The Use of Gadolinium in Musculoskeletal MRI—Time to Rethink?

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Abstract

Magnetic resonance imaging has continued to evolve over the recent decades, in part, due to the evolution of gadolinium-based contrast agents and their use. These were initially thought to have a relatively low-risk profile. However, there is mounting evidence that trace amounts of gadolinium are retained within the body. To ascertain the current use of gadolinium in medical practice, we performed a survey of musculoskeletal radiologists, within the United Kingdom, Europe and India. The survey demonstrated varied practices amongst all radiologists with relatively indiscriminate use of gadolinium. In this review, we discuss the current evidence for and against the use of gadolinium in musculoskeletal magnetic resonance imaging.

Keywords
► gadolinium
► contrast
► musculoskeletal imaging
► brain deposition

Background

Magnetic resonance imaging (MRI) is a sensitive noninvasive modality with several advantages in comparison to other imaging techniques.¹ It is the primary imaging modality for the detailed evaluation of a broad spectrum of musculoskeletal (MSK) disease processes.² This is due to its high-resolution providing unparalleled soft tissue contrast and allowing the visualization of both anatomical structures and pathological processes.³

The modality of choice for tumors and tumor-like conditions is often MRI, owing to its excellent soft tissue contrast, its sensitivity to bone marrow and soft tissue edema, and its multiplanar imaging.¹,⁴ It is key for diagnosing, staging, preoperative work-up, and follow-up of patients with benign and malignant soft tissue neoplasms.³–⁵ Furthermore, it provides detailed tissue characterization and aids in the staging of bone lesions.² MRI is also useful in the evaluation of trauma,²,⁶ infection, and neuromuscular disease.³

The 1980s brought about new advances in MSK MRI with the development of gadolinium-based contrast agents (GBCAs).⁷ In its free form, unpaired gadolinium electrons are highly toxic.⁸ Thus, to reduce their toxicity and improve stability, they are bound to a ligand and administered in chelated forms.⁸,⁹

The use of GBCAs has grown substantially; they are used in approximately one-in-three of all MRI studies worldwide.⁹,¹⁰ GBCAs are used in an attempt to improve diagnostic confidence to influence patient care and management.¹¹ In MSK MRI GBCAs are often used in the assessment of soft tissue sarcomas (STS) prior to histological diagnosis and in follow-up imaging to assess for local recurrence.¹² They help radiologists plan soft tissue biopsies by identifying viable enhancing malignant tissue from cystic/necrotic tissue.¹¹ Contrast can enable the detection of the early stages of soft tissue infection and differentiate phlegmon from normal surrounding tissues.¹³ It is also useful to gauge the extent of the infections and make abscess/collections more conspicuous.¹³ GBCAs can provide accurate representation of
the degree of osseous and nonosseous involvement in complicated extremity infections. Additionally, GBCAs can aid in the diagnosis of infections in septic arthritis, acute-subacute- and chronic osteomyelitis. The role of GBCAs in spinal disease will not be covered in this article.

**Mechanism of Action**

Gadolinium (Gd$^{3+}$), in its raw form is a paramagnetic ion composed of seven unpaired electrons resulting in a highly magnetic effect. Administration of Gd$^{3+}$ falters the rotation frequency of water molecules, shortening both T1 and T2 relaxation times of tissues in which it accumulates, thus allowing differentiation through increasing signal intensity on T1 sequences and decreasing signal on T2 sequences. GBCAs are distributed within the blood and extravascular-extracellular space. They are biologically inert and generally eliminated by the kidneys.

As previously mentioned, Gd$^{3+}$ in its free form is highly toxic. Its structure makes it unstable in vivo and therefore it is bound to a ligand and administered in chelated forms. Pharmacologically GBCAs are classified according to the molecular structure of the chelating ligand to which they are bound. These are classified as linear or macrocyclic. The chelating ligand compounds are designed to minimize dissociation of gadolinium. It is because of this; GBCAs were expected to have high contrast efficiency and safety in addition to their rapid excretion, high stability, low osmolality, and low viscosity.

**Current Practice**

In an attempt to assess current practices regarding the use of GBCAs, we anonymously surveyed members of the British Society of Skeletal Radiologists (BSSR), European Society of Musculoskeletal Radiology (ESSR), and the Musculoskeletal Society of India (MSS) regarding the use of gadolinium use in MSK MRI.

**Table 1** A simple anonymous yes/no web-based survey of eight questions was sent to members of British Society of Skeletal Radiologists (BSSR), European Society of Musculoskeletal Radiology (ESSR), and the Musculoskeletal Society of India (MSS) regarding the use of gadolinium use in MSK MRI.

<table>
<thead>
<tr>
<th>No.</th>
<th>Questions</th>
<th>Choices</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Do you use contrast in musculoskeletal MR imaging?</td>
<td>Yes</td>
</tr>
<tr>
<td>2</td>
<td>Relative proportion of post contrast imaging in your practice?</td>
<td>$&lt;25%$, $25%–50%$, $&gt;50%$</td>
</tr>
<tr>
<td>3</td>
<td>Post contrast imaging in soft tissue lump?</td>
<td>Yes</td>
</tr>
<tr>
<td>4</td>
<td>Post-contrast imaging in bone lesions?</td>
<td>Yes</td>
</tr>
<tr>
<td>5</td>
<td>Post-contrast imaging in soft tissue infection?</td>
<td>Yes</td>
</tr>
<tr>
<td>6</td>
<td>Post-contrast imaging in bone infection?</td>
<td>Yes</td>
</tr>
<tr>
<td>7</td>
<td>Where do you work?</td>
<td>Tertiary referral center, University hospital, District hospital, Private clinic</td>
</tr>
<tr>
<td>8</td>
<td>Do you check eGFR before administration of contrast?</td>
<td>Yes</td>
</tr>
</tbody>
</table>

Abbreviation: eGFR, estimated glomerular filtration rate.

In addition to its key role in the development of nephrogenic systemic fibrosis (NSF), there is evidence of Gd$^{3+}$ deposition in patients receiving GBCAs despite having an intact blood-brain barrier and normal renal function. This was first reported in 2010 by Xia et al with the discovery of insoluble deposits of gadolinium in the biopsies of brain tumor patients, all of whom had at least one CEMRI scan with a linear chelating agent in their past. In 2013, Kanda et al reported an association between GBCAs administration and NSF. Elawad et al found this to be connected with changes of the subcortical gray matter on MRI of the brain. Unenhanced T1-weighted images showed a positive correlation with previous exposure to nonionic

Current Evidence of GBCAs Deposition

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linear chelating type GBCAs, demonstrating areas of high-intensity signals bilaterally in the globus pallidus (GP) and in the dentate nuclei (DN). Extracranial sites of gadolinium deposition have been reported in the liver, skin, and bones. Previously, it has been suggested in studies by Roccatagliata and colleagues, that multiple sclerosis was associated with hyperintense DN presence on MRI. Similarly, Kasahara et al proposed an association between hyperintense DN findings with a history of brain irradiation. However, Kanda et al found these changes were in fact associated with previous
GBCAs administrations, rather than any relation to history of multiple sclerosis or brain irradiation.20 This was further supported by Ranga et al as hyperintense DN findings in irradiated patients were found to be likely related to gadolinium deposition.18

In addition, a positive dose–response correlation between the number of previous GBCAs administrations, and high signal intensity in the DN and GP was established by Kanda et al.20 Similar findings in pediatric patients were illustrated in case reports by Miller et al and Roberts and Holden, in which cumulative doses of administered GBCAs demonstrate significant changes in signal intensity.26,27

Brain specimens of patients with a history of receiving linear GBCAs were evaluated along with a control group. These studies revealed that there were increased gadolinium deposits in the GP and DN compared with other brain regions.28–30 Several studies by Radbruch et al and others support the hypothesis that gadolinium accumulation in the deep brain nuclei is associated with linear GBCAs and not macrocyclic GBCAs.29–33

### Risks and Side Effects of Gadolinium Deposition

Apart from the rare incidence of NSF, the potential impact of long-term Gd$^{3+}$ retention remains unknown. The risk of developing adverse effects following gadolinium deposition in the brain is significantly increased in patients who are subjected to multiple scans throughout their lifetime, using GBCAs.31 Those with chronic conditions who undergo serial surveillance scans or patients who have interval follow-up scans are at increased risks of gadolinium brain deposition.34

In addition, young children due to their age and expected long lifespan, bear considerable risk and should, therefore, be given the appropriate consideration and risk assessment to minimize exposure and deposition.34

A large prospective cohort study with a control group was designed by Parillo et al to evaluate the occurrence of symptoms within 24 hours after GBCA administration. These symptoms have been grouped as gadolinium deposition disease (GDD). Findings showed an increased incidence of new symptoms within the first 24 hours subsequent to GBCA exposure, in comparison to after unenhanced MRI.35 Patients reported symptoms of fatigue, dizziness, mental confusion, and diarrhea.35 In a separate study, performed by Burke et al which consisted of anonymous patient surveys, 66% of respondents self-reported immediate adverse manifestations experienced following GBCA administration, 32% within 6 weeks, and 2% complained of symptoms within a 6-month period.36 More than 77% reported side effects such as headaches, visual changes, auditory changes, and bone/joint pains.36 Skin changes, such as thickening and discoloration, were reported by >50% of those surveyed, while respiratory (difficulty in breathing) and digestive (nausea, vomiting, diarrhea) changes were felt by >40%.36 All respondents related their symptoms to their previous GBCA exposure. Although this survey suggests a temporal relationship between gadolinium and the reported symptoms, the invalidity of self-report surveys indicated that further research is advised.

With the DN being the most noted site of gadolinium deposition, adverse effects relating to its functions of planning, initiation, and control of voluntary movements are expected.21 However, none have been reported in relation to any GBCA exposure.

Well-controlled studies to investigate the adverse biological and/or neurological side effects of GBCAs administration are essential both to conclude the short- and long-term effects of gadolinium deposition in the brain. In addition, data linking these adverse effects to gadolinium deposition in the brain must also be established. Despite being unproven scientifically, there has been an increase of GDD-related litigation and personal injury advertising in the United States of America targeting potential GDD patients.

### Regulatory Changes Related to GBCA

With the emergence of new evidence regarding GBCAs retention in the brain, guidelines have been amended after investigations in 2016 by the Pharmacovigilance Risk Assessment Committee (European Medicines Agency) and submitted recommendations to the Committee of Medicinal Products for Human Use in 2017.37 As a result, the Royal College of Radiologists has updated its guidance on the use of Gd$^{3+}$ (8). The new guidelines highlight the suspension, withdrawal, or alterations in the use of some linear chelate GBCAs.7 The changes highlight the need to reconsider the routine use of gadolinium unless the diagnostic need outweighs possible unknown future complications.

### Gadolinium in Food

Research in Germany has found traces of gadolinium in beverages such as Coca-Cola.38 This has been attributed to gadolinium in the urine excreted by patients post-CEMRI and entering municipal wastewater treatment systems. Gadolinium is not removed or purified by the treatment systems and as a result enters the public water supply and subsequently the environment. Schmidt et al reported six major German cities had gadolinium polluted water systems and present in food and beverages from McDonald’s and Burger King.38 In addition, Thomsen noted that the gadolinium concentration levels are rising slowly and remain persistent in water thus causing a growing concern.39 Currently, no clinical adverse effects have been reported although the long-term implications remain unknown.

### Current Evidence Supporting the Use of Gadolinium

Currently, the use of gadolinium in MSK MR can be classified into three main groups: tumors, infections, and joint pathology.
**Soft Tissue Sarcomas**

In the evaluation of sarcomas and sarcoma like-lesions, gadolinium is thought to improve diagnostic accuracy and provide additional data in staging, biopsy planning, tissue characterization, evaluation of response to chemotherapy, and detection of recurrence. Gadolinium has been reported to increase the sensitivity of recurrent STS by 74% compared with unenhanced MRI. GBCAs allow the assessment of intra- or extra-compartmental extent and involvement of adjacent bone, joint, muscle, or neurovascular involvement. In addition, biopsy planning is made easier by improved tissue enhancement highlighting necrotic and/or cystic areas to be avoided. It can be difficult to distinguish between true cystic lesions and cystic-like solid lesions on noncontrast MRI (e.g., myxoid lesions); gadolinium allows this distinction to be readily made due to the lack of enhancement of true cystic lesions. One relies on tumor enhancement to make tumors more conspicuous when there is significant peritumoral edema. Gadolinium can also be of value in the evaluation of hemorrhagic lesions. It will uncover enhancing tumors masked by the surrounding hemorrhage.

Dynamic contrast enhanced (DCE) MRI is a technique that allows the evaluation of the temporal pattern of enhancement in tumors, monitoring response to neoadjuvant chemotherapy, and assessing for tumor recurrence. Other uses include distinguishing adjacent inflammatory processes and bone tumor perfusion. Postoperative gadolinium is useful in distinguishing underlying collections from surrounding inflammatory change and recurrence.

**Bone Tumors**

With regard to bone tumors MRI is primarily performed for local staging and extent rather than diagnosis and therefore there is no requirement of contrast. Radiological diagnosis is predominantly based on radiography. However, GBCAs may provide information on the assessment of intramedullary extension, the extension to adjacent structures and can be useful in post-surgical follow-up imaging. For selective bone tumors, such as osteosarcoma, gadolinium offers the potential for determining the efficacy of chemotherapy, by evaluating tumor necrosis prior and subsequent to chemotherapy. Sarcomas close to joints, gadolinium may aid in determining whether tumor resection should be intra- or extra-articular. One may need to give contrast when faced with equivocal imaging findings for suspected osteoid osteomas (OO), as dynamic MRI increases nidus conspicuity. It should be noted that in our center of 793 cases of OO over a 12-year period, we have never needed to use dynamic imaging for cases of OO. The results of our survey demonstrate that a large proportion of radiologists are using GBCAs for bone lesions, and this is disproportionately higher in Europe/India.

**Arthritis**

Several studies have shown that GBCA-enhanced MRI is beneficial in discriminating active from dormant arthritis.

For tenosynovitis, sensitivity and specificity are decreased without gadolinium contrast administration. Gadolinium contrast administration increases sensitivity when evaluating synovitis and tenosynovitis in early arthritis. A study by Reiser and coworkers examining both knee and wrist joints showed the use of gadolinium contrast markedly increased the enhancement between pannus and effusion, improving detection. In addition, CEMRI improved the evaluation of the pannus extension in the joint cavity, and into the suprapatellar recess. Furthermore, GBCA was effective in tracking the therapeutic effectiveness of treatment and found to be valuable in the selection process of patients suitable for synovectomy.

**Infections**

Gadolinium’s role was suggested as an aid to clarify the extent of active infections, distinguishing infectious from noninfectious inflammatory lesions, and in highlighting soft tissue abscesses. Characterization of focal collections and differentiation of abscesses from surrounding cellulitis/myositis were both improved by gadolinium. In septic arthritis, CEMRI was found to be useful in the evaluation of synovial hypertrophy. In addition, synovitis was more easily differentiated from simple joint effusion using GBCAs. Hopkins et al showed that the major role of CEMRI lies in diagnosing osteomyelitis and distinguishing it from neuropathic disease. Gd3+ can be especially useful when imaging the diabetic foot and to help differentiate osteomyelitis from Charcot’s arthropathy. Differentiating between the two requires careful evaluation of the patient, including medical history, physical examination, selected laboratory findings, and imaging studies. The use of contrast demonstrates areas of nonenhancement amongst enhancing inflammatory tissue allow necrotic regions in bone or abscesses to become more conspicuous and suggestive of osteomyelitis. However, one must proceed with caution if administering gadolinium due to potential complications with contrast nephropathies in poorly controlled diabetic patients.

**Joint Pathologies**

The evaluation of internal joint pathology with the use of gadolinium has been well established. Intra-articular Gd3+ aids in the assessment of labrum or cartilage. MR arthrography has advantages over conventional MR imaging owing to distention of the joint capsule, outlining the intra-articular structures, and hence delineating the abnormalities. This can be achieved by direct injection into the joint or indirect via intravenous gadolinium administration.

**Current Evidence against the Use of Gadolinium**

**Tumors**

Numerous studies have shown that gadolinium has done little to improve the diagnostic specificity of MRI. May et al
reported CEMRI did not provide additional information in 89% of cases and only led to changes in the management in 10% of patients (Fig. 4). Gadolinium-enhanced imaging did not lead to a reliable distinction between lesions, surrounding edema and fibrovascular tissue present in organizing hematomas which may show enhancement similar to tumor nodules. Additionally, cystic regions have particular signal characteristics on noncontrast images which should be identifiable, particularly if a fluid–fluid level is present (Fig. 5). The associated costs of GBCAs and increased length of examination associated with the acquisition of specific post-contrast sequences must also be considered. Furthermore, patients may not tolerate the increased length of scanning time, risking image degradation from movement artifact. It can be argued that the routine use of GBCAs for the evaluation of soft tissue tumors has negligible benefits. Its effectiveness in the evaluation and staging of MSK neoplasms is controversial. GBCAs should only be administered if it will change management. Indeterminate or aggressive appearing soft tissue lesions will often undergo an image-guided biopsy. If MRI demonstrates cystic/necrotic areas, ultrasound will allow the identification of solid areas to target, therefore, negating the need for GBCA use (Fig. 6). GBCAs rarely provide additional information during the assessment of primary bone lesions (Fig. 7).

**Infections**

Gadolinium does not allow radiologist to reliably distinguish infectious from noninfectious inflammatory conditions. Although evidence has shown gadolinium-enhanced MRI to be a highly sensitive (89–100%) technique in diagnosing MSK infections, the use of GBCAs varies in specificity from 46 to

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**Fig. 4** Axial MRI (a) T1, (b) STIR, and (c) T1FS post-contrast of a 63-year-old female with histologically proven undifferentiated pleomorphic sarcoma. There is an aggressive appearing soft tissue lesion which will require biopsy. The post-contrast image (c) confirms the mass is solid and demonstrates nonenhancing necrotic areas within the tumor. Ultrasound can clearly demonstrate both solid and necrotic components (image not shown) thus allowing a biopsy to be obtained from the most appropriate region of the lesion. The addition of the gadolinium, in this case, did not provide additional useful diagnostic information in either making the diagnosis or deciding if the lesion required a biopsy. MRI, magnetic resonance imaging.

**Fig. 5** Axial MRI (a) T1, (b) T2, and (c) T1FS post-contrast of a 74-year-old male with histologically proven undifferentiated pleomorphic sarcoma. This is a predominantly necrotic lesion with areas of hemorrhage shown as intralesional areas of high T1 signal. The low T1/high T2 areas are necrotic/cystic areas and the enhancing wall confirms the solid component as seen in (c). However, gadolinium use did not provide information on the type of aggressive mass and solid areas for target for biopsy seen on ultrasound (image not shown). MRI, magnetic resonance imaging.
88% and as a result often does not lead to alterations in patient care.\textsuperscript{54}

**Arthritis and Joint Pathologies**

GBCA can be useful in demonstrating the enhancement of the synovium but is unable to differentiate between similar inflammatory lesions.\textsuperscript{56} Rheumatoid and septic arthritis show similar enhancements.\textsuperscript{57} Doppler ultrasound should be considered in the first instance for the assessment of tenosynovitis/synovitis. Diffusion-weighted imaging has shown promising results in detecting synovitis and may be a novel noninvasive approach to contrast-free imaging of synovitis.\textsuperscript{58} The resolution of 3T MRI is such that there is mounting evidence supporting the use of unenhanced 3T MRI to evaluate the hip labrum.\textsuperscript{59,60} Evidence is not as clear for the glenoid labrum.\textsuperscript{61–64} The lack of 3T magnets and expertise in image interpretation means traditional arthrograms are still performed. In the future, arthrograms, particularly of the hip and shoulder, may become obsolete as 3T imaging becomes the standard. 3T MRI allows higher resolution and implements a small field of view strategies to improve spatial resolution, negating the need for contrast.

**Conclusion**

It is important to recognize the role of gadolinium in specific clinical settings such as infection and post-surgical follow-up of soft tissue tumors (\textsuperscript{64}Table 2). However, the use of GBCAs in MSK imaging is not without controversy and as innocuous as previously thought. Despite the unknown clinical implications, the mounting evidence of deposition in the body, contamination of water supplies as well as the food chain, and potential medicolegal implications, one should give due consideration before proceeding with its use. In conclusion, the use of GBCAs should be assessed on a case-by-case basis depending on the clinical scenario and merit rather than routine protocol.
## References


### Table 2 Summary of proposed recommendation for the use of gadolinium in MSK MRI

| Low-level indications for the use of gadolinium: | • Bone tumors (exception of dynamic imaging for osteoid osteoma if required). |
| Intermediate level indications for the use of gadolinium: | • Solid soft tissue tumors (except for necrotic or myxoid cases where ultrasound is not available). |
| High-level indications for the use of gadolinium: | • Trauma or internal derangements (joints). |
| | • Arthritis (need to differentiate synovitis/pannus from effusions). |
| | • Diabetic foot. |
| | • Necrotic soft tissue tumors (to aid biopsy planning if ultrasound not available). |

Abbreviation: MRI, magnetic resonance imaging; MSK, musculoskeletal.

Note: The use of gadolinium has been grouped into low, intermediate, and high-level indications. A low level represents cases where gadolinium should not be used, as no added clinical benefit is seen. Intermediate and high levels represent cases which may provide additional benefit with the use of gadolinium; however, this should be considered on a case-by-case basis.

**Funding**

None.

**Conflict of Interest**

None declared.