intracellular glucose uptake, diffusion imaging primarily represents microstructural information. The FDG uptake in PET for diagnostic purposes and therapy monitoring is typical for several FDG-positive entities such as bronchial cancer, rectal cancer and lymphoma and is advantageous over DWI in evaluating structures such as the mediastinum, the lungs and the spleen [49]. However, a number of tumors such as prostate carcinoma, hepatocellular carcinoma and glial tumors may prove to be FDG-negative depending on their degree of differentiation [50–52]. Precisely in the case of these tumors with a low FDG uptake, which also include neuroendocrine tumors, thyroid gland carcinomas and several types of low-malignancy lymphoma, diffusion images with high b-values are advantageous for detection [53]. Diagnostic advantages of diffusion-weighted imaging are also apparent in anatomical regions that have a heavy accumulation of FDG in healthy tissues such as in the brain, liver, bone marrow and urinary system. Several studies show an inverse correlation between the standardized uptake value (suv) of PET and ADC [54, 55]. Because the two procedures complement one another, combining FDG-PET and whole-body DWI through PET-MRT could constitute a further advancement in the detection and characterization of tumor-associated lesions [56] as well as in the evaluation of response to therapy [57].

Indications for whole-body diffusion-weighted imaging in oncology

Whole-body diffusion-weighted imaging makes it possible to show the primary location and remote metastasis of many types of tumors within the limits of the particular accuracy available [5]. Suitable candidates would appear to be tumors with small cell and cell-dense appearance under histological examination, such as multiple myeloma, malignant myeloma (Fig. 7), neuroendocrine tumors, small cell tumors and pediatric tumors [13]. Whole-body diffusion-weighted imaging is beneficial in the case of bone marrow diseases as well as for evaluating therapy response in patients with mixed osteoplastic-osteolytic metastasis patterns, particularly breast and prostate carcinoma. The method is additionally advantageous for instances in which the use of ionizing radiation must be decreased to a minimum, such as for children and pregnant women.
[58 – 60] as well as when frequent follow-up examinations are necessary. It is likewise imperative to minimize the risk of inducing radiogenic secondary tumors in tumor patients who have the prospect of a long life expectancy following a curative therapy approach. Patients who cannot undergo contrast-enhanced CT examination due to impaired kidney functions or allergies can also benefit from this diagnostic imaging technology. With its higher tumor-to-normal tissue contrast, whole-body diffusion-weighted imaging holds further potential for tumor detection, grading and therapy monitoring, while being easier and more cost-effective to perform. However, validation through multicentric studies is currently limited by the lack of standardization of methods.

Note

In the eFirst-version of this article the sequence of the figures 7b and 7c was wrong. It has been corrected.

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