ONCOLOGY IMAGING

Whole‑body magnetic resonance imaging (WB‑MRI) in oncology: recommendations and key uses

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Abstract

The past decade has witnessed a growing role and increasing use of whole-body magnetic resonance imaging (WB-MRI). Driving these successes are developments in both hardware and software that have reduced overall examination times and significantly improved MR imaging quality. In addition, radiologists and clinicians have continued to find promising new applications of this innovative imaging technique that brings together morphologic and functional characterization of tissues. In oncology, the role of WB-MRI has expanded to the point of being recommended in international guidelines for the assessment of several cancer histotypes (multiple myeloma, melanoma, prostate cancer) and cancer-prone syndromes (Li–Fraumeni and hereditary paraganglioma–pheochromocytoma syndromes). The literature shows growing use of WB-MRI for the staging and follow-up of other cancer histotypes and cancer-related syndromes (including breast cancer, lymphoma, neurofbromatosis, and von Hippel–Lindau syndromes). The main aim of this review is to examine the current scientifc evidence for the use of WB-MRI in oncology.

Keywords Magnetic resonance imaging · Difusion-weighted imaging · Oncology · Whole-body MRI · Cancer screening · Cancer-related syndromes

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Introduction

In 1905, Albert Einstein published an article in *Annalen der Physik* describing the random motions of small particles suspended in a fuid that provided the frst quantitative theory for the natural phenomenon widely known as the *Brownian motion.* Essentially, Einstein's theory allows one to relate the difusion constant to physical quantities, such as the mean squared displacement of a particle in a given interval of time [[1\]](#page-11-0). Denis Le Bihan published the frst article displaying difusion-weighted magnetic resonance images of the central nervous system in 1985 [[2\]](#page-11-1), some 80 years after Einstein's theory on water difusion. Since then, difusion-weighted imaging (DWI) has evolved as a clinical magnetic resonance imaging (MRI) technique; it frst entered practice in neuroradiology for the assessment of cerebral ischemia [[3\]](#page-11-2) and subsequently has extended into oncological applications [\[4](#page-11-3), [5](#page-11-4)].

In 2004, Taro Takahara started applying DWI in a single, whole-body (WB) examination, thus giving birth to the WB-MRI examination as we know it today: used mostly in oncological applications to provide a combination of morphologic and difusion-weighted images from head to mid-thigh. Taking steps to ensure good background body suppression of the signal, this frst whole-body MRI (WB-MRI) with DWI examinations took almost an hour to acquire and required off-line image concatenation to produce an unifed stack of slices for visualization [\[6\]](#page-11-5). As highlighted in recent meta-analyses [\[7](#page-11-6), [8](#page-12-0)], the growing evidence regarding the use of WB-MRI in oncology has come to support recommendations of whole-body MRI in international guidelines [[9–](#page-12-1)[11](#page-12-2)] and widening clinical adoption for diferent cancer histotypes [\[12,](#page-12-3) [13](#page-12-4)].

Our aim is to review the current scientifc evidence for the use of WB-MRI in oncology.

Imaging acquisition protocol

The uptake of WB-MRI as a radiological technique is closely tied to technical developments that have contributed to achieving a good spatial resolution with good signal-to-noise ratio throughout the body, including continuous moving-table acquisitions, multi-channel surface receiver coils, and parallel imaging acquisition. These developments have dramatically reduced the scanning time for WB-MRI, such that DWI can now be performed along with supporting morphologic T1- and T2-weighted images of the whole body in a reasonably short acquisition time (30–45 min).

The anatomical coverage of a WB-MRI examination is usually from the skull base to mid-thigh analogous to positron emission tomography (PET) and computed tomography (CT) scans, though in specifc clinical contexts it may be extended from vertex to feet. Imaging is usually performed with a large field of view $(40-45 \text{ cm})$ [\[14](#page-12-5)].

Morphologic images

Due to their short acquisition times, T1-weighted images of the chest and abdomen are usually performed within breath-holds as this prevents misinterpretation due to motion artifacts. Gradient echo (GE) and Dixon techniques are increasingly used for the T1-weighted images due to their capability to derive multiple images (including in-phase, opposite phase, water, and fat images) in a single acquisition. T1-weighted imaging is usually acquired in the same plane as the DWI images to ensure a good anatomical match. Intravenous contrast agent administration with subsequent post-contrast T1 image acquisition is performed only for specifc clinical requests, such as the detection of brain metastases or characterization of liver masses.

Post-processing of the Dixon images and specifcally derivation of a fat fraction (*F*%) map that describes fat distribution are recommended [\[15](#page-12-6)]. A fat fraction map can be computed on most of the MRI post-processing consoles as:

where *F* and *W* are, respectively, the fat and water images produced by the Dixon technique.

T2-weighted images are acquired in the axial plane without fat suppression using turbo spin-echo (TSE) sequences, with half-Fourier acquisition single-shot turbo spin-echo (HASTE) being preferred. Notably, some centers do not include T2-weighted images in the WB-MRI protocol in order to limit the acquisition time as similar information is provided by low *b*-value DWI images, but at a lower spatial resolution.

T1-weighted images and T2-weighted images with fat suppression via short tau inversion recovery (STIR), acquired the spine in the sagittal plane, are usually warranted for investigation of suspected skeletal metastasis.

Difusion‑weighted images

DWI is usually performed using relatively thick sections (from 5 to 7 mm) in an axial orientation during free breathing using a single-shot spin-echo planar imaging (SSH-EPI) acquisition, to reduce both acquisition time and image distortion. Fat suppression with the STIR is strongly recommended in order to provide homogeneous fat signal suppression at larger felds of view.

In the interest of limiting examination duration, two *b*-values are usually sufficient but three or more *b*-values are likely to more precisely quantify the apparent difusion coefficient (ADC) value, as requested for the tumor response assessment. The lowest *b*-value is usually in the range from 50 to 100 $s/mm²$ to minimize perfusionrelated signal, while the highest is typically between 800 and 1000 s/mm² to allow good detection of hyper-cellular lesions with a good signal-to-noise ratio. Depending on the MR scanner homogeneity and patient size, between four and six stacks of contiguous slices (corresponding to diferent bed positions) are usually necessary to perform WB-MRI from skull base to mid-thigh.

Post-processing of the DWI data is mandatory and should consist of:

- Unification of the high *b*-value images into a single series (consecutive from superior to inferior),
- Generation of maximum intensity projections (MIPs) of the unifed high *b*-value series at small angular increments (typically 3°) rotating around the cranial–caudal axis,
- Generation of coronal multi-planar reconstructions (MPRs) of the unifed high *b*-value series,
- If dedicated T2-weighted images were not acquired, unify the low *b*-value images into a single series,
- Unification of the ADC maps into a single series (consecutive from superior to inferior).

 $100 \times F/(F + W)$

MIP reconstructions are usually displayed with an inverted gray scale, to provide a panoramic view similar to PET examinations. A complete image acquisition protocol is summarized in Table [1,](#page-2-0) and a typical example of WB-MRI imaging, including DWI, is displayed in Fig. [1](#page-4-0).

Other possible WB‑MRI protocols

Other WB-MRI protocols for oncology have been used in the studies published over the last 15 years. Their diferences can be described in terms of the following four concepts: anatomical coverage, imaging planes, MR sequences and reconstructions, and contrast administration.

The *anatomical coverage* can be extended from the vertex to the feet (including upper limbs) when evaluating bone involvement in diseases that frequently involve the long bones, such as multiple myeloma (MM) [[16](#page-12-7), [17\]](#page-12-8), when evaluating soft tissues in melanoma patients [[18,](#page-12-9) [19](#page-12-10)], as well as for cancer screening in patients with Li–Fraumeni syndrome [[20\]](#page-12-11). Extending the anatomical coverage implies longer acquisition times.

As regards *imaging planes*, many studies have made use of the coronal plane for morphologic whole-body images, alone or in combination with axial imaging [[16,](#page-12-7) [21](#page-12-12)[–29](#page-12-13)]. In some cases, this choice may allow a better trade-off between scan time and image coverage, but is also a matter of scanner performance, radiologist preference, local practices, and considerations of readability, for example, the axial skeleton [\[30\]](#page-12-14). The addition of dedicated single body part imaging is worth considering for specifc assessment of brain, liver [[31,](#page-12-15) [32](#page-12-16)], and chest [[32](#page-12-16)]. Sagittal acquisition of the whole spine is included in the majority of WB-MRI published protocols $[21-25, 27, 31-34]$ $[21-25, 27, 31-34]$ $[21-25, 27, 31-34]$ $[21-25, 27, 31-34]$ $[21-25, 27, 31-34]$ $[21-25, 27, 31-34]$ $[21-25, 27, 31-34]$, as it allows rapid acquisition and efficient evaluation of vertebral lesions, spinal cord compression, and vertebral fractures [\[26\]](#page-12-20).

The choice of the *MR sequences and reconstructions* used in WB-MRI protocols has seen considerable evolution, and while not all include DWI $[21-25, 32]$ $[21-25, 32]$ $[21-25, 32]$ $[21-25, 32]$, the majority of studies without DWI were published between 2002 and 2013, when DWI of the whole body was quite time-consuming and quality inconsistent. MR technology developments in the last few years have signifcantly improved the quality and time efficiency of DWI in large volumes, allowing its wider and easier application in WB-MRI protocols. Similarly, the use of *F*% maps that we suggest in this review for bone assessment is a recent introduction to WB-MRI and is only described in two studies published in 2017, using WB-MRI for multiple myeloma [\[17](#page-12-8)] and breast cancer [\[34](#page-12-19), [35\]](#page-12-21) patients. That earlier studies did not make use of *F*% maps is probably due to progressive acceleration of the Dixon acquisition and prior lack of experience with this application in WB-MRI.

Contrast administration is reported primarily in studies where WB-MRI was used for patients with tumors requiring brain assessment, such as melanoma [[18](#page-12-9), [31\]](#page-12-15) and lung adenocarcinoma [[27\]](#page-12-18). The administration of contrast is also described in several studies including patients with MM. However, DWI has largely replaced the administration of contrast agents in many studies performed in patients with osteotropic tumor histotypes [\[26](#page-12-20), [34–](#page-12-19)[36](#page-13-0)] and lymphoma [[29,](#page-12-13) [37](#page-13-1)].

Clinical applications in oncology

Multiple myeloma

Multiple myeloma (MM) is a hematological disorder characterized by the accumulation of neoplastic plasma cells in the bone marrow. The resulting bone disease is characterized by the presence of osteolytic lesions, osteoporosis, or bone fractures and has a signifcant impact on morbidity and mortality

Table 1 WB-MRI acquisition protocol: morphologic and difusion-weighted imaging

Images	Sequences	Image contrast	Fat suppression Plane		Acquisition mode Reconstructions	
Morphologic GE		T1	Dixon	Axial	Breath hold	F%
	HASTE	T ₂				
	TSE	T1			Sagittal Free breathing	
		T ₂	STIR			
DWI	SSE SE EPI	low b-value $50-100$ s/mm ² high b-value $800-1000$ s/mm ²	STIR	Axial	Free breathing	Unification of the high <i>b</i> -value images into a single series MIPs of the unified high <i>b</i> -value series rotating around the cranial- caudal axis Coronal MPRs of the unified high <i>b</i> -value series Unification of the ADC maps into a single series

Fig. 1 a Morphologic images acquired during WB-MRI examination. ◂The anatomical coverage is from the skull base to mid-thigh. Axial scans include multiple T1-weighted images from a single Dixon acquisition (namely in-phase, out of phase, water, and fat images) and T2- weighted images performed without fat suppression. The *F*% map is computed from the T1-weighted Dixon image data. Sagittal T1- and T2-weighted (with STIR fat suppression) images are useful for the investigation of metastases in the spine. **b** Difusion-weighted images acquired as part of a WB-MRI examination. The anatomical coverage is from the skull base to mid-thigh. Two diferent difusion weightings (with *b*-values of 50 s/mm² and 900 s/mm², respectively, in this example) are used to calculate the ADC map (third column). Maximum intensity projections of the high *b*-value images are used to perform rotational reconstructions, here displayed in frontal projection

in multiple myeloma patients. For this reason, the International Myeloma Working Group (IMWG) has affirmed that even the presence of asymptomatic bone disease on conventional radiography should be considered a criterion of symptomatic MM, requiring treatment [[38\]](#page-13-2). In a study conducted by Walker et al. involving 611 MM patients, WB-MRI detected more focal lesions than conventional whole-body x-rays in three of the most common metastatic sites for MM, including spine (78% versus 16%; *p* value 0.001), pelvis (64% versus 28%; *p* value 0.001), and sternum (24% versus 3%; *p* value 0.001) [\[39](#page-13-3)]. Similar results were observed in a prospective cohort study conducted by Baur–Melnyk on 41 newly diagnosed MM patients, in which WB-MRI showed diagnostic performance superior to conventional wholebody computed tomography (CT) in the detection of skeletal lesions (CT understaged 11/41 patients compared to MRI, *p*<0.001) [[40\]](#page-13-4). In consideration of emerging evidence, the IMWG and the British Society for Haematology (BSH) have both recommended WB-MRI for the staging of all forms of multiple myeloma (Grade A recommendation, GR A [[41\]](#page-13-5)), as well as for the follow-up of oligo-secretory and nonsecretory myelomas (Level of Evidence LE 1B) or for patients with extramedullary diseases (LE 1B) [[10](#page-12-22), [38](#page-13-2), [41](#page-13-5)]. Finally, WB-MRI is also recommended for the staging of solitary bone plasmacytoma (SBP), an early-stage malignancy with a clinical course between monoclonal gammopathy of undetermined signifcance (MGUS) and MM [\[38](#page-13-2)].

Melanoma

Although newly discovered immunological treatments for advanced melanoma have signifcantly increased overall survival, most patients with stage III–IV melanoma will still die of the disease [\[42\]](#page-13-6). Therefore, the clinical management of these patients requires frequent monitoring with a technique having good diagnostic performance in assessment of the entire body. In a prospective study conducted by Muller-Horvat et al. involving 41 metastatic melanoma patients, WB-MRI performed with contrast agent administration but

without DWI detected some 40% more lesions than wholebody CT. Moreover, treatment strategy was altered due to the WB-MRI findings in 10 $(24%)$ of the patients [[43\]](#page-13-7). In a later study involving 71 scans, WB-MRI with DWI but without contrast-enhanced scans and WB-DWI without DWI but with contrast agent were seen to have an equivalent diagnostic performance in the detection of extracranial metastases from advanced melanoma [[31](#page-12-15)]. Thus, there is growing evidence of the value of WB-MRI with DWI in both the detection and the staging of advanced melanoma, as illustrated in the cases depicted in Figs. [2](#page-5-0) and [3.](#page-5-1)

Based on these results, the German Dermatological Society (GDS), the Dermatologic Cooperative Oncology Group (DCOG), and the recently updated Swiss guidelines recommend WB-MRI for cross-sectional imaging of advanced melanoma (stage III or worse), indicating the equivalence of this method to whole-body CT and PET/CT. Moreover, WB-MRI is recommended for the follow-up of melanoma patients staged from IIC to IV [[9,](#page-12-1) [44](#page-13-8)].

Prostate cancer

Mortality rates for prostate cancer are low, despite it being the most common cancer among males in Europe [\[45](#page-13-9)]. This is tied to the fact that most prostate cancer diagnoses are made while the disease is still in an early stage, and thus of low–intermediate risk. There exists, however, a subgroup of prostate cancer patients who can be considered to be of high risk because their clinical parameters (prostate serum antigen $(PSA) > 20$ ng/mL or $> cT2c$ or with a Gleason score $GS \ge 8$) are associated with a greater probability of developing locoregional or metastatic lesions [[46\]](#page-13-10).

For the staging of high-risk prostate cancer patients, guidelines developed by the European Association of Urology (EAU) recommend at least cross-sectional abdominopelvic imaging and a bone scan (BS) (GR A, LE 2A) [\[11](#page-12-2)]. They acknowledge, however, that WB-MRI is the most sensitive imaging technique, citing a meta-analysis by Shen et al. [\[47](#page-13-11)] including 1102 metastatic prostate cancer patients, in which WB-MRI was found to be more sensitive (97%) than choline PET/CT (91%) and BS (78%) in the detection of skeletal metastasis (Fig. [4](#page-6-0)).

In metastatic castration-resistant prostate cancer (mCRPC) patients treated with enzalutamide, abiraterone, or radium-223, up to one disease progression in three is detected radiologically in the absence of clinical symptoms or PSA progression [\[15\]](#page-12-6). Furthermore, prostate-specifc membrane antigen (PSMA) PET/CT scans may fail to provide information on tumor viability during androgen receptor inhibition [[48\]](#page-13-12). A recent review conducted by Padhani et al. [[49](#page-13-13)] indicates the potential of WB-MRI to address unmet clinical needs in mCRPC patients. Similarly, the European Organization for Research and Treatment of

Fig. 2 Contrast-enhanced (gadolinium-ethoxybenzyl-diethylenetriamine pentaacetic acid, Gd-EOB-DTPA) WB-MRI with DWI performed on a patient with a stage III melanoma. Hepatobiliary phase 20 min after injection (**a**) reveals the presence of a 9 mm metastasis in the fourth liver segment (white arrow). The same lesion is clearly detectable in the high *b*-value (900 s/mm²) diffusion-weighted image performed in the same session (**b**)

Cancer (EORTC) considers WB-MRI as a "one-size-ftall" solution for evaluating treatment efficacy in advanced prostate cancer patients [[50\]](#page-13-14).

The Advanced Prostate Cancer Consensus Conference (APCCC) has confrmed that PSA alone is not reliable for monitoring disease activity in mCRPC, suggesting the use of a robust imaging technique before deciding to start a new line of treatment. In this respect, their guidelines recognize the superior diagnostic performance of WB-MRI relative to CT and BS in the detection and assessment of skeletal metastasis [[51\]](#page-13-15), though they note limited availability of the WB-MRI technique.

Lymphoma

Fluorodeoxyglucose (FDG)-PET/CT is the imaging technique recommended for the most common lymphomas,

Fig. 3 Follow-up evaluation of a patient with stage IV melanoma using low-dose CT of the lung and WB-MRI. A subcentimetric metastasis in the left inferior lobe of the lung is detected on the axial CT image (arrow in **a**) as well as axial T1-weighted image (arrow in **b**)

including difuse large B cell lymphoma (DLBCL), follicular, and Hodgkin subtypes, all of which are usually characterized by a high glucose metabolism and thus likely to be FDG-avid histotypes [\[52](#page-13-16)]. Consistent with this, the National Comprehensive Cancer Network (NCCN) recommends serial CT or PET/CT examinations for both the staging and follow-up of lymphoma patients [\[53](#page-13-17)]. In lymphoma subtypes that exhibit variable FDG avidity or non-FDG avidity however, FDG-PET/CT may be inefective. The sensitivity of WB-MRI with DWI to hyper-cellular lesions, independent of glucose metabolism, allows a reliable radiological evaluation of these lymphoma subtypes [[54\]](#page-13-18).

In a prospective study conducted by Mayerhoefer et al. [[13](#page-12-4)] on 140 patients, WB-MRI with DWI demonstrated better sensitivity (94.4%) than FDG-PET/CT (60.9%) and contrast-enhanced CT (70.7%) in staging patients with lymphoma subtypes of variable FDG avidity. (The majority were MALT lymphomas.) Further, the same group found WB-MRI with DWI to have a diagnostic performance similar to FDG-PET/CT and CT in FDG-avid lymphomas [\[55](#page-13-19)].

A growing application of WB-MRI in lymphoma involves young patients irrespective of histotype. In a

Fig. 4 A patient admitted with an initial diagnosis of prostate adenocarcinoma (GS $5+4$, biopsy positive in 8/8 cores), $PSA = 29$ ng/ mL, positive digital rectal examination (cT2c). The systemic staging, previously assessed with contrast-enhanced CT examinations of chest, abdomen, and pelvis (CT-CAP) and BS, was negative for metastatic disease. Subsequent systemic staging with WB-MRI with DWI showed two metastases, one in the anterior arch of a left rib (arrow) and the other at level of retroperitoneal lymph node (arrowhead). A multiparametric MRI examination performed in the same session showed a PI-RADS fve lesion (dashed line), visible on T2-weighted axial images, high *b*-value difusion-weighted images, and ADC map

recent study by Brenner et al., the overall 10-year survival for lymphoma patients younger than 35 years of age was seen to be above 90% (94.7% for patients $<$ 24 years and 89.4% for patients comprised between 25 and 35 years, respectively) [\[56\]](#page-13-20). Despite this long survival, the NCCN guidelines still recommend PET/CT or CT for the staging and follow-up of lymphoma patients [[57\]](#page-13-21), with 6–9 examinations to be performed in the frst two years after diagnosis. This is particularly high exposure to ionizing radiation for young patients with a long life expectancy. In justifying the choice of imaging technique, one should consider the growing evidence of equivalent diagnostic performance of WB-MRI in respect of PET scans for both staging and follow-up of lymphoma patients [[13](#page-12-4), [58](#page-13-22)[–65\]](#page-13-23) (Fig. 5). As summarized in Table [2,](#page-7-1) the kappa coefficient of agreement observed between these two imaging techniques is typically in the range from 0.68 to 1.00.

Breast Cancer

In the last two decades, conservative treatments and early detection have substantially improved the prognosis for patients with low-stage breast cancer (BC) [[66](#page-14-0)]. Nevertheless, according to the American Cancer Society, 5-year overall survival for patients with advanced (stage IV) BC remains unfavorably low at just 22% [\[67\]](#page-14-1). A recent epidemiological study by Kwast et al. including 25,336 women with newly diagnosed BC demonstrated bone to be the most common site of metastatic spread, regardless of tumor histological characterization. Notably, bone metastases were present at the frst staging procedure in 49.7% of invasive ductal cancer (IDC) and in 61.7% of invasive lobular cancer (ILC) subtypes, respectively [[68](#page-14-2)]. The widely accepted Response Evaluation Criteria in Solid Tumors (RECIST) are not suited to a proper assessment of bone metastases and in fact consider them to not be measurable. Even the revised RECIST 1.1 version does not fully address the need to assess bone metastases. According to the new criteria, bone metastases are considered measurable only once they have spread to the surrounding soft tissue with an extent larger than 10 mm in diameter, which rarely occurs in clinical practice [[69](#page-14-3)]. This situation points to a signifcant unmet clinical need for means to evaluate bony metastases, with implications beyond breast cancer.

In a 2011 meta-analysis by Yang et al. [[70](#page-14-4)] covering 145 studies with 15,221 metastatic cancer patients, WB-MRI showed diagnostic performance comparable to PET and superior to CT and BS in the detection of skeletal metastases (Table [3\)](#page-8-0). In addition, two recent studies emphasize the potential of WB-MRI in advanced breast cancer. Comparing the fndings of 210 paired WB-MRI and CT-CAP for the follow-up of metastatic BC patients, Kosmin et al. observed that out of the 46 treatment changes made due to progressive disease (PD) detected at imaging, in 34.7% of the cases (16 paired examinations), PD was visible only on WB-MRI and not on the CT-CAP examinations [\[12\]](#page-12-3). Similarly among 40 cases of progressive disease fndings in a group of 58 patients who had both WB-MRI and CT-CAP or an 18 F-FDG-PET/CT within 8 weeks, Zugni et al. [[71\]](#page-14-5) found that all 40 (100%) have been identifed on WB-MRI but only 23 (58%) were also identified by CT-CAP or ${}^{18}F$ -FDG-PET/ CT.

WB-MRI has also shown promising diagnostic performance and established itself as a safe and accurate diagnostic imaging method for the systemic staging of pregnant BC patients; some 40% of whom receive a diagnosis when BC is already at an advanced stage [\[72](#page-14-6)]. In this setting, accurate systemic staging of disease using imaging techniques that avoid ionizing radiation and contrast agent administration is to be preferred (Fig. 6).

Fig. 5 A 22-year-old woman with relapse of non-Hodgkin lymphoma (Ann Arbor stage IVb). After multiple lines of systemic treatments FDG-PET/CT and WB-MRI were performed before and 6 months after initiating chemotherapy under the ESHAP scheme and periph-

eral blood stem cell transplantation. Response assessment with both PET/CT and WB-MRI at the same time points reveals a complete resolution of the abnormal FDG uptake across all body regions, consistent with complete response (CR)

Table 2 Level of agreement observed between WB-MRI and PET/CT techniques for staging not-FDG avid and variable FDG avidity lymphoma subtypes

References	\boldsymbol{N}	Lymphoma histotypes	Kappa coefficient	Agreement	P value
Mayerhoefer et al. [13]	140	FDG-avid/non-FDG-avid lymphomas	0.92/0.89	Excellent	< 0.0001
Albano et al. [58]	104	HL. NHL	0.93	Excellent	< 0.01
Albano et al. [59]	68	HL, NHL	0.88	Excellent	< 0.05
Abdulgadr et al. [60]	31	HL, aggressive NHL, indolent NHL	0.87	Excellent	< 0.0001
Stèphane et al. [61]	23	HL, DLBCL, NHL	1.00	Excellent	< 0.0001
Van Ufford et al. [62]	22	HL. NHL	0.68	Good	< 0.0001
Gu et al. $[63]$	17	HL, NHL	0.82	Excellent	< 0.0001
Lin et al. $[64]$	15	DLBCL	0.85	Excellent	< 0.0001
Wu et al. [65]	8	DLBCL	1.00	Excellent	< 0.0001

 N =number of patients, Kappa coefficient was considered corresponding to the following levels of agreement <0 = no agreement, 0–0.20 = poor agreement, 0.21–0.40=fair agreement, 0.41–0.60=moderate agreement, 0.61–0.80=good agreement, and 0.81–1.00=excellent agreement

Finally, in the last decade the use of WB-MRI has seen progressive growth in the management of BC patients having an ILC subtype [[35](#page-12-21)]. ILC is statistically more likely than IDC to spread to the gastrointestinal organs, the peritoneum and retroperitoneum, the gynecological system, and the pleura (ILC 20% and IDC 8.4%, respectively) [\[68](#page-14-2)], which are notoriously challenging body regions to explore using PET/CT and CT techniques. This may in part be because ILC typically follows a pattern of difuse infltration characterized by "Indian fle" neoplastic growth related to a reduced (or absent) E-cadherin membrane expression that reduces cell-to-cell adhesion and facilitates permeation along tissue planes [\[73\]](#page-14-7). Further, metastases from ILC are less FDG-avid than other BC histotypes and therefore less visible on FDG-PET examinations. Due to the aspecifc nature of difusion-weighted images (hypercellular lesions are always visible at high *b*-values, regardless of the glucose metabolism), WB-MRI with DWI has the capability to depict the presence of neoplastic spread into gastrointestinal organs (Fig. [7](#page-9-0)).

Table 3 Comparison of diagnostic performances (sensitivity and specificity) between WB-MRI, CT, PET, and BS in the detection of bone metastases (based on the meta-analysis in [[70](#page-14-4)])

	WB-MRI $(\%)$	CT(%)	PET $(\%)$	BS(%)
Per-patient analysis				
Sensitivity	90.6	72.9	89.7	86.0
Specificity	95.4	94.8	96.8	81.4
Per-lesion analysis				
Sensitivity	90.4	77.1	86.9	75.1
Specificity	96.0	83.2	97.0	93.6

Values reported on a per-patient and per-lesion analysis, respectively

Fig. 6 WB-MRI for the systemic staging of disease in a 37-year-old woman with locally advanced BC (cT2N1) at 31 weeks of gestation. Multiple skeletal metastases (arrows) in spine, sternum, and pelvis are visible on both morphologic (T1-weighted sagittal TSE) and high *b*-value difusion-weighted images (MIP). Several pathological lymph nodes are visible in the right axillary region on T1-weighted gradientecho Dixon in-phase image (white arrow). High-intensity regions visible in the abdomen (dashed line) on high *b*-value difusion-weighted images correspond to the fetal kidneys and brain. Based on the WB-MRI examination, the patient underwent an early cesarean delivery and was subsequently treated with both chemotherapy and hormonal therapy. The child did not experience adverse events during the prenatal or development periods

Other cancer histotypes

In the last few years, several articles have clearly demonstrated WB-MRI to have excellent diagnostic performance in staging and monitoring of other cancer histotypes. Lung cancer, for example, is usually staged by PET/CT together with dedicated brain MRI examinations, but in a prospective study conducted by Usuda et al. including 81 non-small cell lung cancer (NSCLC) patients, WB-MRI with DWI achieved a diagnostic performance similar to PET/CT and brain MRI in pre-surgical staging. Moreover, they found WB-MRI to provide higher diagnostic accuracy (87.6% 71/81) than PET/ CT (82.7% 67/81) for the staging of hilar and mediastinal lymph nodes (N) [\[74](#page-14-8)]. Similar results emerged from a study by Ohno et al. involving 96 postoperative NSCLC patients that compared WB-MRI with DWI to PET/CT in the assessment of local recurrence and distant metastases. Here, WB-MRI with DWI exhibited lower sensitivity (88.2%) than PET/CT (100%), but superior specifcity (100% and 81%, respectively) in the assessment of regional and distant metastases in NSCLC patients [\[75](#page-14-9)].

WB-MRI with DWI has also proven effective for the staging and preoperative assessment of ovarian cancer. In a prospective cohort study by Michielsen et al. that included 161 patients suspected of having ovarian cancer, WB-MRI with DWI showed a signifcantly superior diagnostic accuracy (93%) compared to CT (82%) for determining the malignant nature of the ovarian mass. Furthermore, WB-MRI with DWI was superior to CT in the detection of cancers having non-ovarian origin (WB-MRI+ DWI = $26/3281\%$, $CT = 10/32$ 31%, *p* value = 0.001) and to assigning the correct FIGO stage to ovarian cancers (WB-MRI+DWI=82/94 87%, CT=33/94 35%) [\[76](#page-14-10)].

Another promising result has emerged from a study conducted by Gorelik et al. [[77\]](#page-14-11) on 33 myxoid liposarcoma (MLS) patients, in whom WB-MRI depicted extrapulmonary metastases from MLS that were not visible at CT in 79% of the cases (7/9 metastatic patients).

Cancer screening

The excellent diagnostic performance of WB-MRI in tumor detection (overall sensitivity 90%) with low false-positive and false-negative rates is manifested in a number of metaanalyses (Table [4](#page-9-1)) [\[7](#page-11-6), [8,](#page-12-0) [47](#page-13-11)]. This, together with the shortening of examination times achieved thanks to modern acquisition protocols and hardware, has extended its application to cancer screening programs. Several articles have demonstrated the potential of WB-MRI in the early detection of cancer in subjects with Li–Fraumeni syndrome (LFS), a highly penetrant, cancer-prone syndrome, caused by germline mutations of the TP53 gene that predispose carriers to

Fig. 7 In a 51-year-old metastatic ILC BC patient, tumor marker progression (CA $15-3=660$ U/mL) occurred during third-line chemotherapy treatment. Contrast-enhanced CT of chest and lung for suspected disease progression showed the absence of secondary lesions. A thickening of the lateroconal and left anterior renal fascia (white arrow) was described as stable compared to the previous controls and interpreted as the result of a previous surgical intervention for ovarian serous cystadenoma. A suspicious thickening of the left anterior renal fascia and the descending colon wall was visible WB-MRI

examination performed within a few days of the CT: the colon fnding is recognizable on high *b*-value difusion-weighted images (axial and MIP images), ADC map, and morphologic in-phase T1-weighted gradient-echo Dixon images (white arrowheads). MIP images of the high *b*-value images showed further metastatic lesions in descending colon (black arrowhead) and subcentimetric skeletal metastasis in the spine and pelvis (black arrows). All fndings were confrmed at the following WB-MRI examinations

Table 4 Principal meta-analyses comparing the diagnostic performances of WB-MRI with PET in the assessment of primary and metastatic disease

N number of patients, *SE* sensitivity, *SP* specifcity

a variety of cancers, including sarcoma, BC, adrenal gland carcinoma, and leukemia [\[78\]](#page-14-12). Around 50% of individuals with LFS will develop cancer before reaching 30 years of age, and the prevalence of LFS-related cancer rises to 95% in female subjects older than 60 years [\[79\]](#page-14-13). In a study by Bojadzieva et al. [[80\]](#page-14-14) including 63 TP53 mutation carriers, WB-MRI with contrast agent disclosed the presence of six asymptomatic tumors and one recurrent metastatic cancer, resulting in a cancer rate of 13.2%. Similar observations were found in the preliminary results from the UK SIGNIFY study, involving 44 TP53 germline mutation carriers, where WB-MRI detected the presence of four cancers that would

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Table 5 Main studies of WB-MRI for cancer screening in asymptomatic individuals, sorted by number of enrolled subjects

References	Subjects	cancer	Positive for Cancer rate $(\%)$
Ballinger et al. [20]	578	39	6.7
Mai et al. [78]	116	5	4.3
Villani et al. [82]	59	15	25.4
Bojadzieva et al. [79]	53	7	13.2
Saya et al. $[80]$	44	4	9.1
Anupindi et al. [83]	24		4.2

Cancer rate is computed as the percentage of subjects with at least one suspected or diagnosed malignancy out of the enrolled subjects

be life-threatening if left untreated, in 5 LFS subjects (cancer rate of 9.1%) [\[81](#page-14-15)]. A meta-analysis recently published has validated the frst statistically robust estimate of the clinical utility of WB-MRI in screening TP53 mutation carriers [\[20](#page-12-11)]. Table [5](#page-10-0) summarizes the main results from available cohort studies including WB-MRI examination for cancer screening in subjects with TP53 mutations [[20](#page-12-11), [79](#page-14-13)[–81](#page-14-15), [83](#page-14-16), [84](#page-14-17)].

On the basis of the above evidence, the NCCN and the American Association for Cancer Research (AACR) have recently included annual WB-MRI examination (along with contrast-enhanced brain MRI and breast MRI for women) in their recommendations for cancer screening of individuals with LFS [[85,](#page-14-18) [86\]](#page-14-19).

Similarly, WB-MRI is now recommended for the baseline surveillance of subjects with hereditary paraganglioma–pheochromocytoma syndromes (HPP), genetic disorders that manifest with rare and usually benign tumors that originate in the nervous system. Aside from paragangliomas and pheochromocytomas, subjects with HPP syndromes can also develop cancers characterized by aggressive behavior, including renal cell carcinoma (RCC), gastrointestinal stromal tumors (GIST), and other rare tumor types [\[87](#page-14-20)]. AACR guidelines recommend biennial screening with WB-MRI for the early detection of cancer onset in subjects with HPP older than 6/8 years of age [[88](#page-14-21)]. Guidelines including WB-MRI for cancer screening in genetically predisposed subjects are also under development by the Response Evaluation in Neurofbromatosis and Schwannomatosis International Collaboration (REiNS) group for early detection of cancer in neurofbromatosis (NF)-related disorders [\[89](#page-14-22)].

Lastly, WB-MRI is starting to see use in cancer screening in those with von Hippel–Lindau syndrome (VHL), a rare autosomal dominant disorder caused by germline mutations of human chromosome 3p25 [[90\]](#page-14-23). Although MRI examinations of diferent body regions are already included in many institutional screening protocols for VHL subjects, no formal consensus exists as to the screening technique of choice for this rare genetic disease.

Table 6 Main studies of WB-MRI for cancer screening in LFS subjects, sorted by number of enrolled subjects

References		Subjects Positive for cancer Cancer rate	
Bamberg (GNC) et al. [96]	30,000	Ongoing	Ongoing
Hegenscheid et al. [90]	2500	$62*$	2.48%
Cieszanowski et al. [95]	666	7	1.05%
Goehde et al. [91]	298	1 ^a	0.33%
Lo et al. $[94]$	132	2	1.5%
Ulus et al. $[92]$	118	2	1.7%
Tarnoki et al. [93]	22		4.5%

Cancer rate is computed as the percentage of subjects with at least one suspected or diagnosed malignancy out of the enrolled subjects ^aHistological confirmation not available in all subjects, GNC=Ger-

man National Cohort

In the last decade, the application of WB-MRI to the early detect of cancer in asymptomatic subjects has been the subject of a number of articles (Table [6](#page-10-1)) [\[91](#page-14-24)[–97\]](#page-14-25). The frst, by Lo et al. [[95\]](#page-14-26), included 132 doctors from the Hong Kong Sanatorium and Hospital who underwent unenhanced WB-MRI without DWI revealed two cancers (one bronchoalveolar carcinoma and one RCC) in two (1.5%) of the participants. Similar results were observed by Cieszanowski et al. who, in a retrospective study of WB-MRI without contrast agent administration that included 666 asymptomatic individuals, found 9 malignant or possibly malignant cancers in 7 individual, corresponding to a cancer rate of 1.05% [[96](#page-14-27)].

The growing interest in the potential of WB-MRI for cancer screening in asymptomatic subjects has led to its inclusion in some large cohort studies, including the German National Cohort (GNC) and UK Biobank. The German Ministry for Education and Research and other local research institutions initiated the GNC in 2015 with the aim of establishing one of the world's largest imaging data repositories, with a target of obtaining some 30,000 WB-MRI in asymptomatic subjects [\[97](#page-14-25)]. Similarly, the UK Biobank, a research project promoted by numerous public and private British research institutions, seeks to create the largest collection of WB-MRI scans along with a range of clinical data with the purpose of promoting the study of a wide range of diseases, including cancer, in the general population [\[98](#page-15-0)].

Pitfalls and limits of WB‑MRI

The combined analysis of both morphologic and functional MR images is aimed at overcoming the limitations related to analysis of a single image type in whole-body MRI examinations, especially in patients with bone marrow involvement [[99\]](#page-15-1).

There are a number of possible sources of false-positive and false-negative fndings when reviewing DWI images alone. False-positive fndings in the bone marrow may be related to hemangiomas, focal areas of red bone marrow, bone marrow edema, and hyperplasia [[100](#page-15-2)]. False-positive fndings in soft tissues may be related to small signal abnormalities attributable to slow fuid motion in vessels, ganglia, or infammatory processes such as infammatory bowel disease or abscesses [[101](#page-15-3)]. Importantly, the appearance of healthy lymph nodes can overlap with that of pathologic lymph nodes. The combined analysis of DWI with ADC maps and morphologic T1 and T2 images is, therefore, strongly recommended [\[101](#page-15-3)].

False-negative fndings in bone marrow may be related to minimal disease infltration or to the invisibility of hyperintense lesions within bone marrow hyperplasia [[102\]](#page-15-4) in young patients or during bone marrow stimulating treatment; in some cases, this limitation can be overcome by combined review of morphologic T1 images and *F*% maps. Falsenegative fndings in soft tissues may occur within organs that physiologically present impeded difusion, such as the central nervous system, salivary glands, spleen, and lymph nodes [[5,](#page-11-4) [101\]](#page-15-3). Furthermore, it should be kept in mind that high signal intensity due to impeded difusion might be suppressed in tumor histotypes characterized by mucinous or cystic structure [[101\]](#page-15-3). Other false-negative fndings in DWI may be related to imaging artifacts, including ghosting, poor fat suppression or magnetic susceptibility efects, that are most common in the neck, lungs, mediastinum, and left liver lobe [[101](#page-15-3)]. As well, lesions in the skull might not be visible due to proximity of the brain [[100\]](#page-15-2).

Some pitfalls in image interpretation have been described in the assessment of response to therapy in patients with bone metastases [\[103\]](#page-15-5). When bone metastases respond to cytotoxic treatments or to radiation therapy, for example, an increase in bone marrow water content related to massive cell death can be observed, resulting in difuse signal intensity reduction in T1 images. This fnding might be wrongly attributed to disease progression (termed "T1-pseudoprogression") when not correlated with the appearance of bone metastases in DWI and ADC maps. Similarly, successfully treated bone metastases might show increased signal intensity in high b-value DWI images, despite showing a marked increase in ADC values; this pattern is known as "T2 shinethrough" and can be recognized by combined review of DWI and corresponding ADC maps. Finally, bone metastases may show reduced signal intensity in high *b*-value images while maintaining stable ADC values; a fnding associated with sclerotic evolution of the metastases, representing either progression (sclerotic progression) or response (sclerotic response) of the disease. This combination of observations should be regarded as indeterminate and requires the radiologist to carefully review morphologic T1 weighted or *F*% images, in order to correctly recognize the underlying pattern.

Conclusion

A number of oncological applications for WB-MRI are already well established and supported by clinical evidence. This evidence has led to guidelines that clearly indicate the central role of WB-MRI with DWI in the clinical management of several cancer histotypes and highlight its potential in regard to others. The lack of radiation exposure and the absence of contrast agent administration in a typical WB-MRI examination with DWI, together with good diagnostic performance, contribute to its attractiveness in application for cancer screening in both cancer-prone syndromes and asymptomatic subjects. Further work is needed to standardize acquisition protocols and interpretation practices, in order to reduce the variation in diagnostic performance. Some initial effort in this direction has been reported $[15]$ $[15]$, but large-scale studies are required.

Compliance with ethical standards

Conflict of interest Summers is part owner of Company QMRI Tech, which offers consulting services in MRI. Author Summers receives consulting fees from Company ASC, Italia. All other authors declare to have no confict of interest.

Ethical approval This article does not contain any studies with human participants or animals performed by any of the authors.

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