Magnetic Resonance Evaluation of Multiple Myeloma at 3.0 Tesla: How Do Bone Marrow Plasma Cell Percentage and Selection of Protocols Affect Lesion Conspicuity?

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Abstract

Purpose: To compare various pulse sequences in terms of percent contrast and contrast-to-noise ratio (CNR) for detection of focal multiple myeloma lesions and to assess the dependence of lesion conspicuity on the bone marrow plasma cell percent (BMPC%).

Materials and Methods: Sagittal T1-weighted FSE, fat-suppressed T2-weighted FSE (FS-T2 FSE), fast STIR and iterative decomposition of water and fat with echo asymmetry and least-squares estimation (IDEAL) imaging of the lumbar spine were performed (n = 45). Bone marrow (BM)-focal myeloma lesion percent contrast and CNR were calculated. Spearman rank correlation coefficients were obtained between percent contrast, CNR and BMPC%. Percent contrasts and CNRs were compared among the three imaging sequences.

Results: BM-focal lesion percent contrasts, CNRs and BMPC% showed significant negative correlations in the three fat-suppression techniques. Percent contrast and CNRs were significantly higher for FS-T2 FSE than for STIR (P<0.01, P<0.05, respectively), but no significant differences were found among the three fat-suppression methods in the low tumor load BM group.

Conclusion: The higher BMPC% was within BM, the less conspicuous the focal lesion was on fat-suppressed MRI. The most effective protocol for detecting focal lesions was FS-T2 FSE. In the high tumor load BM group, no significant differences in lesion conspicuity were identified among the three fat-suppression techniques.

Introduction

Multiple myeloma is a plasma-cell malignancy characterized by the presence of lytic bone disease causing severe bone pain, pathological fractures, spinal cord compression and hypercalcemia [1]. Up to 90% of myeloma patients develop osteolytic lesions during the course of the disease [2]. These lesions occur predominantly in the axial skeleton (i.e., skull, spine, rib cage and pelvis), as well as the proximal areas of the arms and legs [3].

In 2003, the International Myeloma Working Group introduced the Durie-Salmon PLUS staging system [4], which takes into account the number of lesions detected by magnetic resonance imaging (MRI) or 18F-fluorodeoxyglucose positron emission tomography (PET). In patients with active myeloma, the number of lesions on MRI correlates very well with treatment outcomes and overall survival [5]. This excellent correlation with survival outcome is the primary reason for the inclusion of MRI into the Durie-Salmon PLUS system. However, counting focal lesions can be somewhat confusing, with variegated or diffuse patterns of tumor cell infiltration reportedly found in 57% of cases on T1-weighted imaging [6], obstructing detection of focal lesions. Furthermore, the Durie-Salmon PLUS staging system does not include the presence or absence of diffuse infiltration of tumor cells into the bone marrow (BM). This means that focal myeloma lesions must be detected regardless of any abnormality in background BM. The optimal MRI sequence for detecting focal bone lesions thus remains to be determined.

Various MR pulse sequences are available for evaluating the spine, including fast spin-echo (FSE) imaging with and without fat suppression, short inversion time inversion recovery (STIR) imaging, and the iterative decomposition of water and fat with echo asymmetry and least-squares estimation (IDEAL) technique. An important feature to detect focal lesions in the spine is suppression of marrow fat, because subtle high-intensity lesions

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can be obscured by the high signals from marrow fat on routine T1-weighted spin-echo imaging.

The STIR pulse sequence offers high sensitivity for detecting neoplasia due to its ability to show the combined effects of prolonged T1 and T2 relaxation times of these pathological tissues [7–9]. In 2000, Nakatsu et al. [9] reported STIR as superior to T1-weighted FSE with fat saturation for detection of metastatic lesions, in terms of lesion conspicuity.

A newer approach for achieving fat suppression is the simple spectroscopic imaging technique, from which IDEAL was later developed, originally published by Dixon [10]. In the original implementation, Dixon acquired an image with water and fat signals in-phase and another image with water and fat signals 180° out-of-phase. Dixon showed that simple summation and subtraction of the two images yield a water-only image and a fat-only image, respectively. The most serious problem for the Dixon techniques is B0 inhomogeneity, which appears as phase errors in the acquired Dixon images. Without proper phase correction, the simple summation and subtraction approach results in incomplete water and fat separation, thus making the Dixon techniques also sensitive to magnetic field inhomogeneity. After Dixon’s original work, Yeung and Kormos [11], Glover and Schneider [12], and Glover [13] showed that phase errors can actually be determined by acquiring an additional image. This analytical method for water and fat separation involves acquisition of three separate images with different water and fat relative phase angles and determination of water and fat on a per-pixel basis through an iterative least-squares process [14]. Integration of the region-growing and iterative linear least-squares methods has improved water-fat separation compared with using the original iterative process alone [15]. IDEAL is one of the three-point water-fat separation methods that uses asymmetric echoes and least-squares fitting to achieve the maximum possible signal-to-noise ratio (SNR) on MRI [14,16–18]. Calculation of the quantity of magnetic inhomogeneity in each pixel from data using the least-squares method was employed to generate the field map of IDEAL. By correcting phase shift in each pixel using this field map, robust fat suppression can be achieved, resulting in more accurate and confident interpretations in areas of B0 inhomogeneity.

The present study was performed to compare lesion conspicuity between T1-weighted fast spin echo (T1 FSE), fat-suppressed T2-weighted FSE (FS-T2 FSE), STIR, and T2-weighted FSE IDEAL sequences in terms of percent contrast and contrast-to-noise ratio (CNR) between focal myeloma lesion and background BM, and to assess the dependence of lesion conspicuity on myeloma cell mass in background BM in patients with multiple myeloma.

**Materials and Methods**

**Study participants**

All study protocols were approved by the appropriate institutional review boards (University Hospital Medical Information Network Clinical Trials Registry [UMIN- 112 CTR] number, UMIN000003663). Each participant provided written informed consent before undergoing MRI.

Spinal MRI was performed in 45 patients with multiple myeloma between June 2010 and November 2013. The criteria used for diagnosis were taken from the classification of Durie and Salmon [18]. Diagnoses were confirmed when there were more than 10% clonal plasma cells in BM samples of the iliac crest. We excluded patients who had undergone chemo- or radiotherapy. Patients comprised 25 men (mean age, 65.2 years; range, 52–81 years) and 20 women (mean age, 67.3 years; range, 55–78 years).

Of the 45 patients in the staging cohort, 8 had asymptomatic myeloma (Durie-Salmon stage I), and 37 had symptomatic myeloma (Durie-Salmon stage II, n = 21; Durie-Salmon stage III, n = 16). The distinction between symptomatic and asymptomatic myeloma depended on the presence or absence of myeloma-related organ dysfunction according to the criteria of the International Myeloma Working Group [18]. The 37 symptomatic patients had symptoms described in the CRAB criteria (renal insufficiency, n = 9; anemia, n = 24; focal lytic bone lesions, n = 37). Two authors (with 20 years of expertise in spinal imaging and 12 years of experience in hematology) reviewed all medical and clinical records to collect all available data.

**BM examination**

We estimated the percent of BM plasma cells (BMPC%) in BM biopsy specimens obtained from the iliac crest.

**Spinal MRI and quantitative study**

Imaging was performed using a 3.0-T MRI unit (Signa HDxt 3T; GE Healthcare Milwaukee, WI) and the following sequences (Table 1): sagittal T1 FSE; sagittal FS- T2 FSE (with a chemical shift selective (CHESS) technique); sagittal fast STIR imaging; and a sagittal IDEAL T2 FSE sequence. Co-registered water, fat, in-phase (water+fat) and out-of-phase (water-fat) images were generated by the IDEAL software.

Mean signal intensity and standard deviation were calculated by placing operator-determined regions of interest (ROIs) within the focal myeloma lesions and within BM without focal lesions. The ROI for BM was defined manually within the internal part of the L1–L3 vertebral bodies in the midsagittal images, as described elsewhere [19], which did not include focal myeloma lesions, because these spinal levels were less affected by degenerative disc disease compared to lower lumbar elements. Signal intensity values for BM were then calculated as the mean value obtained from the three vertebral bodies and used as background BM. The ROI was placed at the same location on all sequences. Each ROI had an area of 254–519 mm² for BM and 58–222 mm² for myeloma lesion. Focal myeloma lesion was defined as an area of low signal intensity on T1-weighted images and high or intermediate signal intensity on in-phase images of IDEAL with a relatively well-defined margin larger than 0.7 cm in the long axis. All focal myeloma lesions were confirmed to be shown as lytic lesions on computed tomography (CT). The largest focal lesion in each patient was measured. For each MRI examination, the percent contrast and CNR between BM and focal myeloma lesion (n = 45) was measured using the following equations:

\[
\text{Percent contrast} = \frac{(S_a - S_b)}{(S_a + S_b)}
\]

\[
\text{CNR} = \frac{|S_a - S_b|}{\sqrt{(S_{und}^2 + S_{had}^2)}}
\]

where \(S_a\) and \(S_b\) are mean signal intensity and \(S_{und}\) and \(S_{had}\) are the standard deviation of intensities of focal myeloma lesion and BM, respectively.

**Statistical analysis**

All statistical analyses were performed using statistical software (Excel 2007; Microsoft, Redmond, WA).

Spearman rank correlation coefficients (\(p\)) were calculated to investigate correlations between percent contrast, CNR and BMPC%. One-way analysis of variance with Scheffé’s post hoc
test was used to compare percent contrast and CNR among the four different groups (i.e., T1 FSE, FS- T2 FSE, fast STIR, and water image of IDEAL) for 45 patients with focal myeloma lesions. Differences were considered significant at the $P < 0.05$.

**Results**

A significant correlation was seen between percent contrast and BMPC%, with Spearman correlation coefficients of $-0.586$ ($P < 0.0001$), $-0.796$ ($P < 0.0001$), and $-0.494$ ($P < 0.01$) for water image of IDEAL, FS- T2 FSE, and STIR, respectively (Table 2). No significant correlation was evident between percent contrast and CNR with BMPC% for T1 FSE. Negative correlations between percent contrast and CNR with BMPC% for FS- T2 FSE were stronger than those for water image of IDEAL. Correlations between percent contrast and CNR with BMPC% for STIR were the weakest among the three fat-suppression techniques.

This result means that the higher the BMPC%, the less conspicuous the focal lesion is on fat-suppressed MRI. Therefore, in light of the difficulty of detecting focal lesions in low fat-containing marrow, i.e., a high tumor load BM, we categorized patients into two groups: a high tumor load BM group ($n = 15$), with BMPC% $\geq 45\%$; and a low tumor load BM group ($n = 30$), with BMPC% < 45%.

BM-focal lesion percent contrast and CNR were significantly greater for FS- T2 FSE than for STIR in the low tumor load BM group ($P < 0.001$, $P < 0.05$, respectively, Table 3, Fig. 2). In the low tumor load BM group, percent contrast was significantly higher for FS-T2 FSE than for water image of IDEAL ($P < 0.05$), but this was not the case in the comparison of CNR. No significant difference was found among the three fat-suppression methods in the high tumor load BM group, although mean values of BM-focal lesion percent contrast and CNR were highest for FS- T2 FSE and lowest for STIR.

Representative images are shown in Figures 3 and 4.

**Table 1.** Acquisition parameters for MR sequences.

| Sequence            | TR/TE/TI (ms) | NEX | FOV (mm) | Matrix | Slice thickness (mm) | Bandwidth (kHz) | Imaging time (mins) |
|---------------------|---------------|-----|----------|--------|----------------------|-----------------|--------------------|
| Water image of IDEAL | 4000/112.4    | 6   | 300      | 384×192| 4                    | 83.3            | 5:12               |
| Fat-suppressed T2 FSE | 4000/116     | 2   | 300      | 320×288| 4                    | 62.5            | 2:16               |
| Fast STIR           | 4200/106.8/190 | 2   | 300      | 320×224| 4                    | 31.2            | 3.01               |
| T1 FSE              | 700/11.8      | 2   | 300      | 512×224| 4                    | 41.7            | 2:38               |

NEX, number of signal averages.

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**Table 2.** Results of Spearman rank correlation for percent contrast and CNR with BMPC% among the four different sequences in multiple myeloma ($n = 45$).

| Sequence            | Percent contrast | CNR |
|---------------------|------------------|-----|
| IDEAL               | $r = -0.806$     | $-0.664$ |
| FS-T2 FSE           | $r = -0.796$     | $-0.709$ |
| Fast STIR           | $r = -0.494$     | $0.500$ |
| T1 FSE              | $r = -0.086$     | $0.57$ |

IDEAL, water image of IDEAL; FS-T2 FSE, fat-suppressed T2 FSE.

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Figure 1. Graph of the BMPC% versus percent contrast. The linear regression curve is shown. Vertical line corresponds to a BMPC% of 45%.

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Discussion

The presence of focal lesions on MRI has been correlated with shorter overall survival in several studies of patients with multiple myeloma [5,20]. Since the Durie-Salmon PLUS staging system does not mention about diffuse infiltration pattern of tumor cells in the BM, focal myeloma lesions must be detected regardless of any diffuse abnormality in background BM. In this study, BM-focal lesion percent contrasts and BMPC% showed significant negative correlations in the three fat suppression techniques. This means that the higher the BMPC% within BM, the less conspicuous the focal lesion is on fat-suppressed MRI. We attributed the lower lesion percent contrasts and BMPC% to increased signal intensity of background BM, mainly caused by T2 prolongation by diffusely infiltrated myeloma cells, which can reduce the signal intensity contrast between focal lesion and background BM. Furthermore, results of post hoc tests according to groups categorized by tumor load also demonstrated no significant difference among the three fat-suppression methods in the high tumor load BM group. This study did not identify any clearly superior fat-suppression technique for detecting focal myeloma lesions in the high tumor load BM. In clinical settings, other modalities such as CT with multiplanar reconstruction and PET/CT might be helpful to detect focal lesions.

BM-focal lesion percent contrast and CNR for FS-T2 FSE were significantly higher than STIR in the analyses for total lesions and for lesions in the low tumor load BM group. This finding can be explained by the improvement of the saturation pulse of CHESS technique in 3-T MRI. In 2011, Tagliafico et al. [21] compared 1.5- and 3-T MRI of the brachial plexus and demonstrated that CNR in 3-T MRI was significantly better in FS-T2 FSE sequences. On the other hand, Sormaala et al. [22] found no noteworthy differences in the sensitivity of 1.5- and 3-T images when they evaluated bone edema caused by acute bone stress in the foot using STIR. In addition, when imaging at 3 T, the longer T1 of tissue at higher field strengths would decrease lesion contrast if imaging time was the same as that at 1.5 T. Furthermore, even though STIR is insensitive to B1 inhomogeneities, it may be sensitive to B0 inhomogeneities, particularly at higher field strengths, where B1 may be more inhomogeneous. In addition, chemical shift is larger in 3 T than in 1.5 T. These facts may partly explain the opposite result in this study regarding the percent contrast of FS-T2 FSE T2 and STIR.

The present study showed that the percent contrast for FS-T2 FSE was significantly greater than that of water image of IDEAL in the low tumor load group. The exact cause is unclear, but may be related to the inherent homogeneity of magnetic fields in the lumbar spine area. In 2004, Ma et al. [23] reported that the fast three-point water-fat separation technique provides superior fat suppression and lesion conspicuity in the spine by 1.5 T, and can potentially be used as an alternative to T2-weighted imaging of the spine. They analyzed the whole spinal column, including the cervical and thoracic spine, as often being subject to susceptibility artifacts around the shoulder, mediastinum, or diaphragm. On the other hand, we investigated contrast characteristics focused on the lumbar spine, where fat suppression was fairly uniform over a field of view, which reduced the superiority of IDEAL for obtaining fat-suppressed images.

![Table 3. Comparison of percent contrast and CNR among the three fat-suppression sequences in multiple myeloma.](image)

|                    | IDEAL  | FS-T2 FSE | STIR  |
|--------------------|--------|-----------|-------|
| **Percent contrast**|        |           |       |
| Total (n=45)       | 0.52±0.21 | 0.60±0.19** | 0.44±0.22 |
| Low tumor load BM* (n=30) | 0.58±0.21 | 0.69±0.14*** | 0.51±0.24 |
| High tumor load BM** (n=15) | 0.40±0.14 | 0.43±0.15** | 0.32±0.10 |
| **CNR**            |        |           |       |
| Total (n=45)       | 7.17±3.70 | 7.47±3.74* | 5.97±4.24 |
| Low tumor load BM* (n=30) | 8.60±3.61 | 8.87±3.7* | 7.21±4.6 |
| High tumor load BM** (n=15) | 4.32±1.44 | 4.68±1.49** | 3.50±1.20 |

IDEAL, water image of IDEAL; FS-T2 FSE, fat-suppressed T2 FSE; BM, bone marrow.

Data are shown as mean ± standard deviation.
*Low tumor load BM: bone marrow with fat-signal fraction <45%.
**High tumor load BM: bone marrow with fat-signal fraction ≥45%.
*P<0.05, Fat-suppressed T2 FSE vs. water image of IDEAL.
**P<0.01,
***P<0.001, Fat-suppressed T2 FSE vs. fast STIR.

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Figure 2. Percent contrast and CNR comparison among the three different fat-suppression sequences. BM-focal lesion percent contrast (a) and CNR (b) are significantly greater for FS-T2 FSE than for STIR in the low tumor load BM groups (P<0.001, P<0.05, respectively). In the low tumor load BM group, percent contrast is significantly higher for FS-T2 FSE than for water image of IDEAL (P<0.05).

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Our protocol for assessment of bone lesion for multiple myeloma did not include diffusion-weighted imaging. In 2011, Sommer et al. reported the diagnostic potential of diffusion-weighted imaging with background suppression (DWIBS) in the detection of focal BM lesions from multiple myeloma [24]. According to that report, the CNR values provided by DWIBS in patients with high serum concentration of M-component are slightly higher than those of T2-weighted STIR. Since they did not perform CNR analysis for IDEAL or FS-T2 FSE, whether spinal diffusion-weighted imaging could yield better performance in delineation of focal myeloma lesions compared to IDEAL and FS-T2 FSE cannot be determined. However, diffusion-weighted images suffer from susceptibility artifacts and image distortion caused by eddy currents and cannot yet achieve high spatial resolution. Since the Durie–Salomon PLUS staging system does not mention lesion size, we believe that MRI with reasonable spatial resolution and without significant degradation is needed for the purpose of counting focal lesions.

Several limitations to this study must be considered when interpreting the present findings. First, the scanning parameters used for fat-suppression techniques were not identical, to adapt acquisition time to clinical practice. FS-T2 FSE used twice the bandwidth of STIR, presenting a more favorable condition for STIR regarding SNR. IDEAL used three times the NEX of the other two sequences, presenting a more favorable condition for IDEAL. We therefore consider that these differences in imaging parameters are unlikely to have had any substantial effect on our results. Second, biopsy specimens of spinal BM were not obtained in this study; instead, we calculated BMPC% from BM samples of the iliac crest. Neoplastic plasma cells tend to form clusters, which may be small or large [25]. Variability of the histopathological pattern and spatial distribution for multiple myeloma could have resulted in some error in this study. Other factors with possible differences in spatial distribution, including hematopoietic BM and degenerative disc disease, might thus have influenced the fat fraction of BM, resulting in altered signal intensity in fat-suppressed MRI. The normal distribution of hematopoietic BM in the adult, in which only the axial skeleton and proximal shafts of the femurs and humeri contain hematopoietic marrow, is completed by around 25 years old. With advancing age, generally beyond 40 years old, the vertebral BM becomes increasingly replaced with fatty marrow [26]. The BM also undergoes changes due to dietary changes, anemia, chronic hypoxia, chemotherapy, and other medications, through the actions of various cytokines [27,28]. Such interindividual variability of fatty marrow replacement could thus have affected our results. We also acknowledge that MRI in our study was limited to the lumbar vertebrae. The use of specific vertebral bodies might not be appropriate for assessing lesion conspicuity of focal myeloma lesions, which can occur in any bones susceptible to magnetic field inhomogeneity. Further studies on lesion conspicuity on whole-body MRI combined with information on the amount of myeloma cells are warranted.

In conclusion, the higher the BMPC% obtained from biopsy, the less conspicuous the focal lesion on fat-suppressed MRI. To the best of our knowledge, this dependence of lesion conspicuity on myeloma mass in the BM has not been described previously, and could have clinical implications for staging and treatment planning in cases of multiple myeloma. The most effective protocol for detecting focal lesions was FS-T2 FSE. No significant differences in lesion conspicuity were found among fat suppression techniques in the high tumor load BM group.

Author Contributions
Conceived and designed the experiments: MT TT Y. Kuroda YA. Performed the experiments: MT YA KT SD Y. Kaichi AS KA. Analyzed the data: MT TT. Contributed reagents/materials/analysis tools: MT YA TT. Wrote the paper: MT.

References
1. Dimopoulos M, Terpos E, Comenzo RL, Tosi P, Bekac M, et al. (2009) International myeloma working group consensus statement and guidelines regarding the current role of imaging techniques in the diagnosis and monitoring of multiple Myeloma. Leukemia 23:1545–1556.

2. Terpos E, Dimopoulos MA (2005) Myeloma bone disease: pathophysiology and management. Ann Oncol 16:1225–1231.

Figure 3. Sagittal MRI in a 58-year-old woman with focal myeloma lesions in the low tumor load group (BMPC%, 12%). a) Water image of IDEAL; b) FS-T2 FSE; c) STIR; and d) T1 FSE. ROIs of lesion and background BM used for the percent contrast computation are demarcated in the images. Actual ROIs of BM are placed in similar lesion and background BM used for the percent contrast computation are demarcated in the images. Among the three fat-suppressed images, signal hyperintensity of the lesion is less conspicuous in STIR (c) than in water image of IDEAL (a), which seems to be due to the lower SNR in STIR.

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Figure 4. Sagittal MRI in a 65-year-old man with combined focal and diffuse infiltration pattern (BMPC%, 71%). a) Water image of IDEAL; b) FS-T2 FSE; c) STIR; d) T1 FSE; and e) CT. ROIs of lesion and background BM used for the percent contrast computation are demarcated in the images. The actual ROIs of BM are placed in similar locations in the midsagittal images. The focal myeloma lesion in L4 is seen as a slightly high intensity signal on fat-suppressed images (white arrows) but is less conspicuous compared to the lesion in Figure 3. Among the three fat-suppressed images, signal hyperintensity of the lesion is less conspicuous in STIR (c) than in water image of IDEAL (a) or FS-T2 FSE (b). The focal lesion is not detected on T1 FSE (d).

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3. Kyle RA, Gertz MA, Witzig TE, Lust JA, Lacy MQ, et al. (2003) Review of 1027 patients with newly diagnosed multiple myeloma. Mayo Clin Proc 78:21–33.

4. Durie BG, Kyle RA, Belch A, Bensinger W, Blade J, et al. (2003) Myeloma management guidelines: a consensus report from the Scientific Advisors of the International Myeloma Foundation. Hematol J 4:379–398.

5. Walker R, Barlogie B, Haeseler J, Tricot G, Anaisie E, et al. (2007) Magnetic resonance imaging in multiple myeloma: diagnostic and clinical implications. J Clin Oncol 25:1121–1129.

6. Bau-Melnik A, Buhlmann S, Durr HR, Reiser M (2005) Role of MRI for the diagnosis and prognosis of multiple myeloma. Eur J Radiology 55:56–63.

7. Fleckenstein JL, Archer BT, Barker BA, Vaughan JT, Parkey RW, et al. (1991) Fast short-tau inversion-recovery MR imaging. Radiology 179:99–104.

8. Rahmouni A, Divine M, Mathieu D, Golli M, Dao TH, et al. (1993) Detection of multiple myeloma involving the spine: efficacy of fat suppression and contrast enhanced MR imaging. AJR Aus J Roentgenol 160:1049–1052.

9. Nakatsu M, Hatabu H, Itoh H, Morikawa K, Miki Y, et al. (2000) Comparison of Short Inversion Time Inversion Recovery (STIR) and Fat-Saturated (chemsat) Techniques for Background Fat Intensity Suppression in Cervical and Thoracic MR Imaging. J Mag Reson Imaging 11:56–60.

10. Dixon WT (1984) Simple proton spectroscopic imaging. Radiology 1984;153:189–194.

11. Yeung HN, Korman DW (1986) Separation of true fat and water images by correcting magnetic field inhomogeneity in situ. Radiology 159:783–786.

12. Glover GH, Schneider E (1991) Three-point Dixon technique for true water/fat decomposition with B0 inhomogeneity correction. Magn Reson Med 18:371–383.

13. Reeder SB, Wen Z, Yu H, Pineda AR, Gold GE, et al. (2004) Multicoil Dixon chemical species separation with an iterative least-squares estimation method. Magn Reson Imaging 1:521–530.

14. Reeder SB, Chen YF, Shimakawa A, Brittain JH, Pele N, et al. (2005) Field map estimation with a region growing scheme for iterative 3-point water-fat decomposition. Magn Reson Med 54:1032–1039.

15. Ricci C, Costa M, Kang YS, Yang A, Rahmouni A, et al. (1990) Normal age-related patterns of cellular and fatty bone marrow distribution in the axial skeleton. Magn Reson Imaging 18:371–383.

16. Travlos GS (1985) Red-yellow marrow conversion: its effect on the location of some solitary bone lesions. Skeletal Radiol 14:10–19.

17. Gerdes CM, Kijowski R, Reeder SB (2007) IDEAL imaging of the musculoskeletal system: robust water fat separation for uniform fat suppression, marrow evaluation, and cartilage imaging. AJR Am J Roentgenol 189:W204–W291.

18. International Myeloma Working Group (2003) Criteria for the classification of monoclonal gammopathies, multiple myeloma and related disorders: a report of the International Myeloma Working Group. Br J Haematol 121:749–757.

19. Takasu M, Tani C, Sakoda Y, Ishikawa M, Taniki K, et al. (2012) Iterative decomposition of water and fat with echo asymmetry and least-squares estimation (IDEAL) imaging of multiple myeloma: initial clinical efficiency results. Eur Radiol 22:1114–1121.

20. Moulopoulos LA, Gika D, Anagnostopoulos A, Delassalle K, Weber D, et al. (2005) Prognostic significance of magnetic resonance imaging of BM in previously untreated patients with multiple myeloma. Ann Oncol 16:1824–1829.

21. Tagliafaico A, Saccio G, Neumaier CE, Serafini G, Ghidara M, et al. (2011) MR imaging of the brachial plexus: comparison between 1.5-T and 3-T MR imaging: preliminary experience. Skeletal Radiol 40:717–724.

22. Normolla MJ, Ruchhola JP, Mattila VM, Koskinen SK, Pihlajamaki HK (2011) Comparison of 1.5T and 3T MRI scanners in evaluation of acute bone stress in the foot. BMC Musculoskelet Disord 12:128.

23. Ma J, Singh SK, Kumar AJ, Leed ES, Zhan J (2004) T2-weighted spine imaging with a fast three-point dixon technique: comparison with chemical shift selective fat suppression. J Magn Reson Imaging 20:1025–1029.

24. Sommer G, Klarhoefer M, Lenz C, Scheffler K, Bongartz G, et al. (2011) 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 21:857–862.

25. Hanrahan CJ, Christensen CR, Cram JR (2010) Current Concepts in the Evaluation of Multiple Myeloma with MR Imaging and FDG PET/CT. Radiographics 30:127–142.

26. Ricci C, Costa M, Kang YS, Yang A, Rahmouni A, et al. (1990) Normal age-related patterns of cellular and fatty bone marrow distribution in the axial skeleton. Magn Reson Imaging 18:371–383.

27. Kricun ME (1985) Red-yellow marrow conversion: its effect on the location of some solitary bone lesions. Skeletal Radiol 14:10–19.

28. Travlos GS (2006) Normal structure, function, and histology of the bone marrow. Toxicol Pathol 34:548–565.