Review

Advanced Imaging in Multiple Myeloma: New Frontiers for MRI

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Abstract: Plasma cell dyscrasias are estimated to newly affect almost 40,000 people in 2022. They fall on a spectrum of diseases ranging from relatively benign to malignant, the malignant end of the spectrum being multiple myeloma (MM). The International Myeloma Working Group (IMWG) has traditionally outlined the diagnostic criteria and therapeutic management of MM. In the last two decades, novel imaging techniques have been employed for MM to provide more information that can guide not only diagnosis and staging, but also treatment efficacy. These imaging techniques, due to their low invasiveness and high reliability, have gained significant clinical attention and have already changed the clinical practice. The development of functional MRI sequences such as diffusion weighted imaging (DWI) or intravoxel incoherent motion (IVIM) has made the functional assessment of lesions feasible. Moreover, the growing availability of positron emission tomography (PET)–magnetic resonance imaging (MRI) scanners is leading to the potential combination of sensitive anatomical and functional information in a single step. This paper provides an organized framework for evaluating the benefits and challenges of novel and more functional imaging techniques used for the management of patients with plasma cell dyscrasias, notably MM.

Keywords: multiple myeloma; plasma cell dyscrasia; diffusion weighted imaging; whole body MRI

1. Introduction

Multiple myeloma (MM), the second most prevalent hematopoietic malignancy, is a plasma cell dyscrasia where monoclonal plasma cells infiltrate bone marrow, resulting in marrow failure and/or bone destruction, mainly in the axial skeleton and proximal appendicular bones [1,2]. MM exists on the end of a spectrum of disease progression, from an asymptomatic premalignant state, namely monoclonal gammopathy of undetermined significance (MGUS), to smoldering multiple myeloma (SMM), and finally to symptomatic MM, commonly with MM-associated end-organ damage, also known as CRAB (hyperCalcemia, Renal failure, Anemia, and Bone disease) [3,4].

Currently, MM is identified and staged according to the International Myeloma Working Group (IMWG) criteria. IMWG criteria outlines the MM diagnosis as clonal bone marrow plasma cells > 10%, or biopsy-proven bony or extramedullary plasmacytoma, in addition to one or more of the following: evidence of end organ damage (hypercalcemia, renal insufficiency, anemia and/or osteolytic bone lesions) and/or biomarkers positive for malignancy (serum-free light chain ratio, and clonal plasma cell proportion), and the presence of more than one focal marrow lesions on MRI [5]. There are several limitations to some of the tests involved in these criteria. The biomarkers outlined above are of limited
value in non-secretory or extramedullary MM cases [3–5]. While bone marrow biopsy is the gold standard diagnostic test and is required for the assessment of treatment response, it is invasive, with a risk of local hemorrhage and infection, and it is not representative of the whole spectrum of the marrow [6].

Earlier in 2009, the consensus statement of the IMWG recommended conventional radiography (CR) as the standard imaging modality to stage MM, both in new cases and for relapsed patients, being widely available and inexpensive [5]. However, radiographs lack sensitivity and specificity, and they poorly detect extra-osseous lesions and diffuse medullary invasion. Therefore, skeletal surveys have been replaced with more sophisticated imaging techniques (computed tomography (CT), magnetic resonance imaging (MRI), and positron emission tomography (PET)) in many tertiary centers [7,8]. Newer staging systems incorporate more sensitive imaging modalities, such as whole-body MRI (WB-MRI) or \(^{18}\)F-fluorodeoxyglucose (FDG) PET/CT into the staging system [9]. The need for whole-body imaging in MM is related to the fact that myelomatous lesions can potentially affect any bony segment in the body.

Once it has been diagnosed and staged, as with most other malignancies, MM treatment management is also dependent on bone imaging to determine the extent of tumor cell burden, both for prognostic stratification and to assess post-therapy changes [10,11]. This paper describes available advanced imaging techniques in an assessment of MM, with a greater focus on whole-body diffusion weighted imaging (WB-DWI).

2. MRI

2.1. Conventional Whole-Body MRI (WB-MRI)

According to IMWG guidelines, MRI should be considered as the complementary adequate imaging modality for the diagnosis of MM, as it is more sensitive than FDG-PET/CT [12,13]. As opposed to computed tomography (CT) or positron emission tomography (PET), an advantage of MRI is that it causes no radiation exposure. In contrast to \(^{18}\)F-FDG PET/CT, MRI is widely available, relatively faster, has no pre-scan diet requirement, and is not reliant on the metabolic activity of tumor cells. This is specifically important in the case of MM, as frequent follow-up imaging is necessary, and survival time is increasing in light of recent therapeutic advancements. Conventional MRI can relatively accurately measure the size and extension of MM pathology, but it takes longer to interpret and requires more training to read than other imaging modalities [14–17].

Nearly half of MM lesions may be missed when imaging only covers the spine [14]. Thus, for a more sensitive assessment, MRI should cover the entire axial skeleton and the proximal appendicular skeleton, defining the idea of the “whole body” MRI technique. WB-MRI has gained popularity since the last decade for diagnosing and assessing the treatment response in MM by providing morphological information on tumor spread. Around one-third to a half of MM patients show diffuse infiltration or focal deposits on WB-MRI [18,19].

The Myeloma Response Assessment and Diagnosis System (MY-RADS) comprehensively characterizes the myeloma state at diagnosis, initiation of treatment, and during follow-up, as the disease course changes in response to therapy. MY-RADS recommendations will help to improve response assessments by increasing standardization, and by decreasing the variations seen in the acquisition, interpretation, and reporting of whole-body MRI. For response assessments, the classified response assessment category (RAC) is according to anatomical regions. For each region, the RAC should use a five-point scale as follows: (1) highly likely to be responding; (2) likely to be responding; (3) stable; (4) likely to be progressing; and (5) highly likely to be progressing. MY-RADS functions to categorize patients with regard to specific disease patterns to aid in clinical trial stratification [20].

Different MRI sequences are being used based on imaging protocols of the radiology department [21–24]. Fat-suppressed T2-weighted (FS T2W), Short Tau Inversion recovery (STIR), and T1-weighted (T1W) MRI sequences are most commonly utilized for MM [25–27]. The signal intensity on FS T2W and STIR images correlates with the plasma cell concen-
tration in the bone marrow. However, it lacks enough accuracy to differentiate between hyperplastic/red marrow and myeloma marrow. Changes on T1W images occur relatively late in the course of disease, and might differentiate MM from MGUS and SMM, although to a lesser extent compared to FS T2-WI, but they offer increased specificity [28]. While the T1W, STIR, and T2W sequences are more frequently used, diffusion weighted imaging (DWI) is the most promising MRI sequence for distinguishing MM from MGUS and SMM through visual assessment, and for the qualitative evaluation of lesion activity and treatment response [28].

2.2. Whole-Body Diffusion Weighted Imaging (WB-DWI)

DWI is a functional MRI sequence that can quantitatively evaluate tissue cellularity by measuring the random thermal movements of water molecules (Brownian motion) using the apparent diffusion coefficient (ADC) map [29]. Although DWI MRI was limited to brain imaging for many years due to its sensitivity to motion, its use has been recently expanded to include other anatomic locations, due to the introduction of echoplanar imaging and the use of fast acquisition techniques capable of capturing images during breath holds [30]. Thus, DWI MRI, including WB-DWI, have revolutionized the assessment of myeloma lesions from simple evaluations based on size to quantitative data based on free water molecule movement and tissue cellularity [29,31].

From 2010 on, the utilization of WB-DWI in MM has steadily increased [30–33]. WB-DWI can be performed relatively rapidly with low technical and operational efforts when added to the standard WB-MRI protocol. WB-DWI keeps the aforementioned advantages of WB-MRI, and adds further details to morphological imaging, as compared to the conventional sequences of WB-MRI during the assessment of treatment response and extramedullary disease, which are critical for MM management [21].

ADC mapping, which is derived from DW images, can distinguish bone marrow changes in active myeloma from those in remission, providing clinically relevant data on tumor viability [34]. ADC values above 600–700 µm²/s in a non-treated and newly diagnosed patient with multiple myeloma could be used to increase confidence for the diagnosis of diffuse marrow involvement, while normal marrow ADC value mostly falls below 600 µm²/s, with even lower values in elderly patients with prominent fatty marrow [32,35–37]. Koutoulidis et al. reported a higher ADC value for diffuse MM patterns in imaging comparing to focal lesions, and they found that an ADC value of >548 µmm²/s shows 100% sensitivity and 98% specificity for comparing a diffuse pattern of myeloma infiltration, than normal marrow [36]. Messiou et al. reported a significant decrease in the ADC values of MM patients who were responders from 4 weeks to 20 weeks after treatment, while patients with stable or progressive disease did not show a significant decrease in ADC value within the same time period [35].

If cellularity was the dominant factor in determining the ADC value in myeloma marrow, there should be a negative correlation between the ADC value and marrow cellularity. However, this is not the case. For this reason, cellularity is not the main factor affecting the DWI-ADC image signal in MM. Recent studies on liver fibrosis and pancreatic cancer have suggested that the perfusion effect on the measured ADC value dominates over hypercellularity [30,38,39]. This has been suggested for MM as well [40], as the ADC number decreases with hypercellularity and increases with hypervascularity, which parallels hypercellularity in the myeloma marrow [30]: the net effect is an increase of the ADC value in MM relative to the normal reference. This explains the Intravoxel Incoherent Motion (IVIM) MRI advantage for MM diagnosis/monitoring, which will be discussed in the following section. Employing lower b-values offer a better Signal to Noise Ratio (SNR), whereas higher b-values are more accurate for detecting MM lesions (Figures 1 and 2) [41]. Most of the studies on DWI in MM have been conducted with a high b-value of 600–800, which offers a good MM lesion detection rate at a reasonable SNR [42–44].
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Figure 1. A 45-year-old man with hyposecretory IgG kappa multiple myeloma (MM) and a history of autologous stem cell transplantation 3 years before imaging. Patient had received radiation therapy to L1 vertebral body 2 years ago, and has been on maintenance therapy (carfilzomob, pomalidomide, and dexamethasone) for 6 months. WB-MRI images failed to demonstrate any osseous lesion, but showed a 12 mm extramedullary lesion deep to right parotid gland (arrow) on coronal high b-value (b = 800) DWI MIP image (A). Fused coronal DWI MIP and T1-weighted Dixon image (B) confirms the finding (arrow). Axial ADC map (C) confirms diffusion restriction (mean value of $0.8 \times 10^{-3} \text{ mm}^2/\text{s}$). Note areas of marrow infarction at bilateral proximal tibial metaphysis (arrowheads, (B)).

One of the main advantages to WB-DWI is that it provides an excellent visual contrast between the normal marrow and bone marrow lesions, differentiating them with a higher sensitivity than the conventional MRI sequence, radiologic skeletal survey, or PET/CT scan [45–47]. For example, lesion conspicuity is greater in DWI as compared to the conventional T1-MRI and STIR sequences, and has a higher lesion detection rate compared to PET/CT [48,49].

DWI was able to identify 11% more patients than PET/CT in a cohort of 227 patients, and had a sensitivity of 77% versus 47% in PET/CT compared to a conventional MRI and CT in a smaller study of 24 patients [50]. WB-DWI also offers an excellent interobserver agreement for quantifying the disease burden in MM, both for the whole-body assessment and regional evaluation for any location across the body [48,51]. This superiority offers a substantial impact on treatment planning and patient classification, and it allows clinicians to make better decisions [35,52]. Importantly, WB-DWI has also been found to be useful in differentiating the stages of monoclonal gammopathies [35,52]. In addition, WB-DWI provides more reliable differentiation between benign tumors and pathological vertebral compression fractures [49].
collapse from malignant fractures with a reported sensitivity and specificity of 95.6% and 90%, respectively [36,48].

Figure 2. WB-MRI in a 75-year-old woman with mixed active and treated disease. (A) Sagittal T1-WI shows a lesion (black arrow) in the posterior and inferior L2 vertebral body. (B) Sagittal STIR image shows the L2 lesion (black arrow), as well as at several other sites of disease (thin white arrows) that were not obvious on the T1-WI. (C) Coronal T1 Dixon Fat image shows the L2 lesion (black arrow) and another lesion in the right ischium (white arrow). (D) Fused b800-Dixon image shows increased signal at L2 (black arrow) and to a lesser extent at the ischial lesion (white arrow).

DWI-MRI is also useful in evaluating treatment efficacy, remission, and prognostication. ADC images, similar to other functional imaging modalities, account for the heterogeneous pathologic distribution and patchy marrow infiltration in MM (which is accentuated after relapse) [53], and thus can help to monitor the treatment response in MM patients.

With effective treatment, responders show an increased ADC value at 4–6 weeks post-treatment: hemorrhage, edema, vascular congestion, and liquefaction necrosis from tumor cell death contribute to an increased diffusivity [35,46]. ADC then decreases at 20 weeks after therapy, due to a reappearance of normal fatty marrow. Conversely, patients who are resistant to therapy show a persistent marrow hyperintensity on DWI, and hypointensity on ADC images [35,37,54,55].

DWI images have also been proven to detect prognostically relevant residual focal lesions with a higher sensitivity than PET and PET/CT [56,57]. For a diffuse pattern of myeloma infiltration, there are controversies regarding the correlation between the ADC signal and therapy response, as some studies support its use [47] while others do not [47,58]. A previous study has related this issue to the slower transition from replaced to recovered fatty marrow in focal MM lesions that make it possible to be captured when imaged post-treatment [36]. WB-MRI with DWI also offers a quantitative analysis of the entire bone marrow after treatment, which is invaluable for clinicians directing further need for therapies and remission.
The limitations of WB-DWI include some issues with a definitive identification of lesions, and with resolution. The ADC value of MM lesions increases early after effective treatment, and for an interpreter who is blind to previous exams, this might resemble a hemangioma. However, when previous images are not available, corresponding T1 and T2-weighted imaging solves this issue. Additionally, DWI suffers from some limitations in resolution, rendering, and field of view (FOV). Therefore, when interpreted with the available anatomical data, DWI is an important support for MM, and should be added to morphologic imaging sequences in WB protocols; and it currently cannot be relied on as a stand-alone imaging modality [49].

2.3. IVIM MRI

IVIM theory was first introduced by Le Bihan et al. more than two decades ago, describing the role of the perfusion effects on the significant signal decay at b-values of less than 300 s/mm² [59]. The recorded ADC value is influenced by a combination of tissue microarchitecture/cellularity and perfusion. Given that the routine clinical implementation of DWI does not include a complete set of b-values (low, intermediate, and high), it does not provide information on the perfusion component of the signal derived from the imaged tissues [59].

As previously mentioned, in MM, the impact of the perfusion component on the ADC value seems to be larger than the effect of hypercellularity; hence, IVIM may be even more promising than DWI for MM patients [59]. IVIM decontaminates the ADC maps from the so-called “pseudo-diffusion” (D*), and marks out a part of the ADC signal that arises from any process other than diffusion, which is predominantly microperfusion at the capillary level. By taking three or more DWI images that are set at low, intermediate, and high b-values, two or more ADC maps could be generated, where ADC maps of low to moderate b-values reflect a combination of perfusion (D*: pseudo or enhanced diffusivity) and true diffusion (D). Subtracting these ADC maps will achieve tissue vascularity, which would be of added value to predict prognosis and assess treatment response in MM.

IVIM parameter D, the molecular diffusion coefficient, has been reported to be significantly higher in a diffuse pattern of myeloma marrow infiltration [60]. IVIM parameters are also associated with serum levels of biomarkers, showing that IVIM could be another useful tool for the prognostication of disease activity in MM [61].

2.4. Marrow Fat Quantification Techniques

Recently, gradient echo-based Dixon MRI as a fat quantification method approach has been evolved to include the generation of four separate image types: in-phase (IP), out-of-phase (OP), water-only (WO), and fat-only (FO) [62–64]. This method has been used for anatomical WB-MRI in MM, and it has several benefits over conventional T1- or T2-weighted imaging. The fat fraction is obtained using OP and IP images, and it can be efficient for both the lesion detection and the response assessment of bone lesions in MM patients [62]. It also has been shown that T2 Dixon fat-only Dixon images are more efficient by providing higher lesion detection rates compared to in-phase images alone in multiple myeloma [65].

2.5. Whole-Body Dynamic Contrast-Enhanced MRI (WB-DCE MRI)

WB-DCE MRI refers to the acquisition of serial images pre- and post-contrast administration, which can provide functional data on marrow infiltration in MM. WB-DCE MRI data can be computed into time-signal intensity curves providing a quantitative assessment of myeloma marrow infiltration, which could be valuable for diagnosis, prognosis, and monitoring treatment response. With regard to diagnosis, higher peaks and steeper slopes of WB-DCE MRI curves are associated with a higher percentage of plasma cell infiltration [66–68]. Prior studies have also found WB-DCE MRI curves to be well-correlated with MM disease activity and serum biomarkers [67]. In terms of prognosis, the relative signal enhancement of marrow in SMM predicts its rapid progression into symptomatic disease,
and more severe enhancement in progressive MM predicting a shorter progression-free survival [69,70]. After treatment, persistent abnormally elevated peaks of marrow enhancement and the foci of early enhancement portend disease progression or relapse (poor treatment response) [71]. Due to the paucity of research on standard acquisition protocols and reporting systems, WB-DCE MRI is not presently recommended for daily practice [72].

3. 18F-Fluorodeoxyglucose Positron Emission Tomography (18FDG PET)

3.1. 18F PET

18F-fluorodeoxyglucose (FDG) is the most commonly used radiotracer in PET imaging for MM [73]. FDG uptake, which is widely semi-quantitatively assessed via the maximum standardized uptake value (SUVmax), is known to be correlated with biological tumor aggressiveness [74]. 18FDG PET may over- or underestimate MM disease activity; for example, over-calling pathologic fractures or under-identifying small sized lesions. Moreover, SUVmax measures 18FDG uptake in a single-voxel region of interest, which might emit a distorted signal due to noise, reconstruction protocols, and post-processing (if applicable). Additionally, a single-voxel measurement may not be representative of marrow disease in its entirety [75]. Reporting SUVmean and SUVpeak instead of SUVmax addresses these setbacks, but is rarely performed in clinical practice [76]. Additionally, recent studies have suggested that measuring total lesion glycolysis (TLG) and metabolic tumor volume (MTV) may represent the tumor burden/activity and the patient’s prognosis more accurately [77,78]. However, these measurements can be time-consuming and not routinely performed in clinical practice. Hybrid imaging and adding the morphological data from CT or MRI into the functional data from PET imaging resolves these issues to a considerable extent.

3.2. 18FDG PET/CT

18FDG PET/CT is a sensitive diagnostic modality for both medullary and extramedullary plasma cell dyscrasias, and can accurately detect MM lesions, assess treatment response, and predict prognosis and progression in MM [77,79–81]. The wider employment of 18FDG PET/CT has also resulted in a significantly increased detection rate for extramedullary MM at diagnosis compared to older imaging techniques [82].

The International Myeloma Working Group (IMWG) recently introduced the evaluation of minimal residual disease (MRD) within the multiple myeloma (MM) response criteria. Currently the most powerful predictor of favorable outcomes over long-term follow-up, MRD negativity can be assessed both inside and outside of the bone marrow. Functional imaging techniques such as PET/CT and magnetic resonance imaging (MRI) serve in sensitive response assessment, and have been shown to be promising in terms of evaluating the response to treatment. Not only have they helped to assess MRD status in MM patients, but they also provide a global representation of the tumor burden by including several prognostic markers in addition to lesion type [83].

The sensitivity of 18FDG PET/CT for focal MM lesions has been reported to be more than WB-MRI, roughly estimated at 75% [84]; however, a diffuse pattern of marrow infiltration is better detected using WB-MRI. Additionally, 18FDG PET/CT was shown to have more promising results in having a higher impact on clinical decisions than WB-MRI in MM patients in terms of prognosis and management [85]. In a recent meta-analysis, a high 18FDG uptake significantly and independently predicted a shorter overall and progression-free survival in MM patients [81] when considering whole-body burden disease [81].

WB-MRI with DWI provides a non-invasive and quantitative assessment of the entire bone marrow after treatment. Based on a study by Torkian et al., DWI had a pooled sensitivity of 78% (95% CI: 72–83) and a specificity of 73% (95% CI: 61–83) in distinguishing responders from non-responders, emphasizing the prominent role of DWI for treatment response assessment in patients with MM [86]. In a cohort of 49 cases, WB DWI has been shown to be more sensitive than 18FDG PET/CT for detecting intramedullary lesions in all regions except the skull, both in patients with a new diagnosis and previously treated
patients. Additionally, WB DWI has been shown to have a sensitivity equivalent to that of 18F-DG PET/CT for detecting extramedullary lesions [87].

3.3. 18FDG PET/MRI

Finally, hybrid 18FDG PET/MRI techniques combine the morphological information provided by MRI and the metabolic and functional data furnished via 18FDG PET imaging, which allows for both the ability to detect the marrow foci of myeloma infiltration and to assess prognosis and treatment response. 18FDG PET/MRI increases the visibility of focal MM lesions at diagnosis and initial staging, and localizes residual disease activity after treatment [12]. 18FDG PET/MRI has been reported to have a higher lesion detectability rate than 18FDG PET/CT in evaluating skeletal lesions [88]. However, Sachpekidis et al. concluded that these two techniques are equally sensitive in detecting MM lesions [89]. As with WB-DCE MRI, there are few studies on a standard acquisition protocol and reporting system in 18FDG PET/MRI for MM; thus, further studies are warranted to test repeatability and validity before 18FDG PET/MRI can be considered as a viable tool in a routine imaging work-up of MM.

4. Conclusions

In the last two decades, novel imaging techniques have been developed or employed for MM to provide more information that can guide not only diagnosis and staging, but also treatment efficacy. These imaging techniques, due to their low invasiveness and high reliability, have gained significant clinical attention and have changed the clinical practice. The development of functional MRI sequences such as diffusion weighted imaging (DWI) and Intravoxel Incoherent Motion (IVIM) has made the functional assessment of lesions feasible.

DWI-MRI provides a high utility tool for not only the diagnosis and initial staging of plasma cell dyscrasias such as MM, but also for the evaluation of treatment efficacy and for re-staging. It can reliably detect tissue hypercellularity with high sensitivity, challenging the use of other procedures or imaging modalities for the management of MM. Therefore, DWI-MRI presents a promising option for clinicians engaged in the management of plasma cell dyscrasias like MM.

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References

1. Dimopoulos, M.; Terpos, E.; Comenzo, R.L.; Tosi, P.; Bekscak, M.; Sezer, O.; Siegel, D.; Lokhorst, H.; Kumar, S.; Rajkumar, S.V.; et al. International myeloma working group consensus statement and guidelines regarding the current role of imaging techniques in the diagnosis and monitoring of multiple Myeloma. Leukemia 2009, 23, 1545–1556. [CrossRef] [PubMed]
2. Bataille, R.; Harousseau, J.-L. Multiple Myeloma. N. Engl. J. Med. 1997, 336, 1657–1664. [CrossRef] [PubMed]
3. Kyle, R.A.; Durie, B.G.M.; Rajkumar, S.V.; Landgren, O.; Blade, J.; Merlini, G.; Kröger, N.; Einsele, H.; Vesole, D.H.; Dimopoulos, M.; et al. Monoclonal gammopathy of undetermined significance (MGUS) and smoldering (asymptomatic) multiple myeloma: IMWG consensus perspectives risk factors for progression and guidelines for monitoring and management. Leukemia 2010, 24, 1121–1127. [CrossRef] [PubMed]
4. Smith, D.; Yong, K. Multiple myeloma. BMJ 2013, 346, f3863. [CrossRef] [PubMed]
Diagnostics 2022, 12, 2182

5. Rajkumar, S.V.; Dimopoulos, M.A.; Palumbo, A.; Blade, J.; Merlini, G.; Mateos, M.V.; Kumar, S.; Hillengass, J.; Kastritis, E.; Richardson, P.; et al. International Myeloma Working Group updated criteria for the diagnosis of multiple myeloma. Lancet Oncol. 2014, 15, e538–e548. [CrossRef]

6. Durie, B.G.; Harousseau, J.L.; Miguel, J.S.; Bladé, J.; Barlogie, B.; Anderson, K.; Gertz, M.; Dimopoulos, M.; Westin, J.; Sonneveld, P.; et al. International uniform response criteria for multiple myeloma. Leukemia 2006, 20, 1467–1473. [CrossRef] [PubMed]

7. Ormond Filho, A.G.; Carneiro, B.C.; Pastore, D.; Silva, I.P.; Yamashita, S.R.; Consolo, F.D.; Hungria, V.T.M.; Sandes, A.F.; Rizzatti, E.G.; Nico, M.A.C. Whole-Body Imaging of Multiple Myeloma: Diagnostic Criteria. Radiographics 2019, 39, 1077–1097. [CrossRef] [PubMed]

8. Hotta, T. Classification, staging and prognostic indices for multiple myeloma. Nihon Rinsho Jpn. J. Clin. Med. 2007, 65, 2161–2166.

9. Durie, B.G. The role of anatomic and functional staging in myeloma: Description of Durie/Salmon plus staging system. Eur. J. Cancer 2006, 42, 1539–1543. [CrossRef]

10. Hillengass, J.; Fechtner, K.; Weber, M.A.; Bäuerle, T.; Ayyaz, S.; Heiss, C.; Hielscher, T.; Egerer, G.; Neben, K.; et al. Prognostic significance of focal lesions in whole-body magnetic resonance imaging in patients with asymptomatic multiple myeloma. J. Clin. Oncol. Off. J. Am. Soc. Clin. Oncol. 2010, 28, 1606–1610. [CrossRef]

11. Walker, R.; Barlogie, B.; Haessler, J.; Tricot, G.; Anaissie, E.; Shaughnessy, J.D., Jr.; Epstein, J.; van Hemert, R.; Erdem, E.; Hoering, A.; et al. Magnetic resonance imaging in multiple myeloma: Clinical and diagnostic implications. J. Clin. Oncol. Off. J. Am. Soc. Clin. Oncol. 2007, 25, 1121–1128. [CrossRef][PubMed]

12. Dimopoulos, M.A.; Hillengass, J.; Usmani, S.; Zamagni, E.; Lentzsch, S.; Davies, F.E.; Raje, N.; Sezer, O.; Zweegman, S.; Shah, J.; et al. Role of magnetic resonance imaging in the management of patients with multiple myeloma: A consensus statement. J. Clin. Oncol. Off. J. Am. Soc. Clin. Oncol. 2015, 33, 657–664. [CrossRef] [PubMed]

13. Pawlyn, C.; Fowkes, L.; Otero, S.; Jones, J.R.; Boyd, K.D.; Davies, F.E.; Morgan, G.J.; Collins, D.J.; Sharma, B.; Riddell, A.; et al. Whole-body diffusion-weighted MRI: A new gold standard for assessing disease burden in patients with multiple myeloma? Leukemia 2016, 30, 1446–1448. [CrossRef]

14. Bäuerle, T.; Hillengass, J.; Fechtner, K.; Zechmann, C.M.; Grenacher, L.; Moehler, T.M.; Christiane, H.; Wagner-Gund, B.; Neben, K.; Kauczor, H.U.; et al. Multiple myeloma and monoclonal gammopathy of undetermined significance: Importance of whole-body versus spinal MR imaging. Radiology 2009, 252, 477–483. [CrossRef] [PubMed]

15. Baur, A.; Stäbler, A.; Nagel, D.; Lamerz, R.; Bartl, R.; Hiller, E.; Wendtner, C.; Bachner, F.; Reiser, M. Magnetic resonance imaging as a supplement for the clinical staging system of Durie and Salmon? Cancer 2002, 95, 1334–1345. [CrossRef]

16. Kusumoto, S.; Jinnai, I.; Itòh, K.; Kawai, N.; Sakata, T.; Matsuda, A.; Tominaga, K.; Murohashi, I.; Bessho, M.; Harashima, K.; et al. Magnetic resonance imaging patterns in patients with multiple myeloma. Br. J. Haematol. 1997, 99, 649–655. [CrossRef] [PubMed]

17. Moulopoulos, L.A.; Gika, D.; Anagnostopoulo, A.; Delasalle, K.; Weber, D.; Alexanian, R.; Dimopoulos, M.A. Prognostic significance of magnetic resonance imaging of bone marrow in previously untreated patients with multiple myeloma. Ann. Oncol. Off. J. Eur. Soc. Med. Oncol. 2005, 16, 1824–1828.

18. Bäuerle, T.; Hillengass, J.; Fechtner, K.; Zechmann, C.M.; Grenacher, L.; Moehler, T.M.; Christiane, H.; Wagner-Gund, B.; Neben, K.; Kauczor, H.U.; et al. Whole-body and monoclonal gammopathy of undetermined significance: Importance of whole-body versus spinal MR imaging. Radiology 2009, 252, 477–483. [CrossRef] [PubMed]

19. Baur, A.; Stäbler, A.; Nagel, D.; Lamerz, R.; Bartl, R.; Hiller, E.; Wendtner, C.; Bachner, F.; Reiser, M. Magnetic resonance imaging as a supplement for the clinical staging system of Durie and Salmon? Cancer 2002, 95, 1334–1345. [CrossRef]

20. Kusumoto, S.; Jinnai, I.; Itòh, K.; Kawai, N.; Sakata, T.; Matsuda, A.; Tominaga, K.; Murohashi, I.; Bessho, M.; Harashima, K.; et al. Magnetic resonance imaging patterns in patients with multiple myeloma. Br. J. Haematol. 1997, 99, 649–655. [CrossRef] [PubMed]

21. Moulopoulos, L.A.; Gika, D.; Anagnostopoulo, A.; Delasalle, K.; Weber, D.; Alexanian, R.; Dimopoulos, M.A. Prognostic significance of magnetic resonance imaging of bone marrow in previously untreated patients with multiple myeloma. Ann. Oncol. Off. J. Eur. Soc. Med. Oncol. 2005, 16, 1824–1828.

22. Weininger, M.; Lauterbach, B.; Knop, S.; Pabst, T.; Penn, W.; Hahn, D.; Beissert, M. Whole-body MRI of multiple myeloma: Comparison of different MRI sequences in assessment of different growth patterns. Eur. J. Radiol. 2009, 69, 339–345. [CrossRef] [PubMed]

23. Albano, D.; Stecco, A.; Micci, G.; Sconfienza, L.M.; Colagrande, S.; Reginelli, A.; Grassi, R.; Carriero, A.; Midiri, M.; Lagalla, R.; et al. Whole-body magnetic resonance imaging (WB-MRI) in oncology: An Italian survey. Radiographics 2019, 39, 1077–1097. [CrossRef] [PubMed]

24. Feldhaus, J.M.; Garner, H.W.; Wessell, D.E. Society of skeletal radiology member utilization and performance of whole-body MRI in adults. Skelet. Radiol. 2020, 49, 1731–1736. [CrossRef]

25. Dinter, D.J.; Neff, W.K.; Klaus, J.; Böhm, C.; Haslka, J.; Weiss, C.; Schoenberg, S.O.; Metzgeroth, G. Comparison of whole-body MR imaging and conventional X-ray examination in patients with multiple myeloma and implications for therapy. Ann. Hematol. 2009, 88, 457–464. [CrossRef] [PubMed]
27. Ghanem, N.; Lohrmann, C.; Engelhardt, M.; Pache, G.; Uhl, M.; Saueressig, U.; Kotter, E.; Langer, M. Whole-body MRI in the detection of bone marrow infiltration in patients with plasma cell neoplasms in comparison to the radiological skeletal survey. *Eur. Radiol.* 2006, 16, 1005–1014. [CrossRef]

28. Dutoit, J.C.; Vanderkerken, M.A.; Anthonissen, J.; Dochy, F.; Verstraete, K.L. The diagnostic value of SE MRI and DWI of the spine in patients with monoclonal gammopathy of undetermined significance, smouldering myeloma and multiple myeloma. *Eur. Radiol.* 2014, 24, 2754–2765. [CrossRef]

29. Jacobs, M.A.; Pan, L.; Macura, K.J. Whole-body diffusion-weighted and proton imaging: A review of this emerging technology for monitoring metastatic cancer. *Semin. Roentgenol.* 2009, 44, 111–122. [CrossRef]

30. Hillengass, J.; Bäuerle, T.; Bartl, R.; Andruis, M.; McClanahan, F.; Laun, F.B.; Zechmann, C.M.; Shah, R.; Wagner-Gund, B.; Simon, D.; et al. Diffusion-weighted imaging for non-invasive and quantitative monitoring of bone marrow infiltration in patients with monoclonal plasma cell disease: A comparative study with histology. *Br. J. Haematol.* 2011, 153, 721–728. [CrossRef]

31. Horger, M.; Weisel, K.; Horger, W.; Mroue, A.; Fenchel, M.; Lichy, M. Whole-body diffusion-weighted MRI with apparent diffusion coefficient mapping for early response monitoring in multiple myeloma: Preliminary results. *AJR Am. J. Roentgenol.* 2011, 196, 6. [CrossRef]

32. Messiou, C.; Collins, D.J.; Morgan, V.A.; Desouza, N.M. Optimising diffusion weighted MRI for imaging metastatic and myeloma bone disease and assessing reproducibility. *Eur. Radiol.* 2011, 21, 1713–1718. [CrossRef] [PubMed]

33. Sommer, G.; Klarhöfer, M.; Lenz, C.; Scheffler, K.; Bongartz, G.; Winter, L. Signal characteristics of focal bone marrow lesions in patients with multiple myeloma using whole body T1w-TSE, T2w-STIR and diffusion-weighted imaging with background suppression. *Eur. Radiol.* 2011, 21, 857–862. [CrossRef]

34. Fenchel, M.; Konakchieva, M.; Weisel, K.; Kraus, S.; Claussen, C.D.; Horger, M. Response assessment in patients with multiple myeloma during antiangiogenic therapy using arterial spin labeling and diffusion-weighted imaging: A feasibility study. *Acad. Radiol.* 2010, 17, 1326–1333. [CrossRef]

35. Messiou, C.; Giles, S.; Collins, D.J.; West, S.; Davies, F.E.; Morgan, G.J.; Desouza, N.M. Assessing response of myeloma bone disease with diffusion-weighted MRI. *Br. J. Radiol.* 2012, 85, 1020. [CrossRef]

36. Koutoulidis, V.; Fontara, S.; Terpos, E.; Zagouri, F.; Matsaridis, D.; Christoulas, D.; Panourgias, E.; Kastritis, E.; Dimopoulos, M.A.; Moulopoulos, L.A. Quantitative Diffusion-weighted Imaging of the Bone Marrow: An Adjunct Tool for the Diagnosis of a Diffuse MR Imaging Pattern in Patients with Multiple Myeloma. *Radiology* 2017, 282, 484–493. [CrossRef]

37. Giles, S.L.; Messiou, C.; Collins, D.J.; Simpkin, C.J.; West, S.; Davies, F.E.; Morgan, G.J.; deSouza, N.M. Whole-Body Diffusion-weighted MR Imaging for Assessment of Treatment Response in Myeloma. *Radiology* 2014, 271, 785–794. [CrossRef]

38. Lemke, A.; Laun, F.B.; Klaus, M.; Re, T.J.; Simon, D.; Delorme, S.; Schad, L.R.; Stieltjes, B. Differentiation of pancreas carcinoma from healthy pancreatic tissue using multiple b-values: Comparison of apparent diffusion coefficient and intravoxel incoherent motion derived parameters. *Investig. Radiol.* 2009, 44, 769–775. [CrossRef]

39. Luciani, A.; Vignaud, A.; Cavet, M.; Nihieu, J.T.; Mallat, A.; Ruel, L.; Laurent, A.; Deux, J.F.; Brugieres, P.; Rahmouni, A. Liver cirrhosis: Intravoxel incoherent motion MR imaging—Pilot study. *Radiology* 2008, 249, 891–899. [CrossRef]

40. Sezer, O.; Niemöller, K.; Jakob, C.; Zavrski, I.; Heider, U.; Eucker, J.; Kaufmann, O.; Possinger, K. Relationship between bone marrow angiogenesis and plasma cell infiltration and serum beta2-microglobulin levels in patients with multiple myeloma. *Ann. Hematol.* 2001, 80, 598–601. [CrossRef]

41. Ajit, M.; Maruvaneni, S.; Kumar, A.; Ullal, S.; Fernandes, M. Role of Diffusion Weighted Imaging in Differentiating Benign from Pathological Vertebral Collapse using ADC Values. *J. Clin. Diagn. Res.* 2019, 13, 1–5.

42. Biffar, A.; Baur-Melnyk, A.; Schmidt, G.P.; Reiser, M.F.; Dietrich, O. Multiparameter MRI assessment of normal-appearing and diseased vertebral bone marrow. *Eur. Radiol.* 2010, 20, 2679–2689. [CrossRef]

43. Plank, C.; Koller, A.; Mueller-Mang, C.; Bammer, R.; Thurnher, M.M. Diffusion-weighted MR imaging (DWI) in the evaluation of epidural spinal lesions. *Neuroradiology* 2007, 49, 977–985. [CrossRef] [PubMed]

44. Pearce, T.; Philip, S.; Brown, J.; Koh, D.M.; Burn, P.R. Bone metastases from prostate, breast and multiple myeloma: Differences in lesion conspicuity at short-tau inversion recovery and diffusion-weighted MRI. *Br. J. Radiol.* 2012, 85, 1102–1106. [CrossRef] [PubMed]

45. Padhani, A.R.; Koh, D.M.; Collins, D.J. Whole-body diffusion-weighted MR imaging in cancer: Current status and research directions. *Radiology* 2011, 261, 701–718. [CrossRef]

46. Lacognata, C.; Crimi, F.; Guolo, A.; Varin, C.; De March, E.; Vio, S.; Ponzoni, A.; Barilì, G.; Lico, A.; Branca, A.; et al. Diffusion-weighted whole-body MRI for evaluation of early response in multiple myeloma. *Clin. Radiol.* 2017, 72, 850–857. [CrossRef] [PubMed]

47. Giles, S.L.; deSouza, N.M.; Collins, D.J.; Morgan, V.A.; West, S.; Davies, F.E.; Morgan, G.J.; Messiou, C. Assessing myeloma bone disease with whole-body diffusion-weighted imaging: Comparison with x-ray skeletal survey by region and relationship with laboratory estimates of disease burden. *Clin. Radiol.* 2015, 70, 614–621. [CrossRef]
Diagnostics 2022, 12, 2182

49. Sachpekidis, C.; Mosebach, J.; Freitag, M.T.; Wilhelm, T.; Mai, E.K.; Goldschmidt, H.; Haberkorn, U.; Schlemmer, H.-P.; Delorme, S.; Dimitrakopoulou-Strauss, A. Application of (18)FDG PET and diffusion weighted imaging (DWI) in multiple myeloma: Comparison of functional imaging modalities. Am. J. Nucl. Med. Mol. Imaging 2015, 5, 479–492.

50. Rasche, L.; Angtuaco, E.; McDonald, J.E.; Buros, A.; Stein, C.; Pawlyn, C.; Thanendrarajan, S.; Schinke, C.; Samant, R.; Yaccoby, S.; et al. Low expression of hexokinase-2 is associated with false-negative FDG-potison emission tomography in multiple myeloma. Blood 2017, 130, 30–34. [CrossRef]

51. Lai, A.Y.T.; Riddell, A.; Barwick, T.; Boyd, K.; Rockall, A.; Kaiser, M.; Koh, D.M.; Saffar, H.; Yusuf, S.; Messiou, C. Interobserver agreement of whole-body magnetic resonance imaging is superior to whole-body computed tomography for assessing disease burden in patients with multiple myeloma. Eur. Radiol. 2020, 30, 320–327. [CrossRef] [PubMed]

52. Hernandez, J.; Montesinos, O.; Mateo, A.G.; Queizán, J.A.; Peral, G.S.d.; Olivier, C.; Fisac, R.M.; Coca, A.G.d.; López, R.; Ocio, E.M.; et al. Usefulness of Whole-Body Diffusion-Weighted MRI (WB-MRI) With Apparent Diffusion Coefficient (ADC) In The Differentiation of Monoclonal Gammapathies. Clin. Lymphoma Myeloma Leuk. 2015, 15, e90–e91. [CrossRef]

53. Rasche, L.; Schinke, C.D.; Alapat, D.; Gershner, G.; Johnson, S.K.; Thanendrarajan, S.; Epstein, J.; van Rhee, F.; Zangari, M.; McDonald, J.E.; et al. Functional Imaging Detects Residual Disease in MRD-Negative Multiple Myeloma Patients Who Subsequently Relapse. Blood 2017, 130, 4510.

54. Lecouvet, F.E.; Larbi, A.; Pasogliou, V.; Omoumi, P.; Tombal, B.; Michoux, N.; Malghem, J.; Lhomel, R.; Vande Berg, B.C. MRI for the evaluation of bone lesions in patients with multiple myeloma. Radiology 2013, 266, 710–717. [CrossRef]

55. Khoo, M.M.; Tyler, P.A.; Saifuddin, A.; Padhani, A.R. Diffusion-weighted imaging (DWI) in musculoskeletal MRI: A critical review. Skelet. Radiol. 2011, 40, 665–681. [CrossRef] [PubMed]

56. Koutoulidis, V.; Terpos, E.; Papanikolaou, N.; Fontara, S.; Seimenis, I.; Gavriatopoulou, M.; Ntanasis-Stathopoulos, I.; Bourgioti, C.; Dimitrakopoulou-Strauss, A. Application of (18)FDG PET and diffusion weighted imaging (DWI) in multiple myeloma: Comparison of functional imaging modalities. Am. J. Nucl. Med. Mol. Imaging 2015, 5, 479–492.

57. Le Bihan, D.; Breton, E.; Lallemand, D.; Aubin, M.L.; Vignaud, J.; Laval-Jeantet, M. Separation of diffusion and perfusion in magnetic resonance imaging. J. Magn. Reson. Imaging JMRI 1988, 1, 211–220. [CrossRef]

58. Koutoulidis, V.; Terpos, E.; Papanikolaou, N.; Fontara, S.; Seimenis, I.; Gavriatopoulou, M.; Ntanasis-Stathopoulos, I.; Bourgioti, C.; Santinini, J.; Moreira, J.M.; et al. Comparison of MRI Features of Fat Fraction and ADC for Early Treatment Response Assessment in Patients with Multiple Myeloma. Radiology 2022, 304, 211388. [CrossRef] [PubMed]

59. Le Bihan, D.; Breton, E.; Lallemand, D.; Aubin, M.L.; Vignaud, J.; Laval-Jeantet, M. Separation of diffusion and perfusion in magnetic resonance imaging. J. Magn. Reson. Imaging JMRI 1988, 1, 211–220. [CrossRef]

60. Bourillon, C.; Rahmouni, A.; Lin, C.; Belhadj, K.; Beaussart, P.; Vignaud, A.; Zerib, P.; Pigneur, F.; Cuend, C.A.; Bessalem, H.; et al. Intravoxel Incoherent Motion Diffusion-weighted Imaging of Multiple Myeloma Lesions: Correlation with whole-body dynamic contrast agent-enhanced MRI. Radiology 2015, 277, 773–783. [CrossRef]

61. Shah, R.; Stieltjes, B.; Andrusils, M.; Pfeiffer, R.; Sumkauskaite, M.; Delorme, S.; Schlemmer, H.P.; Goldschmidt, H.; Landgren, O.; Hillengass, J. Intravoxel incoherent motion imaging for assessment of bone marrow infiltration of monoclonal plasma cell diseases. Ann. Hematol. 2013, 92, 1553–1557. [CrossRef]

62. Bray, T.J.P.; Singh, S.; Latifoltojar, A.; Rajesperan, K.; Rahman, F.; Narayanan, P.; Naaseri, S.; Lopes, A.; Bainbridge, A.; Punwani, S.; et al. Diagnostic utility of whole body Dixon MRI in multiple myeloma: A multi-reader study. PLoS ONE 2017, 12, e0180562. [CrossRef] [PubMed]

63. Koutoulidis, V.; Terpos, E.; Papanikolaou, N.; Fontara, S.; Seimenis, I.; Gavriatopoulou, M.; Ntanasis-Stathopoulos, I.; Bourgioti, C.; Santinini, J.; Moreira, J.M.; et al. Comparison of MRI Features of Fat Fraction and ADC for Early Treatment Response Assessment in Patients with Multiple Myeloma. Radiology 2022, 304, 211388. [CrossRef] [PubMed]

64. Bourillon, C.; Rahmouni, A.; Lin, C.; Belhadj, K.; Beaussart, P.; Vignaud, A.; Zerib, P.; Pigneur, F.; Cuend, C.A.; Bessalem, H.; et al. Intravoxel Incoherent Motion Diffusion-weighted Imaging of Multiple Myeloma Lesions: Correlation with whole-body dynamic contrast agent-enhanced MRI. Radiology 2015, 277, 773–783. [CrossRef]

65. Shah, R.; Stieltjes, B.; Andrusils, M.; Pfeiffer, R.; Sumkauskaite, M.; Delorme, S.; Schlemmer, H.P.; Goldschmidt, H.; Landgren, O.; Hillengass, J. Intravoxel incoherent motion imaging for assessment of bone marrow infiltration of monoclonal plasma cell diseases. Ann. Hematol. 2013, 92, 1553–1557. [CrossRef]

66. Bray, T.J.P.; Singh, S.; Latifoltojar, A.; Rajesperan, K.; Rahman, F.; Narayanan, P.; Naaseri, S.; Lopes, A.; Bainbridge, A.; Punwani, S.; et al. Diagnostic utility of whole body Dixon MRI in multiple myeloma: A multi-reader study. PLoS ONE 2017, 12, e0180562. [CrossRef] [PubMed]

67. Koutoulidis, V.; Terpos, E.; Papanikolaou, N.; Fontara, S.; Seimenis, I.; Gavriatopoulou, M.; Ntanasis-Stathopoulos, I.; Bourgioti, C.; Santinini, J.; Moreira, J.M.; et al. Comparison of MRI Features of Fat Fraction and ADC for Early Treatment Response Assessment in Patients with Multiple Myeloma. Radiology 2022, 304, 211388. [CrossRef] [PubMed]

68. Berardo, S.; Sukhovei, L.; Andorno, S.; Carriero, A.; Stecco, A. Quantitative bone marrow magnetic resonance imaging through apparent diffusion coefficient and fat fraction in multiple myeloma patients. Le Radiol. Med. 2021, 126, 445–452. [CrossRef]

69. Danner, A.; Brumpt, E.; Alliet, M.; Tio, G.; Omoumi, P.; Aubry, S. Improved contrast for myeloma focal lesions with T2-weighted Dixon images compared to T1-weighted images. Diagn. Interv. Imaging DIPII 2019, 100, 513–519. [CrossRef]

70. Lin, C.; Luciani, A.; Belhadj, K.; Maison, P.; Vignaud, A.; Deux, J.F.; Zerib, P.; Pigneur, F.; Itti, E.; Koberite, H.; et al. Patients with plasma cell disorders examined at whole-body dynamic contrast-enhanced MR imaging: Initial experience. Radiology 2009, 250, 905–915. [CrossRef] [PubMed]

71. Nosás-Garcia, S.; Moehler, T.; Wasser, K.; Kiessling, F.; Bartl, R.; Zuna, I.; Hillengass, J.; Goldschmidt, H.; Kauczor, H.U.; Delorme, S. Dynamic contrast-enhanced MRI for assessing the disease activity of multiple myeloma: A comparative study with histology and clinical markers. J. Magn. Reson. Imaging JMRI 2005, 22, 154–162. [CrossRef]

72. Rahmouni, A.; Montazel, J.L.; Divine, M.; Lepage, E.; Belhadj, K.; Gaultard, P.; Bouanane, M.; Golli, M.; Koberite, H. Bone marrow with diffuse tumor infiltration in patients with lymphoproliferative diseases: Dynamic gadolinium-enhanced MR imaging. Radiology 2003, 229, 710–717. [CrossRef]

73. Hillengass, J.; Ritsch, J.; Merz, M.; Wagner, B.; Kunz, C.; Hielscher, T.; Laue, H.; Bäuerle, T.; Zechmann, C.M.; Ho, A.D.; et al. Increased microcirculation detected by dynamic contrast-enhanced magnetic resonance imaging is of prognostic significance in asymptomatic myeloma. Br. J. Haematol. 2016, 174, 127–135. [CrossRef]
70. Hillengass, J.; Wasser, K.; Delorme, S.; Kiessling, F.; Zechmann, C.; Benner, A.; Kauczor, H.U.; Ho, A.D.; Goldschmidt, H.; Moehler, T.M. Lumbar bone marrow microcirculation measurements from dynamic contrast-enhanced magnetic resonance imaging is a predictor of event-free survival in progressive multiple myeloma. *Clin. Cancer Res. Off. J. Am. Assoc. Cancer Res.* 2007, 13, 475–481. [CrossRef]

71. Lin, C.; Luciani, A.; Belhadj, K.; Deux, J.F.; Kuhnowski, F.; Maatouk, M.; Beaussart, P.; Cuenod, C.A.; Haioun, C.; Rahmouni, A. Multiple myeloma treatment assessment with whole-body dynamic contrast-enhanced MR imaging. *Radiology* 2010, 254, 521–531. [CrossRef] [PubMed]

72. Mulé, S.; Reizine, E.; Blanc-Durand, P.; Baranes, L.; Zerbib, P.; Burns, R.; Nouri, R.; Itti, E.; Luciani, A. Whole-Body Functional MRI and PET/MRI in Multiple Myeloma. *Cancers* 2020, 12, 3155. [CrossRef]

73. Matteucci, F.; Paganelli, G.; Martinelli, G.; Cerchione, C. PET/CT in Multiple Myeloma: Beyond FDG. *Front. Oncol.* 2021, 10, 622501. [CrossRef]

74. Bartel, T.B.; Haessler, J.; Brown, T.L.; Shaughnessy, J.D., Jr.; van Rhee, F.; Anaissie, E.; Alpe, T.; Angtuaco, E.; Walker, R.; Epstein, J.; et al. F18-fluorodeoxyglucose positron emission tomography in the context of other imaging techniques and prognostic factors in multiple myeloma. *Blood* 2009, 114, 2068–2076. [CrossRef] [PubMed]

75. Soret, M.; Bacharach, S.L.; Buvat, I. Partial-volume effect in PET tumor imaging. Journal of nuclear medicine: Official publication. *Soc. Nucl. Med.* 2007, 48, 932–945. [CrossRef] [PubMed]

76. Wahl, R.L.; Jacene, H.; Kasamon, Y.; Lodge, M.A. From RECIST to PERCIST: Evolving Considerations for PET response criteria in solid tumors. *J. Nucl. Med. Off. Publ. Soc. Nucl. Med.* 2009, 50, 122s–150s. [CrossRef]

77. Fonti, R.; Larobina, M.; Del Vecchio, S.; De Luca, S.; Fabbricini, R.; Catalano, L.; Pane, F.; Salvatore, M.; Pace, L. Metabolic tumor volume assessed by 18F-FDG PET/CT for the prediction of outcome in patients with multiple myeloma. *J. Nucl. Med. Off. Publ. Soc. Nucl. Med.* 2012, 53, 1829–1835. [CrossRef]

78. McDonald, J.E.; Kessler, M.M.; Gardner, M.W.; Bueros, A.F.; Ntambi, J.A.; Waheed, S.; van Rhee, F.; Zangari, M.; Heuck, C.J.; Petty, N.; et al. Assessment of Total Lesion Glycolysis by (18)F FDG PET/CT Significantly Improves Prognostic Value of GEP and ISS in Myeloma. *Clin. Cancer Res. Off. J. Am. Assoc. Cancer Res.* 2017, 23, 1981–1987. [CrossRef]

79. Derlin, T.; Bannas, P. Imaging of multiple myeloma: Current concepts. *World J. Orthop.* 2014, 5, 272–282. [CrossRef]

80. Dimitrakopoulou-Strauss, A.; Hoffmann, M.; Bergner, R.; Uppenkamp, M.; Haberkorn, U.; Strauss, L.G. Prediction of progression-free survival in patients with multiple myeloma following anthracycline-based chemotherapy based on dynamic FDG-PET. *Clin. Nucl. Med.* 2009, 34, 576–584. [CrossRef]

81. Han, S.; Woo, S.; Kim, Y.I.; Yoon, D.H.; Ryu, J.S. Prognostic value of (18)F-fluorodeoxyglucose positron emission tomography/computed tomography in newly diagnosed multiple myeloma: A systematic review and meta-analysis. *Eur. Radiol.* 2021, 31, 152–162. [CrossRef]

82. Varettoni, M.; Corso, A.; Pica, G.; Mangiacavalli, S.; Pascutto, C.; Lazzarino, M. Incidence, presenting features and outcome of extramedullary disease in multiple myeloma: A longitudinal study on 1003 consecutive patients. *Ann. Oncol. Off. J. Eur. Soc. Med. Oncol.* 2010, 21, 325–330. [CrossRef] [PubMed]

83. Zamagni, E.; Tacchetti, P.; Barbato, S.; Cavo, M. Role of Imaging in the Evaluation of Minimal Residual Disease in Multiple Myeloma Patients. *J. Clin. Med.* 2020, 9, 3519. [CrossRef] [PubMed]

84. Hillengass, J.; Usmani, S.; Rajkumar, S.V.; Durie, B.G.M.; Mateos, M.V.; Lonial, S.; Joao, C.; Anderson, K.C.; García-Sanz, R.; Riva, E.; et al. International myeloma working group consensus recommendations on imaging in monoclonal plasma cell disorders. *Lancet Oncol.* 2019, 20, e302–e312. [CrossRef]

85. Lecouvet, F.E.; Boyadzhiev, D.; Collette, L.; Berckmans, M.; Michoux, N.; Triqueneaux, P.; Pasogliou, V.; Jambar, F.; Vekemans, M.C. MRI versus (18)F-FDG-PET/CT for detecting bone marrow involvement in multiple myeloma: Diagnostic performance and clinical relevance. *Eur. Radiol.* 2020, 30, 1927–1937. [CrossRef] [PubMed]

86. Torkian, P.; Mansoori, B.; Hillengass, J.; Azadbakht, J.; Rashedi, S.; Lee, S.S.; Amini, B.; Bonaffini, P.A.; Chalian, M. Diffusion-weighted imaging (DWI) in diagnosis, staging, and treatment response assessment of multiple myeloma: A systematic review and meta-analysis. *Skelet. Radiol.* 2022, 51, 622–635. [CrossRef] [PubMed]

87. Chen, J.; Li, C.; Tian, Y.; Xiao, Q.; Deng, M.; Hu, H.; Wen, B.; He, Y. Comparison of Whole-Body DWI and 18F-FDG PET/CT for Detecting Intramedullary and Extramedullary Lesions in Multiple Myeloma. *Am. J. Roentgenol.* 2019, 213, 514–523. [CrossRef]

88. Beiderwellen, K.; Huebner, M.; Heusch, P.; Grueneisen, J.; Ruhlmann, V.; Nersaa, F.; Kuehl, H.; Umultu, L.; Rosenbaum-Krumme, S.; Lauenstein, T.C. Whole-body 18F-FDG PET/MRI vs. PET/CT in the assessment of bone lesions in oncological patients: Initial results. *Eur. Radiol.* 2014, 24, 2023–2030. [CrossRef]

89. Sachpikidis, C.; Hillengass, J.; Goldschmidt, H.; Mosebach, J.; Pan, L.; Schlemmer, H.-P.; Haberkorn, U.; Dimitrakopoulou-Strauss, A. Comparison of (18)F-FDG PET/CT and PET/MRI in patients with multiple myeloma. *Am. J. Nucl. Med. Mol. Imaging* 2015, 5, 469–478.