

# Whole-Body Diffusion-Weighted MRI: Tips, Tricks, and Pitfalls

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**OBJECTIVE.** We examine the clinical impetus for whole-body diffusion-weighted MRI and discuss how to implement the technique with clinical MRI systems. We include practical tips and tricks to optimize image quality and reduce artifacts. The interpretative pitfalls are enumerated, and potential challenges are highlighted.

**CONCLUSION.** Whole-body diffusion-weighted MRI can be used for tumor staging and assessment of treatment response. Meticulous technique and knowledge of potential interpretative pitfalls will help to avoid mistakes and establish this modality in radiologic practice.

**E**xtracranial diffusion-weighted MRI (DWI) is increasingly used for disease evaluation in oncology [1]. DWI depicts differences in the mobility of water in tissues. Cellular tumors exhibit greater impeded water diffusion, reflected as high signal intensity on images with high b values, and return lower apparent diffusion coefficient (ADC) values [2, 3]. This mechanism of image contrast generation is being harnessed in clinical practice and in research for detecting malignancy in different tumor types and organ systems to inform tumor staging and management decisions.

DWI can be performed relatively quickly with a free-breathing spin-echo echo-planar technique [1, 4]. Advances in magnet design, receiver coil technology, and acquisition mean that DWI can be performed over multiple imaging stations in a relatively short time. The concept of whole-body imaging with DWI was initially proposed by Takahara et al. in 2004 [5]. Those authors described whole-body DWI with background signal suppression. Since then, interest in advancing the utility of the technique has continued to grow, particularly in the field of oncologic imaging.

The idea of whole-body MRI acquisition in multiple image stacks of the entire body is not new. In the 1990s, whole-body MRI with conventional T1-weighted and STIR sequences was proposed and investigated for tumor detection [6–8]. These studies had relatively high sensitivity, specificity, and diagnostic accuracy for visualization of a variety of tumor types [6, 7, 9–12]. However, whole-body MRI

has not gained widespread adoption despite its apparent usefulness. One of the negative factors was likely the relatively long examination time. Typically, these studies took 1 hour or longer to perform and resulted in a multitude of images, which posed challenges for image viewing and interpretation before sophisticated workstations became available.

Since the initial studies, substantial improvements have been made in MRI hardware design and software, which allow performance of multipart MRI examinations with relative ease [13, 14]. The advent of continuous moving table technology allows even more efficient image acquisition [12, 13, 15, 16]. Dedicated workstation software also streamlines onsite and remote viewing of whole-body datasets. These software tools are furnished with pushbutton functions that stitch together images acquired at different anatomic stations and displayed as a single image to facilitate reading. Whole-body DWI was developed amid these innovations, making it possible to add DWI to conventional whole-body imaging protocols. With modern state-of-the-art MRI systems, whole-body MRI from vertex to the midhigh (analogous to the coverage for a typical body PET/CT examination) can be accomplished with T1-weighted, fat-suppressed T2-weighted, or STIR and DWI sequences in 40–60 minutes, making it a viable tool for clinical deployment. For this article, an understanding of the basic principles of DWI is assumed and is not discussed. A number of review articles are available [1–3, 17, 18].

**Keywords:** diffusion-weighted imaging, MRI, oncology, pitfalls, technique, whole body

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## Whole-Body Diffusion-Weighted MRI

### Clinical Impetus for Whole-Body Diffusion-Weighted MRI

There has been a resurgence of interest in whole-body MRI, stimulated in part by clinical needs and concerns about radiation safety in CT [19, 20]. Ionizing radiation from CT studies can result in a substantial increase in the lifetime risk of cancer [19]. As a result, CT manufacturers have responded by designing more efficient CT systems that will reduce radiation exposure for each study. Another option, however, is radiation avoidance by performing MRI instead of CT when appropriate, as in the care of young adults and children [21].

From the clinical perspective, there is emerging evidence that whole-body MRI may be a solution to current unmet needs

in cancer staging. A difficult area in tumor evaluation has been assessment of metastatic bone disease and the response to hormonal, chemotherapeutic, and radiation treatment. The difficulty arises because bone disease confined to the marrow cavity is not considered measurable according to conventional measurement criteria (e.g., the Response Evaluation Criteria in Solid Tumors) [22], although measurement criteria have been proposed for bone disease associated with a soft-tissue component [23].

The most widely prescribed imaging test for evaluation of bone disease is radionuclide bone scintigraphy with technetium-labeled tracers. However, the results can be false-negative in patients with active bone disease because lytic bone diseases may not induce an

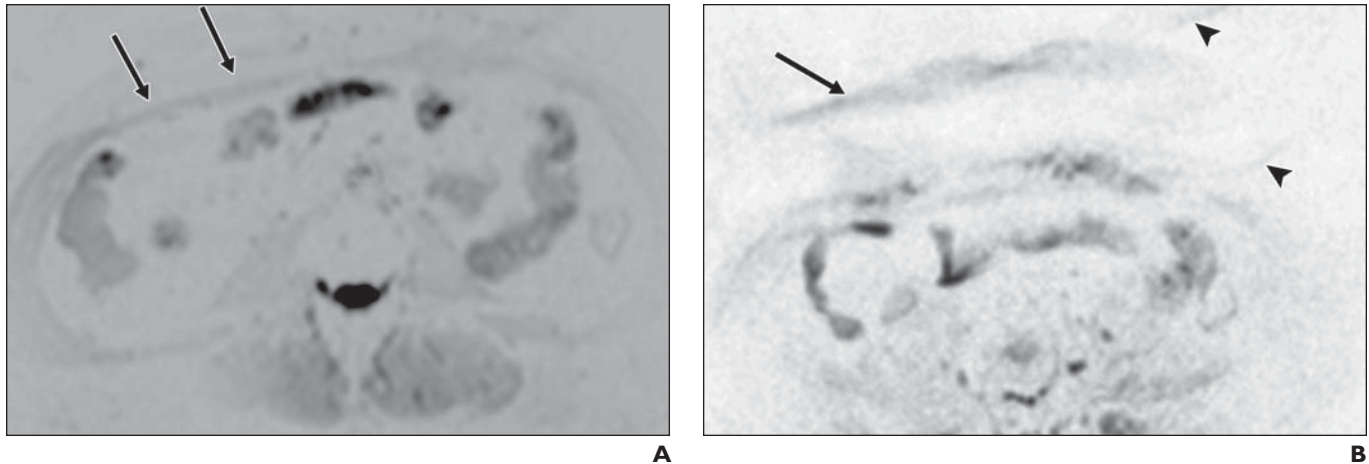
osteoblastic reaction [24, 25]. Furthermore, there is a global shortage of the  $^{99m}\text{Tc}$  isotope used for bone scintigraphy due to uncertainty over its supply [26]. The search is ongoing for an imaging technique to fill this gap. Although it is possible to replace  $^{99m}\text{Tc}$ -radiolabeled bone scintigraphy with  $^{18}\text{F}$ -labeled PET/CT [24], it is unlikely that there will be sufficient numbers of PET/CT scanners worldwide to cope with the workload and demand. Whole-body MRI is poised to become a viable alternative. Studies of whole-body MRI have shown high diagnostic sensitivity, compared with bone scintigraphy and FDG PET/CT, for detecting bone metastasis [6, 9, 24, 27–32].

Literature findings [33–35] suggest that whole-body MRI that includes DWI can be developed to detect diagnostic and response

**TABLE 1: Selected Studies of Whole-Body Diffusion-Weighted MRI (DWI) for Varied Indications**

Study	Year	No. of Patients	Whole-Body MRI Technique	Comparison Modality	Findings
<b>Lymphoma</b>					
van Ufford et al. [36]	2011	22	DWI, STIR and T1-weighted	FDG PET/CT	77% concordance with PET for staging; whole-body DWI understaging rate was 0% but overstaging, 23% (5/22)
Lin et al. [38]	2011	15	Respiratory-gated DWI	FDG PET/CT	ADC increased in nodes after chemotherapy; use of ADC and size criteria reduced false-positive rate for residual nodal masses
Lin et al. [39]	2010	15	Respiratory gated DWI	FDG PET/CT	90% sensitivity, 94% specificity in assessment of nodal involvement by size; 81% sensitivity, 100% specificity with ADC and size criteria
Kwee et al. [49]	2009	31	DWI	CT	DWI similar to CT for disease staging
<b>Mixed disease</b>					
Fisher et al. [60]	2011	68	T2-weighted and DWI	FDG PET/CT	High detection rate and PPV with T2-weighted DWI, evaluated side by side (72%, 89%) and by fusion (74%, 91%)
Gutzeit et al. [44]	2010	36	DWI	FDG PET/CT	DWI had higher sensitivity (97%) than PET/CT (91%) for patients with > 10 skeletal lesions
Nakanishi et al. [52]	2007	30	DWI, STIR, T1-weighted	Bone scintigraphy, CT	STIR and T1-weighted DWI had higher sensitivity (96%) and PPV (98%) than bone scintigraphy and whole-body MRI without DWI
<b>Malignant melanoma</b>					
Laurent et al. [40]	2010	35	T1-weighted, STIR, and DWI	FDG PET/CT	82% sensitivity, 97% specificity for DWI; 73% sensitivity, 93% specificity for PET/CT
<b>Breast cancer</b>					
Heusner et al. [43]	2010	20	DWI	FDG PET/CT	91% sensitivity, 72% specificity for DWI; 94% sensitivity, 99% specificity for PET/CT
<b>Lung cancer</b>					
Chen et al. [46]	2010	56	DWI	FDG PET/CT	Lymph node metastasis: 91% sensitivity, 90%, specificity for DWI; 98% sensitivity, 97% specificity for PET/CT. Other metastases: 90% sensitivity, 95% specificity for DWI; 98% sensitivity, 100% specificity for PET/CT
Takenaka et al. [47]	2009	115	DWI, T1-weighted, STIR	FDG PET/CT, $^{99m}\text{Tc}$ bone scintigraphy	Specificity and accuracy of whole-body MRI with DWI significantly better than scintigraphy for bone metastasis
Ohno et al. [50]	2008	203	DWI, T1-weighted, STIR	FDG PET/CT	Accuracy of whole-body-MRI with DWI ( $A_z$ , 0.87) similar to that of PET/CT ( $A_z$ , 0.89).

Note—ADC = apparent diffusion coefficient, PPV = positive predictive value,  $A_z$  = area under the curve.



**Fig. 1**—45-year-old healthy man. Potential problems with diffusion-weighted MRI at 3 T. **A**, Diffusion-weighted MR image ( $b = 750 \text{ s/mm}^2$ ) shows eddy current–induced geometric distortion (arrows) resulting in distorted appearance of right anterior abdominal wall. **B**, Diffusion-weighted MR image ( $b = 750 \text{ s/mm}^2$ ) shows Nyquist ghosting artifact (arrowheads). Arrow indicates area of distortion in **A**. **C**, Sagittal 3-T MRI reformats of three image stacks ( $b = 750 \text{ s/mm}^2$ ) show misalignment of spine (arrows).

biomarkers of malignant bone and soft-tissue diseases. Table 1 summarizes selected studies on the use of whole-body DWI for disease staging and response assessment in the care of oncology patients [36–52].

### Implementation of Whole-Body Diffusion-Weighted MRI

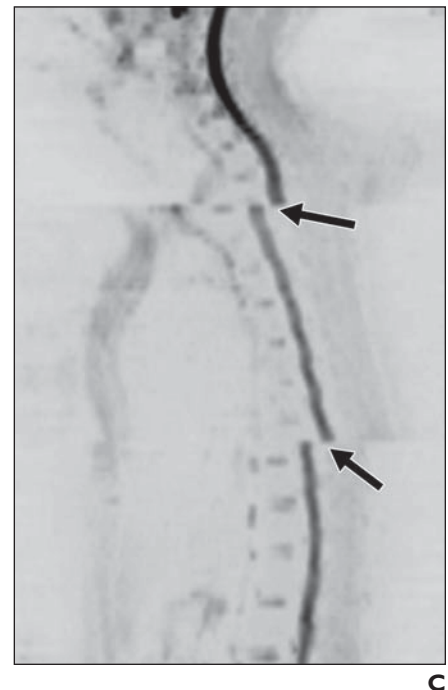
The use of MRI for oncologic imaging is continuing to grow. Imaging platforms being introduced have technologies to optimize multistation body imaging that includes DWI. This capability increases the efficiency of sequential imaging at more than one anatomic station without manual repositioning of the patient or the surface receiver coils between acquisitions. We discuss the tips, tricks, and pitfalls of implementing whole-body DWI in current clinical MRI systems. Included are the choice of field strength, patient positioning on the unit, optimization of the imaging sequence, image processing and viewing, and image interpretation.

#### Choice of MRI System: 1.5 T Versus 3 T

With current technology, whole-body DWI is easier to implement with 1.5-T than with 3-T systems. Whole-body DWI at 3 T is often more difficult because artifacts can be more challenging to control and minimize. In addition to the general issues at 3 T, such as dielectric effects and tissue-specific absorption rates [53, 54], there are challenges related to the DWI measurements. First, eddy currents induced by rapid switching of the magnetic gradient field with echo-planar imaging cause residual mag-

netization that results in geometric distortion and image shearing [4] (Fig. 1A). This distortion can make it more difficult to register and align DW images with corresponding morphologic T1- and T2-weighted images. Second, because of the greater  $B_1$  field inhomogeneity at 3 T, it can be difficult to achieve uniform fat suppression across large fields of view, and the result is chemical-shift and ghosting artifacts (Fig. 1B). Third, because the MR frequency offsets applied can differ markedly between anatomic stations, voxel shift can occur in the phase-encoding direction and cause misalignment of structures (e.g., the spinal cord) between imaging stations (Fig. 1C). The result is difficulty in accurate alignment of individual imaging stations to produce smooth composite images (e.g., whole-body sagittal images), even though images from individual anatomic stations are of high quality. This limitation may not affect the diagnostic quality of the acquired images but falls short of producing pleasing reformatted images for clinicians and patients to review. Coronal reformats are usually less affected because the phase-encoding direction is often anteroposterior.

Despite the limitations, it is clear that imaging at 3 T has the intrinsic advantage of a higher signal-to-noise ratio (SNR), which can enhance lesion detection. As imaging technology continues to improve, a variety of implementations are being explored to minimize eddy current effects and to optimize fat suppression across a large FOV at 3 T. These features are discussed later. Post-



processing techniques are being developed for image alignment and registration. In the current state of the art, whole-body DWI technique is more robust at 1.5 T, but it is conceivable that with the current pace of imaging improvement, the technique may soon be equally successful at 3 T.

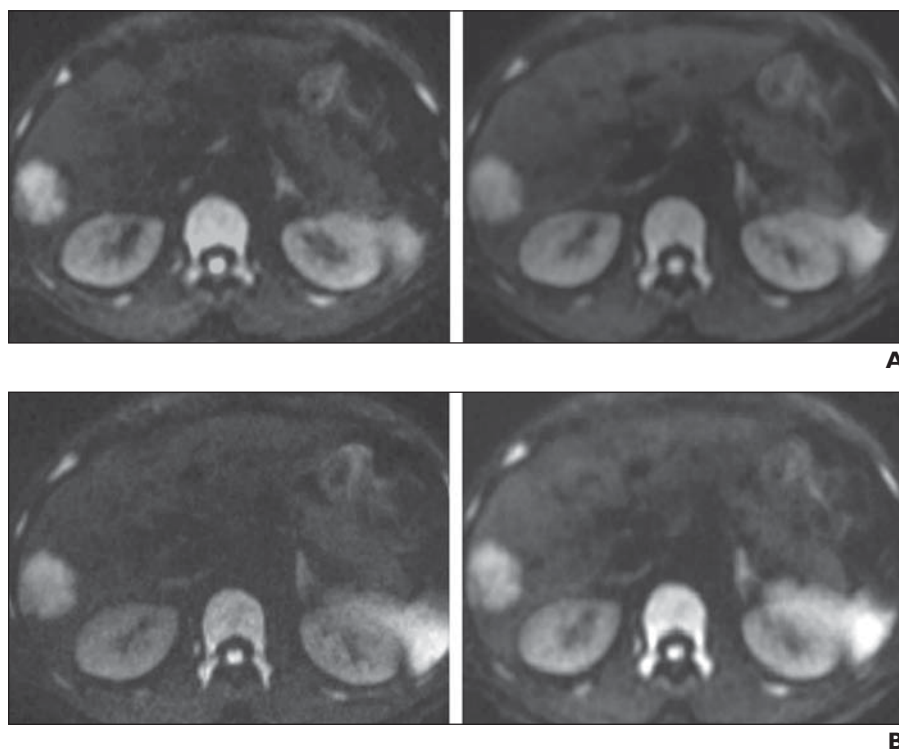
#### Patient Setup

The success of a whole-body DWI study relies in part on patient cooperation to minimize movement during the study. As such, patients should be told of the approximate examination time (typically 40–60 minutes) and be warned that the study can be noisy owing to the vibration of the MRI bed. The patient should be offered the opportunity for bladder emptying

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**TABLE 2: Typical 1.5-T Diffusion-Weighted Protocol for Whole-Body MRI With Three 1.5-T Systems**

Protocol	Avanto (Siemens Healthcare)	Intera (Philips Healthcare)	Signa (GE Healthcare)
Imaging plane	Axial	Axial	Axial
FOV (cm)	380 × 380	400 × 280	440 × 440
Matrix Size	150 × 256	128 × 96	128 × 88
TR	14,000	8322	6625
TE	72	70	64.6
Echo-planar imaging factor	150	37	
Parallel imaging factor	2	2	
No. of signals averaged	4	4 (b = 0), 12 (b = 1000)	3
Section thickness (mm)	5 contiguous	5 contiguous	8
Direction of motion probing gradients	3-scan trace	Tetrahedral encoding	
Receiver bandwidth	1800 Hz/pixel	7.757 (water-fat shift/pixel)	
Fat suppression	STIR (inversion time, 180 ms)	STIR (inversion time, 180 ms)	STIR (inversion time, 160 ms)
b value (s/mm <sup>2</sup> )	Typically 0–100 and 600–1000	0, 1000	Single b value, 600
Acquisition time per station	4 min 30 s	4 min 2 s	1 min 28 s



**Fig. 2**—46-year-old man with liver metastasis of colorectal cancer. Example of effects of decreasing TE and matrix size.

**A**, Diffusion-weighted MR images acquired with TEs of 90 ms (*left*) and 76 ms (*right*) with all other imaging parameters equal show use of shorter TE improves signal-to-noise ratio.

**B**, Diffusion-weighted MR images acquired with matrix sizes of 256 × 256 (*left*) and 128 × 128 (*right*) with all other imaging parameters equal shows reducing matrix size improves signal-to-noise ratio.

before the examination to minimize discomfort during the examination.

Many MRI systems are migrating to the use of multiple surface receiver coil arrays deployed over the body to achieve the best SNR.

These coils should be deployed contiguously to each other or with a slight degree of overlap (up to 5 cm). Using multiple receiver coil arrays also helps to streamline workflow, improve patient throughput, and minimize exam-

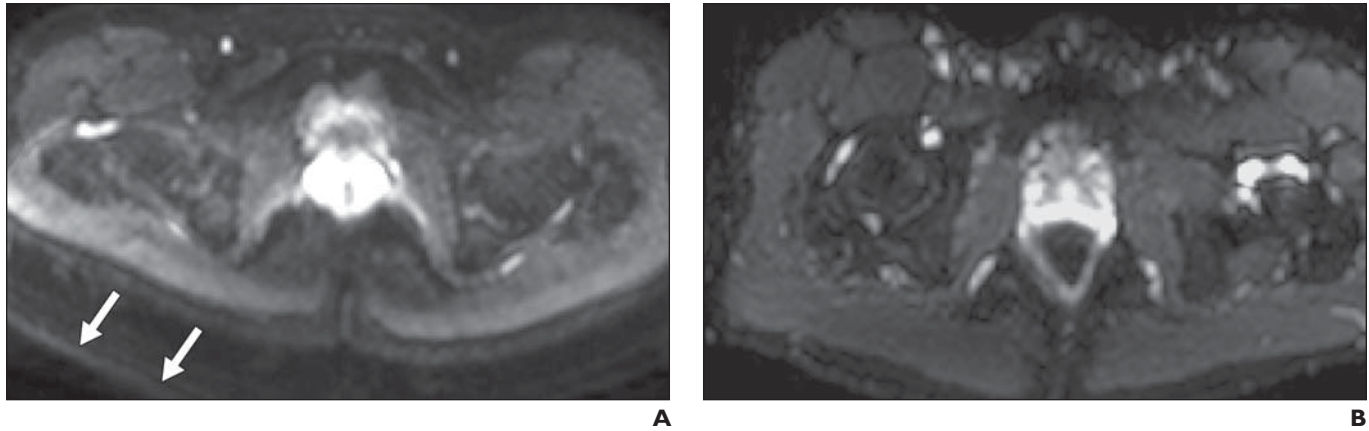
ination times. With some older MRI systems, it may be possible to perform whole-body DWI with just a single phased-array receiver coil. Takahara et al. [55] called this method the sliding coil technique. Where surface coils are not available, whole-body DWI can be performed with the receiver elements in the main magnet bore. However, these images have lower spatial resolution and SNR and cannot be acquired with parallel imaging.

The optimum patient setup for a whole-body DWI examination depends on the local MRI hardware configuration, which has a bearing on the efficiency of acquisition and the quality of the images obtained.

### Imaging Parameters

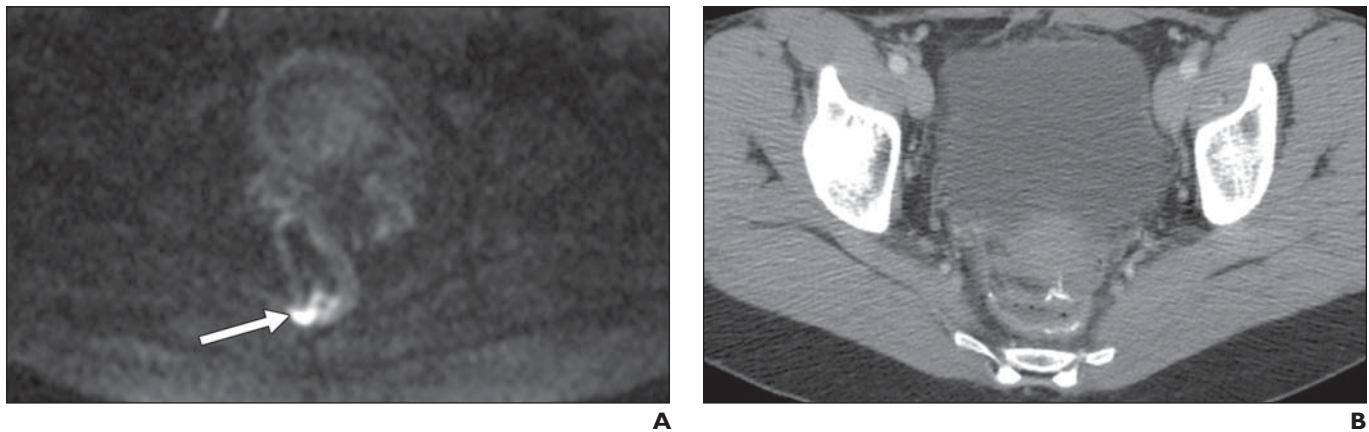
The principles of obtaining high-quality whole-body DW images are similar to those for regional examinations. Acquisition and parameter selection should be streamlined to optimize SNR and to minimize artifacts [4]. Table 2 summarizes typical imaging parameters for whole-body DWI with 1.5-T MRI systems from three vendors.

In addition to axially acquired DW images, conventional T1-weighted and STIR and T2-weighted fat-suppressed images are usually obtained in the axial or coronal plane. Breath-hold 3D T1-weighted gradient-echo sequences allow rapid acquisition of T1-weighted images [56]. For assessment of suspected metastatic bone disease, T1-weighted and STIR and T2-weighted fat-suppressed sagittal images of the whole spine are valuable. Coronal DWI acquisitions with higher parallel imaging factors are being evaluated for reducing acquisi-



**Fig. 3**—35-year-old healthy man. Example of fat suppression at 3 T.

**A**, Axial STIR fat-suppressed MR image ( $b = 750 \text{ s/mm}^2$ ) shows substantial chemical-shift ghosting (arrows) due to poor fat suppression of posterior abdominal wall. **B**, Axial MR image ( $b = 750 \text{ s/mm}^2$ ) acquired with combinatorial fat suppression with spectral attenuated inversion recovery and slice selective gradient reversal shows improvement of fat suppression without marked image artifacts.



**Fig. 4**—56-year-old woman with history of anterior resection for rectal carcinoma. Example of susceptibility artifacts.

**A**, MR image ( $b = 900 \text{ s/mm}^2$ ) appears to show area of impeded diffusion (arrow) in posterior rectal wall. However, finding is due to susceptibility artifact from surgical material in area.

**B**, CT image shows high-attenuation material related to previous surgery.

tion time to less than 30 minutes. The development of whole-body DWI with continuous table movement may further streamline workflow [16]. With continuous table movement, imaging is always performed in the isocenter of the magnet, avoiding the problem of peripheral field image distortion.

With regard to the choice of  $b$  value, two  $b$  values usually suffice because increasing the number of  $b$  values prolongs examination time. Because the technique is usually used as a contrast mechanism to identify cellular tissues, the higher  $b$  value chosen is usually 600–1000  $\text{s/mm}^2$ . This choice renders the diffusion sequence for the optimal SNR for detecting lesions with ADC values in the range of 500–900  $\text{mm}^2/\text{s}$ , typical of malignancy. The lower  $b$  value can be 0–100  $\text{s/mm}^2$ . Using a lower  $b$  value of 50–100  $\text{s/mm}^2$  results in vascular sig-

nal suppression (black-blood images), which can aid disease detection but at the expense of a longer acquisition time compared with  $b = 0 \text{ s/mm}^2$ . When two  $b$  values are used, the ADC can be computed for disease quantification or to support image interpretation (e.g., identifying T2 shine-through).

In the performance of DWI, consideration should be paid to maximizing image SNR. One simple rule is to minimize the TE to as low as achievable [4] (Fig. 2). A common error is to use too high a spatial resolution to match those of the morphologic images, leading to lower SNR. Use of lower spatial resolution due to a coarser imaging matrix (e.g.,  $180 \times 180$ ) helps to improve SNR, particularly at higher  $b$  values (Fig. 3). When technically possible, free-breathing image acquisition is a robust technique because it allows multiple

signal averages, which improve SNR. However, when this technique is suboptimal, multiple breath-hold acquisitions may be considered. Using multiple averaging also allows thinner image partitions (typically 4–5 mm) for multiplanar image reformats [4].

Many types of artifacts can degrade image quality. Common clinical artifacts include poor fat suppression, which results in chemical shift artifacts; eddy current–induced geometric distortions and image shearing; and Nyquist, or  $N/2$ , ghosting.

To optimize fat suppression, the use of non-selective fat suppression techniques such as STIR is preferable over chemical-selective sequences such as spectral attenuated inversion recovery, spectral presaturation by inversion recovery (SPIR), and chemical fat suppression over large fields of view [5]. Another method

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**TABLE 3: Problems and Suggested Solutions in Body Diffusion-Weighted MRI With Spin-Echo Echo-Planar Technique**

Problem	Solution
Poor signal-to-noise ratio	Use shortest TE achievable Use multiple averaging method Choose coarser matrix size Use thicker partition thickness Increase FOV Choose lower b values Use parallel imaging Optimize receiver bandwidth
Relatively long TE (e.g., > 100 ms)	Try to shorten TE in one of the following ways: Use simultaneous gradient application schemes (e.g., 3-scan trace) Increase receiver bandwidth Use parallel imaging
Poor fat suppression and chemical-shift artifact	Use non-chemical-selective fat-suppression scheme over the wide FOV (e.g., STIR) Increase receiver bandwidth Use combinatorial fat suppression techniques at 3 T
Eddy current effects causing geometric distortion and image shearing	Optimize or increase receiver bandwidth Use simultaneous gradient application schemes to decrease peak gradient amplitudes
Nyquist (N/2) ghosting	Optimize or decrease receiver bandwidth
G noise	Reduce parallel imaging factor Use larger FOV
Susceptibility artifacts	Avoid scanning over areas with implants
Poor alignment of image stacks between imaging stations	At 1.5 T, omit image shimming and apply center frequency of one imaging station to all subsequent stations Consider continuous table motion acquisition Apply software postprocessing

is spectral-spatial water-only excitation. These approaches may be successful at 1.5 T but still be unsatisfactory at 3 T. Performing DWI at 3 T may require combinatorial fat suppression to ensure uniform fat suppression across the entire FOV [57] (Fig. 4). For example, it may be advantageous to combine STIR with chemical fat suppression (e.g., SPIR). Another fat-suppression technique found useful at 3 T is the slice selective gradient reversal (SSGR) technique [57]. However, use of SSGR, either alone or in combination with STIR imaging, can result in undesired off-resonance signal suppression from water protons (water suppression), which can obscure lesions.

Use of echo-planar imaging, the rapid switching of magnetic gradients, results in residual magnetic fields, which induce eddy currents and cause image distortion. Use of simultaneous gradient application schemes (e.g., three-scan trace and tetrahedral encoding) reduces eddy currents by reducing the peak amplitude of the applied diffusion gradient.

One effective way of dealing with  $B_0$ -induced distortions is to adjust the receiver bandwidth. The receiver bandwidth is the range of frequencies used to sample the MR signal. Increasing the receiver bandwidth reduces chemical-shift artifacts, decreases the TE of signal acquisition, and reduces geometric distortion, all of which improve image quality. The disadvantages of increasing the receiver bandwidth, however, include reduction of SNR and an increase in Nyquist ghosting (see later). Thus the receiver bandwidth should be adjusted to balance these effects. For DWI of the whole body, the optimal receiver bandwidth is usually much higher than those used for conventional MRI sequences. Typically, a receiver bandwidth of approximately 1600–2000 Hz/pixel is used for body DWI. The receiver bandwidth is expressed differently on different MRI systems. Adjusting the receiver bandwidth may mean changing the receiver bandwidth range (expressed in kilohertz) (GE Healthcare), the water-fat shift (expressed in pixels) (Philips

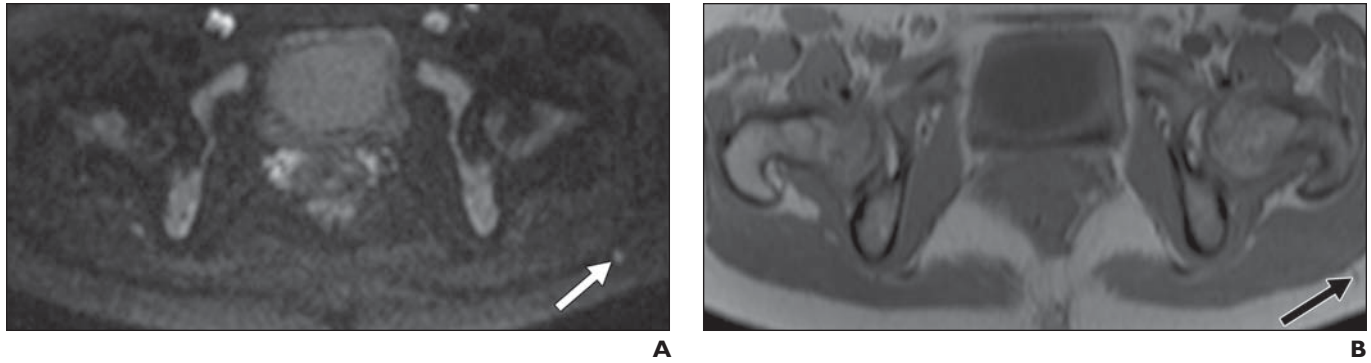
Healthcare), or the receiver bandwidth per pixel (Hz/pixel) (Siemens Healthcare).

The use of echo-planar imaging can result in a sampling artifact, known as Nyquist, or N/2, ghosting. During echo-planar imaging, adjacent lines of k-space are sampled with opposite readout gradients. Misalignment in the sampling from positive and negative gradients leads to additional line modulation in k-space, which is observed as ghosting of the image. Nyquist ghosting can be reduced by applying appropriate phase correction to the data. However, optimizing the receiver bandwidth and echo spacing is also an effective measure.

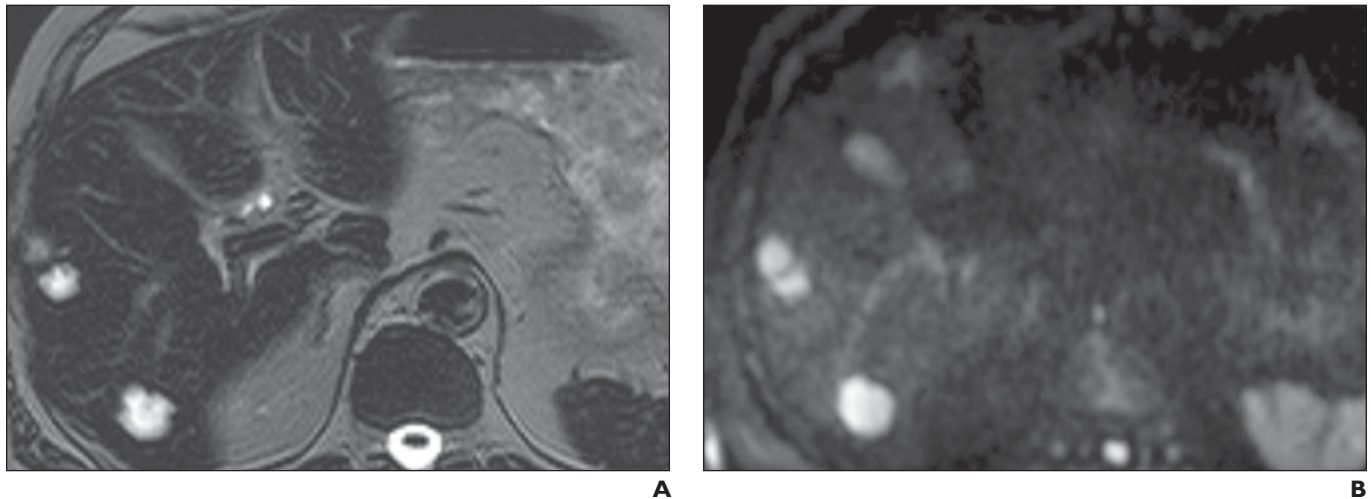
The common technical issues encountered at body DWI and their solutions are summarized in Table 3. These broad principles should help to guide parameter adjustments in clinical practices.

### *Image Processing and Viewing*

For viewing and reporting of whole-body DWI studies, it is desirable to reformat the



**Fig. 5**—62-year-old woman with history of ovarian cancer.  
**A**, MR image ( $b = 900 \text{ s/mm}^2$ ) through lower pelvis shows focus of impeded diffusion (*arrow*) over left gluteal area.  
**B**, MR image shows low-signal-intensity tubular structure (*arrow*) in left gluteal area. Apparent impeded diffusion most likely results from slow flow in presumed small venule.



**Fig. 6**—55-year-old man with colorectal liver metastasis. Example of T2 shine-through.  
**A**, T2-weighted MR image (TE, 240 ms) shows three high-signal-intensity mucinous metastatic lesions in right lobe of liver.  
**B**, MR image ( $b = 750 \text{ s/mm}^2$ ) shows apparent high signal intensity, suggesting impeded diffusion of metastatic lesions.  
**C**, MR image shows high apparent diffusion coefficient of lesions consistent with T2 shine-through effect.

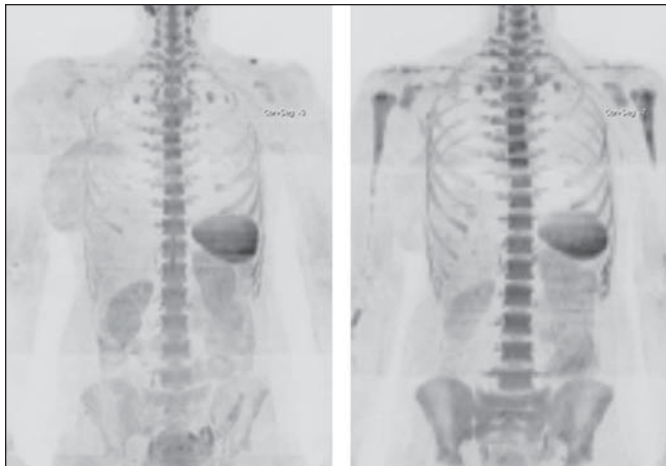
high  $b$  value DW images in the axial, coronal, and sagittal planes for multiplanar assessment. Whole-body DW images are usually displayed with an inverted gray scale. However, to compose DW images from different anatomic sections, it is important to ensure that data acquisition is performed in a way that minimizes voxel shifts between imaging stations for accurate stack alignment.

At 1.5 T, one method used to minimize stack misalignment is to omit the volume shimming at individual anatomic stations. Instead, shimming is performed at the first imaging station (e.g., head and neck area), and the same frequency offset is applied to all sub-

sequent imaging volumes. This method minimizes voxel shifts and misalignment between imaging stations [58] without compromising image quality. However, it should be noted that this approach works well at 1.5 T but

cannot be applied successfully at 3 T because the frequency-dependent voxel shift is more marked at higher field strength. At 3 T, omitting image shimming for individual stations often results in suboptimal fat suppression and

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**Fig. 7**—40-year-old woman who has undergone left mastectomy and axillary nodal dissection for triple-negative breast cancer and is receiving adjuvant chemotherapy with granulocyte colony-stimulating factor (G-CSF) support. Inverted gray-scale whole-body diffusion-weighted maximum-intensity-projection images obtained before (*left*) and at 6 months after (*right*) cyclic G-CSF administration show generalized and symmetric increase in marrow signal intensity evident 6 months after treatment can mimic diffuse marrow infiltration.

substantial chemical-shift artifacts. Therefore, other solutions, including better image post-processing, are being devised for display and analysis at 3 T.

### Image Interpretation

At qualitative visual appraisal of high b value images, disease is identified as areas of impeded diffusion. With inverted gray scale for image display, these areas appear as dark regions against a normal white background. Whole-body DW images should be reviewed with conventional T1-weighted or STIR or fat-suppressed T2-weighted images to ensure accurate interpretation. In oncology, DWI findings are not specific for malignancy because other cellular processes, such as inflammation, can result in similar findings. A clear clinical history should aid the weighted interpretation of abnormalities.

Readers should also be aware of interpretive pitfalls, which can give rise to false-positive and false-negative results.

**False-positive findings**—The following features on whole-body DW images can be misinterpreted as disease.

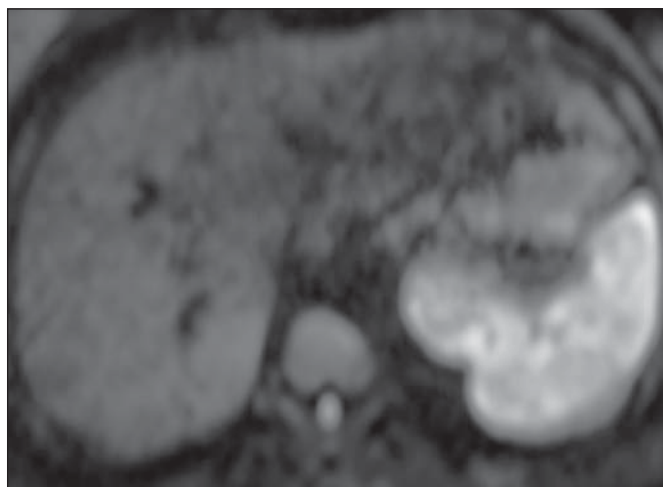
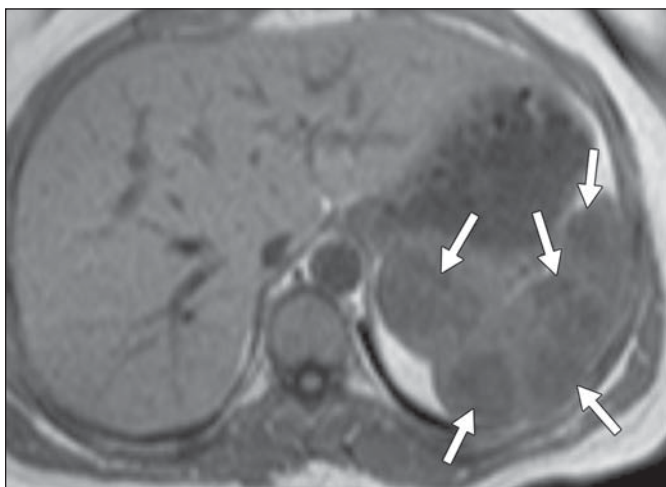
Artifacts resulting from image ghosting, poor fat suppression, or susceptibility effects should not be misinterpreted as disease (Fig. 5). The clue to artifacts resulting from image ghosting and poor fat suppression is that they may appear as recapitulation of structures seen elsewhere on the image or appear at boundaries between fat and water interfaces (e.g., abdominal wall).

Normal lymph nodes return impeded diffusion to different extents during DWI. Although malignant nodes tend to have more impeded diffusion, this characteristic can be difficult to identify, particularly if the im-

age brightness and contrast are not equalized across anatomic stations. Quantitative ADC measurements reportedly show that malignant nodes have lower ADC values than malignant lymph nodes but with substantial overlap. As such, current criteria for diagnosing nodal involvement often rely on a combination of signs, including nodal signal intensity (on its own or compared with the primary tumor), ADC value, and size criteria (short axis diameter > 1 cm).

Tiny foci (typically 1–2 mm) of impeded diffusion are sometimes detected on b value images that are difficult to correlate with structures on the corresponding T1-weighted or STIR images. Some of these foci may represent slow flow in small venules (Fig. 6). In other cases, however, the cause cannot be easily confirmed but may represent small nerves, ganglia, lymphatic vessels, blood vessels, or lymph nodes and exhibit varying degrees of impeded diffusion.

In T2 shine-through, an area of high signal intensity (or low signal intensity on the inverted gray-scale images) is visible that is not due to impeded water diffusion but is caused by the long tissue T2 relaxation time. In these cases, reviewing the corresponding ADC map would enable correct interpretation (Fig. 7). For this reason, it is recommended that

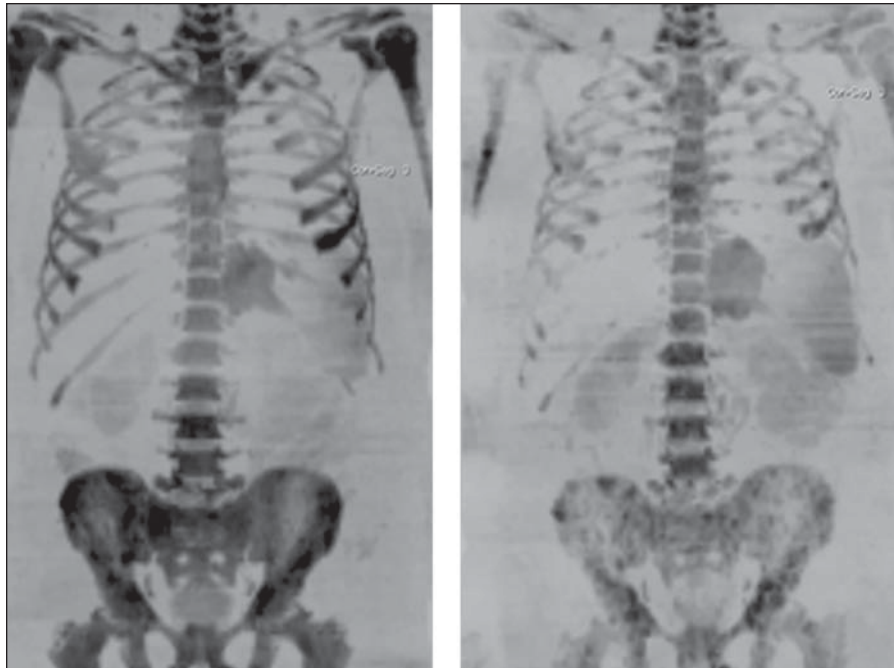


**Fig. 8**—38-year-old woman with splenic angiosarcoma. Example of possible false-negative findings.

**A**, Axial T1-weighted MR image shows several low-signal-intensity lesions (*arrows*) in spleen.

**B**, MR image ( $b = 750 \text{ s/mm}^2$ ) shows that lesions, because they originate in spleen, which normally exhibits impeded diffusion, appear less conspicuous.





**Fig. 9**—54-year-old man with diffuse multiple myeloma and extraosseous myeloma in left paravertebral region. **A**, Inverted gray-scale diffusion-weighted maximum-intensity-projection images before (*left*) and 9 months after (*right*) initiation of cyclophosphamide, dexamethasone, thalidomide, and bortezomib treatment. Image 9 months after treatment shows responding disease in bony pelvis, proximal femurs, humeri, and spine. Extraosseous disease is unchanged.

**B**, Apparent diffusion coefficient (ADC) histograms obtained over bony pelvis before (*left*) and after (*right*) treatment show mean ADC increase in bone disease and right shift of histogram distribution after treatment (*orange*) compared with before treatment (*blue*). Substantial proportion of voxels show increase in ADC values beyond 95th percentile of baseline histogram distribution (*red, top right*) after therapy. In lower left, yellow indicates ADC values between the 5th to 95th percentiles of the baseline distribution; green, below the 5th percentile; red, above the 95th percentile.

normal red marrow results in better contrast between the tumor and the yellow marrow replaced by signal-suppressed fat.

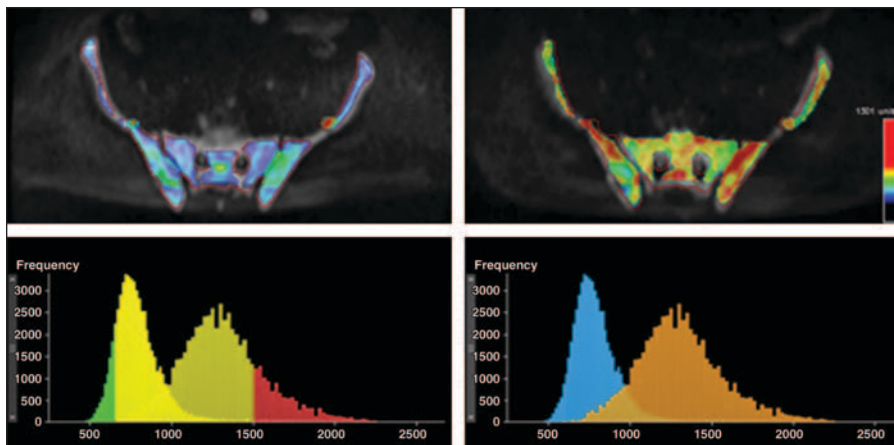
Tumors that are cystic or mucinous or have an intrinsically high ADC (e.g., clear cell renal cancer) can have relative signal suppression on high b value whole-body DW images and therefore be missed. In this regard, knowledge of the clinical history is important, particularly the histologic subtype of any known malignancy at image interpretation.

Certain areas of the body are prone to imaging artifacts, which can obscure lesions. These artifacts appear in the root of the neck, the lower central mediastinum, the left lobe of the liver, and the lungs. In addition, medical implants (e.g., silicone breast implants) and orthopedic prostheses (e.g., hip and knee replacements) can result in ghosting and susceptibility artifacts. As such, lesions lying adjacent to these areas can be missed on DW images. Lesions close to metal implants may also be obscured by artifacts.

For quantitative assessment, regions of interests can be drawn on ADC maps and the tissue diffusivity recorded. Comparison of the mean or median ADC values of the same lesion before and after treatment can be used to observe tumor response as ADC changes (usually increases) after therapy. However, the measurement reproducibility of ADC values obtained with whole-body DWI technique have not been established, particularly with regard to ADC variations across anatomic imaging stations.

### Challenges

Although whole-body DWI appears highly promising, more data are needed to confirm its utility in a variety of diseases and clinical situations. There are inherent disadvantages



**A**

**B**

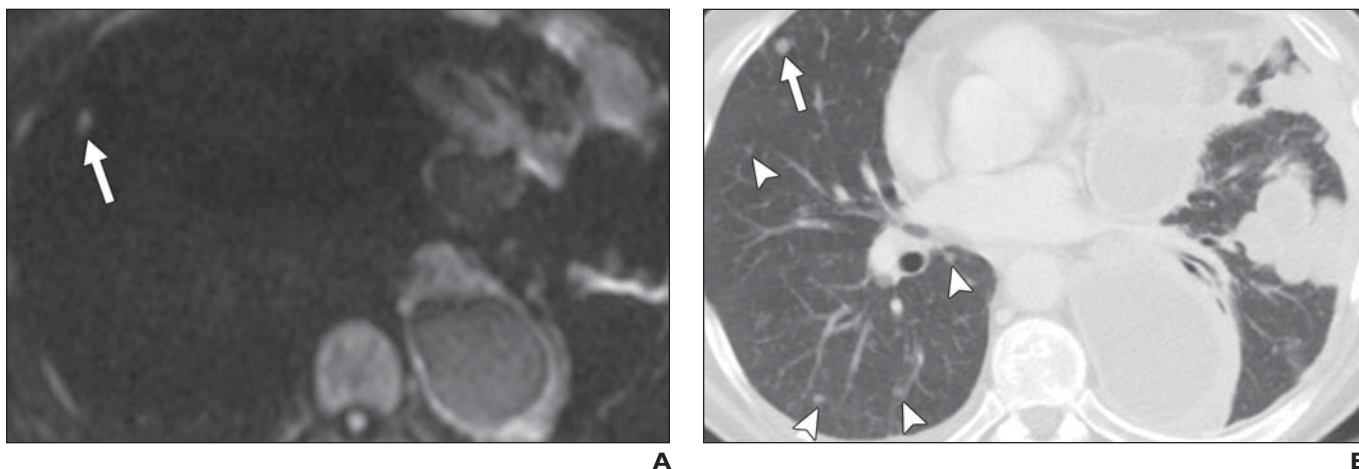
whole-body DWI not be performed with only a single high b value; a minimum of two b values is suggested.

A variety of nonmalignant conditions can exhibit impeded diffusion on DW images, mimicking malignant disease. Included are inflammatory conditions of the abdomen and pelvis, such as inflammatory bowel disease and abscesses. In particular, abscesses can have marked impeded water diffusion and return low ADC values [59]. However, the clinical presentation and symptoms are usually helpful in making the distinction. Diffusion marrow hyperplasia, which can occur after therapy (e.g., with granulocyte colony-stimulating factor), can cause spurious inter-

pretation of increasing bone marrow infiltration (Fig. 8).

**False-negative findings**—Disease in normal structures can exhibit impeded water diffusion. Normal anatomic structures—such as the salivary glands, lymph nodes, spleen, spinal cord, ovaries, testes, red marrow, endometrial lining, bowel wall, peripheral nerves, and neural ganglia—have varying degrees of impeded water diffusion at DWI. As such, disease processes that originate in or involve these structures can be missed, leading to false-negative findings (Fig. 9). For this reason, whole-body DWI studies are more likely to be useful for identification of bone metastasis in older patients, in whom the paucity of

## Whole-Body Diffusion-Weighted MRI



**Fig. 10**—45-year-old man with left mesothelioma.

**A**, Axial MR image ( $b = 750 \text{ s/mm}^2$ ) of thorax shows small metastatic lesion (arrow) in right lung.

**B**, CT image shows MRI is less sensitive than CT for detecting small metastatic lesions (arrowheads). Arrow indicates lesion in **A**.

of whole-body MRI, the most obvious being relative insensitivity to small lung lesions, the clinical implication of which has to be considered and debated (Fig. 10).

Accurate validation of whole-body DWI findings remains a challenge because focal features highlighted with the technique can be difficult to corroborate with findings with other imaging modalities and inaccessible for tissue sampling. When possible, a panel of reference studies, including PET, and follow-up imaging can be used for comparison. Even then, a level of uncertainty may remain regarding small lesions detected with whole-body DWI, which could require follow-up for accurate categorization.

Tools for qualitative and quantitative analysis of DW images are only beginning to emerge. Sophisticated methods are being explored to derive better descriptors of the tissue diffusion processes. The use of DWI to detect bone marrow metastasis and infiltration appears highly promising. However, more work is needed to understand how DWI signal intensity and ADC vary with age and sex in healthy persons and how these parameters change with treatment, which modulates marrow activity. There is also a need to understand the biologic variations of normal and diseased bone marrow with physiologic changes and with a range of treatments. A concerted effort in body DWI research would help to further establish the technique as an important tool in oncologic practice.

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