GUIDELINES

The future of MRI in radiation therapy: Challenges and opportunities for the MR community

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Abstract
Radiation therapy is a major component of cancer treatment pathways worldwide. The main aim of this treatment is to achieve tumor control through the delivery of ionizing radiation while preserving healthy tissues for minimal radiation toxicity. Because radiation therapy relies on accurate localization of the target and surrounding tissues, imaging plays a crucial role throughout the treatment chain. In the treatment planning phase, radiological images are essential for defining target volumes and organs-at-risk, as well as providing elemental composition (e.g., electron density) information for radiation dose calculations. At treatment, onboard imaging informs patient setup and could be used to guide radiation dose placement for sites affected by motion. Imaging is also an important tool for treatment response assessment and treatment plan adaptation. MRI, with its excellent soft tissue contrast and capacity to probe functional tissue properties, holds great untapped potential for transforming treatment paradigms in radiation therapy. The MR in Radiation Therapy ISMRM Study Group was established to provide a forum within the MR community to discuss the unmet needs and fuel opportunities for further advancement of MRI for radiation therapy applications. During the summer of 2021, the study group organized its first virtual workshop, attended by a diverse international group of clinicians, scientists, and clinical physicists, to explore our predictions for the future of MRI in radiation therapy for the next 25 years. This article reviews the main findings from the event and considers the opportunities and challenges of reaching our vision for the future in this expanding field.

KEYWORDS
future, MR, radiation therapy, ISMRM workshop

On behalf of the ISMRM MR in Radiation Therapy Study Group.
For affiliation refer to page 11.

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1 | INTRODUCTION

Radiation therapy (RT), prescribed to ~50% of cancer patients, is a major component of cancer treatment pathways.1,2 The aim of RT is to deliver a sufficiently high dose of ionizing radiation to the tumor to control disease while limiting the dose to healthy tissues for minimal radiation toxicity. The most common RT modality, external-beam RT, delivers megavoltage beams of X-rays via a linear accelerator (Linac) mounted on a gantry that rotates around the patient. Carefully optimized treatment plans tailor beam profiles and photon intensities to focus the prescribed dose to the target volume(s) and minimize exposure to surrounding healthy tissue.3,4 Typically, an RT course is run over 5 to 30 treatment sessions, called fractions that span 1 to 9 weeks. This article focuses on external-beam RT, but some aspects are also applicable to other forms of RT, such as brachytherapy and proton therapy.

Imaging is performed at multiple points in the RT treatment chain with multiple objectives, which can be broadly categorized as: “delineation and dosing,” “guidance and targeting,” and “response and adaptation.”5 Delineation and dosing goals are met by acquiring CT simulation (CT-Sim) images, often with complementary MRI or PET scans. Using these images, radiation oncologists delineate targets and radiosensitive normal tissues. Delineated targets and critical structures inform the treatment planning process that simulates and optimizes the planned dose distribution. Imaging is also used to inform guidance and targeting during treatment delivery. Onboard imaging is a vital component of modern Linac systems, which typically use mounted X-ray systems capable of planar images and cone-beam CT (CBCT) to align the tumor target with the position specified in the treatment plan through rigid couch adjustments each day of treatment.6 In some cases, imaging may also be used to adjust the radiation beams to compensate for internal anatomic changes,7 a process referred to as adaptive therapy. Because courses of RT span multiple fractions, there is room for tumor response assessment with CT, MRI, or PET to inform online or offline treatment plan adaptation.8,9 These imaging techniques for response and adaptation objectives are active areas of clinical development.

Although X-ray and CT-based technologies currently dominate imaging in RT, MRI is quickly growing in this application and has significant untapped potential to improve the field. Figure 1 illustrates conventional, state-of-the-art, and our vision for the future of RT workflows. MRI’s superior soft-tissue visualization in comparison to CT will improve delineation and dosing, allowing for reduced uncertainty in target localization and, therefore, more accurate treatments with the ability to safely deliver higher doses.10–14 The availability of functional MRI, or quantitate MRI (qMRI), techniques, such as DWI and oxygen-enhanced MRI (OE-MRI) could add useful contrasts and may be able to identify high-risk regions of the target that would benefit from dose boosts.15 New standards for guidance and targeting have been created by the recent availability of commercial MR-Linac systems that allow concurrent imaging during treatment for MRI-guided RT (MRgRT).11,16–19 Last, response and adaptation could be advanced through MRI’s potential for quantifying tumor radiation sensitivity.20 Adapting dosing to treatment response between fractions may improve patient outcomes and could be achieved either using diagnostic MR systems21 or MR-Linacs.22

This article is a summary of the findings from a 2021 workshop held by the ISMRM MR in RT study group on the future of MRI in RT. The aim of the workshop was to explore study group members’ predictions for the future of this expanding field in 25 years’ time. An international assembly of MRI and RT scientists, clinicians, and clinical physicists met virtually to discuss the long-term opportunities and challenges of this expanding field of research. Given the nature of this topic, some predictions are based on current literature, but we also consider the consensus of expert opinions at the workshop, for which limited literature is available to cite. Here, we reach out to the MR community to elucidate the unmet needs that must be addressed for our vision in Figure 1 to become a reality.

It should be noted that because of the tremendous flexibility of machine learning approaches, these will often be at the core of this article’s suggestions for overcoming the challenges of MRI in RT. However, where machine learning is applied in future clinical workflows, a consensus should be established on the validation, testing, and quality assurance (QA) of the techniques.23 Despite its great potential, machine learning should not be thought of as a silver bullet. It is well documented that machine learning solutions can behave poorly, for instance, when input data are out of distribution.24,25 It is, therefore, important we are prepared to address this secondary set of challenges for clinical implementation.

2 | DELINEATION

Delineation, or contouring, is performed by radiation oncologists using a combination of CT-Sim and co-registered MRI or PET scans to outline a set of 3D contours (i.e., treatment targets and critical structures). These contours are defined by The International Commission on Radiation Units and Measurements26 guidelines, and include the gross tumor volume (GTV), clinical target volume (CTV), planning target volume (PTV), and selected normal tissues, termed the organs at risk (OARs).27 As
The role of imaging for radiation therapy (RT) in conventional, state-of-the-art, and future workflows. The conventional workflow (top row) begins the pre-treatment phase by scanning the patient in a computed tomography simulator (CT-Sim), where the patient setup for treatment is simulated using the same flat-top couch and positioning devices. MR scans are also acquired and registered to the CT images. Target volumes are delineated manually on MRI and dose distributions are simulated and optimized using the CT images. At treatment, patient setup on a Linac system is aided by onboard cone-beam CT (CBCT) or planar X-ray. The patient must return daily for repeated treatment fractions over the course of several weeks. The middle row illustrates a state-of-the-art RT treatment chain. This MR-only workflow replaces CT-Sim with MR-Sim, reducing the burden on hospitals and patients. Artificial intelligence (AI) assisted contouring increases the efficiency and reliability of delineation (2 DELINEATION). Treatment plans are calculated using synthetic CT generated from MR-Sim images, eliminating CT-MRI registration errors (3 DOSE CALCULATION). At treatment, hybrid MR-Linac systems (6 HARDWARE) will facilitate the safe reduction of treatment margins via MRI-informed adaptation to the daily anatomy and gated deliveries for moving targets (4 IMAGE GUIDANCE). Treatment sessions are more labor intensive than conventional treatments, but could lead to fewer patient visits overall (7 REDUCING PATIENT BURDEN). In the future (bottom row), an MR-Linac-only workflow without a pre-treatment workup may be possible, where planning and treatment delivery is performed within minutes on the same system. Functional and structural MR imaging could inform AI-driven algorithms to generate plans without input from clinicians. MR-derived biomarkers (5 QUANTITATIVE MRI) hold the potential to establish new, contourless dose planning approaches, with information now available to inform the safe delivery of high-dose boosts to targeted regions. Treatment plans could be delivered rapidly via real-time MR-guided tracking to continuously irradiate the target and safely (precisely) deliver dose distributions with steep spatial gradients. The presented workflow would greatly reduce patient and clinical burden (8 IMPLEMENTATION AND DISSEMINATION).

Illustrated in Figure 2, the GTV represents the extent of the primary tumor revealed by imaging and is outlined manually. The CTV extends the GTV to account for invisible, sub-clinical spread. A predefined GTV-to-CTV margin is typically used, although, depending on the treatment site, the CTV may instead be manually contoured or defined by the entire involved tissue (e.g., the prostate). The purpose of the PTV is to ensure CTV coverage, and it is built by adding margins that account for uncertainties in delineation, patient setup, physiological motion, and treatment delivery.

2.1 Autocontouring

Despite being labor-intensive\textsuperscript{28,29} and frequently resulting in large inter- and intra-observer variations,\textsuperscript{30,31} delineation of target volumes and OAR structures is conventionally performed manually. Machine learning-based, automated contouring (autocontouring) is one popular solution\textsuperscript{32,33} that is currently being introduced in daily practice. It promises greater reproducibility and accuracy of delineated structures, with a substantial reduction in clinical burden. Today, the feasibility of autocontouring treatment targets\textsuperscript{28,34–39} and OARs\textsuperscript{34,40–42} using MRI has been demonstrated and commercial solutions are rapidly being released. However, the success of autocontouring over the coming years will rest on balancing clinical, industrial, and regulatory interests.\textsuperscript{43}

Automated contouring must be robust and flexible for clinical implementation to be feasible. For instance, autocontouring models should be able to rapidly adapt to new imaging protocols, without the need to obtain and annotate a new training set. Future solutions may include generating large sets of synthetic data with the desired contrast for training using generative adversarial networks\textsuperscript{41,44–46}.
2.2 New contouring paradigms

An inherent limitation of RT delineation is that treatment volumes depend on the imaging modality or contrast used. Specifically, GTVs extend only as far as what can be revealed by the imaging and GTV-to-CTV margins must provide a conservative estimate of undetected microscopic expansion, sometimes several cm in magnitude. In the future, advanced application of cutting-edge MRI methods (see 5 Quantitative MRI) may hold the key to safely reducing GTV-CTV margins through improved visualization and/or understanding of the underlying biology. These advances might also allow novel contouring concepts to be implemented clinically, such as probabilistic margin optimisation or even contourless planning. With changes in delineation concepts (including autocontouring), new methods for performing clinical evaluation of contours will also be needed. Geometric measures for evaluating contours (e.g., Dice similarity index) will no longer be clinically relevant. Alternatively, dosimetric comparisons could be made compared with plans generated with ground truth reference contours.

The clinical implementation of new contouring paradigms relies on histopathological validation and large clinical trials so that standards and guidelines may be developed. In transitioning to new contouring paradigms, the MR and RT communities will need to first determine target definitions by asking what should be delineated and why. Second, we must establish whether a delineation task can reliably be achieved. Third, how new contours are used for dose prescription must be addressed. All three aspects would ideally form one consistent and robust clinical strategy.

2.3 Unmet needs

- Development and optimization of autocontouring methods that function robustly with heterogeneous inter-institutional data;
- Clinical QA solutions for safe autocontouring for mid-treatment tumor tracking; and
- A re-examination of how target delineations are defined, in collaboration with the RT community.

3 DOSE CALCULATION

Dose calculation is the computation of energy deposited by ionizing radiation in the patient (i.e., radiation “dose”). Following delineation, treatment planning software is used to simulate the interaction between the patient and
the planned treatment X-ray beams. An iterative process is used to update the beams to optimize the calculated dose in tumor targets and OARs. To model photon scatter and absorption within the patient, information is needed on the tissue elemental composition (e.g., electron density), where this is conventionally derived from dedicated CT imaging. In state-of-the-art RT and in the future, so-called synthetic CT (sCT)—CT-like images derived from MRI—will facilitate MRI-only workflows and adaptive replanning on MR-Linac systems by providing up-to-date sCT maps free from registration errors. In addition, hybrid PET/MR systems give rise to a similar need for attenuation correction, where the culmination of research in both areas advances sCT generation techniques.

Although MRI cannot directly measure X-ray attenuation, many techniques for generating sCT have been proposed in the literature and vendor-provided solutions already exist for brain and pelvis. However, commercial sCT solutions are not available for more complex anatomies, such as the thorax, or tumor sites close to abnormal bony anatomy. Where most vendor solutions are based on bulk density overrides or atlases, recent sCT approaches in research use machine learning architectures, such as generative adversarial networks that may provide solutions for more challenging datasets and anatomic sites. As discussed in the previous section, successful clinical implementation of machine learning-based methods for use in treatment planning will depend on their robustness to clinical variability, such that they meet the quality standards defined by consensus guidelines, and the development of suitable QA phantoms for end-to-end testing.

Today, sCT image volumes are designed to match the resolution and axial orientation of CT scans that are anticipated by treatment planning software. However, planning systems may soon be adapted to more conveniently handle other orientations that are facilitated by MRI. Later, 4D sCT or MRI-based motion signals may inform the simulation software for more advanced treatment planning in moving anatomies.

Looking further ahead, sCT may only be a stepping stone on the way to a new RT paradigm. Future pipelines may not directly reconstruct or display sCTs but, instead, feed k-space data directly to the planning system algorithm to generate a treatment plan using predefined library matching or machine learning approaches. Conversely, reconstruction of sCT images might never leave treatment chains totally since intermediate representations could generate optimal performance. Furthermore, supervision of key intermediate steps is needed for QA purposes and so is likely to remain desirable for years to come.

3.1 | Unmet needs

- 4D-sCT methods suitable for adaptive MRgRT in complex anatomies, such as thoracic sites.

4 | IMAGE GUIDANCE

Image guidance is the process of using imaging at the treatment phase to inform up-to-date localization of tumor targets and healthy tissues. Modern Linacs typically house onboard CBCT to facilitate alignment of the targets to the treatment plan model at the start of each fraction. However, conventional image guidance is limited by poor soft tissue contrast and lack of online motion characterization. Residual targeting errors are generally accounted for by the CTV-to-PTV margins, although large margins limit the dose that can be safely delivered while sparing nearby OARs. Hybrid MR-Linac systems promise to allow reduced PTV sizes through the superior localization and targeting afforded by onboard MRI. Accurate, low-latency motion characterization will facilitate gated treatment, tumor tracking during irradiation, and could enable real-time adaptive replanning. Such precise treatments, delivered to smaller PTVs, will permit safe dose escalation and hypofractionation to improve patient outcomes and clinical efficiency. Furthermore, management of bulk patient motion with real-time MRI could remove the need for uncomfortable immobilization devices.

4.1 | 4D-MRI

In the RT context today, 4D-MRI generally refers to respiratory-correlated 3D-MRI, with image volumes acquired over several breathing cycles and retrospectively binned into respiratory phases. Potential applications of motion characterization using 4D-MRI include onboard treatment plan adaptation and retrospective dose calculations, where 4D-MRI serves as a precursor to volumetric real-time imaging.

In the future, respiratory-correlated 4D-MRI could be replaced by truly time-resolved 4D-MRI (i.e., volumetric real-time imaging), with potential applications in tracking, gating, and real-time dose monitoring. Current developments include using motion models built from prospectively acquired 4D-MRI to rapidly generate synthetic 4D-MRI updated by 2D imaging of the motion perpendicular to the treatment beam. Alternatively, the use of higher-order surrogate signals can resolve signal characteristics beyond respiratory motion, enabling simultaneous resolution of peristaltic motion or cardiac
motion to aid cardiac gating for MRgRT of ventricular tachycardia.93

4.2 | Real-time MRI

To fully realize the potential of motion management for MR-guided adaptive targeting, low-latency, high-fidelity data for precise spatial–temporal localization is desirable. However, low-latency goal differs for each motion type. For instance, cardiac motion is on the sub-second scale, whereas organ filling extends over minutes.76 Recommendations have been made for MRI latencies of 200-500 ms for respiratory motion,94 although how fast this could be and still make a clinical impact is an open question that must be revisited as research progresses. Currently, when mid treatment adaptation is desired, time-resolved 2D-MRI images are often obtained, rather than 4D-MRI. Real-time adaptive image processing for MRI is an area of ongoing research.95 To minimize latency, the amount of acquired data per frame must be reduced. Potential solutions include the use of accurate spatiotemporal motion models,96,97 suitable low-rank subspace constraints,98 or sparsely sampled k-space data interpreted by compressed sensing.99 However, for most of these accelerated MRI acquisition techniques, the gain in acquisition time results in longer reconstruction times. Fortunately, machine learning approaches that transfer computational processing to offline training of a neural network99–102 may overcome long reconstruction times of accelerated acquisitions. In the future, latency for treatment planning and image guidance could be further reduced through use of patient representations composited from models that extract various representative states and their probabilistic variations. Tighter integration may gradually lift the need to exchange information between MRI scanners and Linacs in the form of images, opening opportunities for reducing latency through direct contour tracking from raw MRI data.58,103

Opportunities and challenges of real-time imaging for MRgRT are shared by interventional MRI.104 We should, therefore, ensure that these fields learn from one another as solutions are explored in the coming years. In addition, MR-guided proton therapy will benefit from advances in MRgRT, where a full characterization of target and OAR motion is crucial because steep dose gradients exist not only perpendicular to the beam, but also along the direction of the beam.105

4.3 | Unmet needs

• Rapid online reconstruction of highly undersampled MRI data; and
• A tighter integration of MRI and RT systems for adaptive planning informed directly by k-space data.

5 | QUANTITATIVE MRI

Biomarkers derived from qMRI techniques allow for non-invasive assessment of morphological, biological, and functional processes in tissue and so promise several key roles throughout RT workflows.106 First, qMRI could improve visualization for delineation by incorporating advanced contrast mechanisms. Second, qMRI biomarkers promise to provide metrics for RT response to allow adaptive treatment based on physiological responses9 (e.g., necrosis) that manifest earlier than anatomical imaging features.21,107 For instance, changes in cell density—a well-established marker for early response detection—can be measured indirectly using DWI for early response detection.9 Third, qMRI techniques may offer a surrogate for tissue dose sensitivity, such that treatment dose boosts can be informed and adapted according to baseline measurements.108 Several recent articles have been published on the use and level of evidence for different qMRI techniques in RT.103,106,109 We particularly refer to Table 1 in van Houdt et al.103

5.1 | MRI-derived biomarkers

An active area of MRI research that works to detect radiation sensitivity is the investigation of hypoxia, a well-established and important prognostic marker for radioresistance. Hypoxic tissues require up to 3-fold greater doses to achieve the same biological effectiveness.110 Although there are several MRI approaches for assessing oxygenation (pO$_2$), they are predominately indirect. Tissue water T$_1$ is sensitive to pO$_2$ because the oxygen molecule is paramagnetic. The effect is small, but recent pre-clinical work demonstrated the feasibility of stratifying tumors based on pre-irradiation oxygen gas breathing to predict long term tumor control following radiation.108 Meanwhile, T$_2^*$ is strongly influenced by the concentration of deoxyhemoglobin. Perfusion is also an indirect marker for hypoxia, which can be measured using DCE, arterial spin labelling, or intravoxel incoherent motion. A more direct way of measuring pO$_2$ is with dynamic oxygen-17 MRI111; however, this technique is expensive and suffers from weak SNR, so has not been commonly investigated. Other commonly investigated qMRI techniques for RT include CEST and MR spectroscopy.103 Recent analysis suggests radiation dose could be effectively adapted using a genomic-adjusted radiation dose model112 and...
active investigations seek similar capabilities based on radiomics. Ultimately, we believe a combination of several techniques will allow complimentary information to be sampled on the state of the tumor. These data will allow clinicians to generate better personalized treatment plans than ever before, targeting dose to (hypoxic) radioresistant tumor regions and reducing dose to regions it is no longer needed.

We expect that the impact of qMRI development for RT will not only improve RT outcomes, but allow RT in cases that are currently considered unsuitable. For instance, lung cancer patients with severe lung function loss are often limited to surgery because of the risk of damage to remaining healthy tissue. However, with qMRI in combination with ventilation of hyperpolarized gases, functional regions of the lung can be clearly identified and considered, enabling RT as viable treatment in these patients.

### 5.2 From research tool to clinical tool

Currently, qMRI for RT is predominantly a research tool, with most work focusing on establishing a link between MRI and treatment response. To translate qMRI to clinical use, the next step will be establishing quantitative imaging biomarkers (QIBs) from qMRI parameters. The general imaging biomarker roadmap of O’Connor et al. provides a useful framework for these next steps, where there must be a transition from a promising QIB, to a potential QIB, and ultimately toward a clinically validated QIB.

Today, evidence for QIBs in RT is limited. Complex logistics and the added patient burden of extra MRI examinations mean that analyses are often based on small patient cohorts or very few time points. To overcome these difficulties, functional imaging data for QIB studies could be collected on MRgRT systems at the time of treatment. Through systematic measurement of qMRI across treatment courses, large collaborative libraries could be built to detect which qMRI techniques generate truly prognostic QIBs. Such an initiative would require large, collaborative networks that include experts from both MRI and RT communities, such as the Elekta MR-Linac consortium, to collect data prospectively and systematically over many years. To supplement this, robust data-science frameworks should be established, which are often overlooked in qMRI studies.

When the prognostic value of a set of qMRI parameters has been systematically demonstrated in a large cohort, the next step of clinical validation is confirmation that the qMRI method also has predictive value (i.e., can be used to modify treatment). Investigations into predictive value can be conducted using interventional trials that adhere to the RT idea, development, exploration, assessment, and long-term evaluation framework. Although we must first ensure that any unknowns are first solved, like how qMRI parameter maps are translated into the dose prescription.

To systematically study the relation between qMRI parameters, dose, and treatment response requires comparing qMRI with clinical outcome measures at different treatment dose levels. Some insight can be gained by comparing results between periods where guidelines for dose prescriptions changed or countries that prescribe differing treatment doses. Ultimately, however, qMRI validation requires randomized trials with variable dose. Setting up such trials is challenging because current dose levels are the accepted clinical standard. Changing doses could benefit some patients but could result in a worse outcome for others. Therefore, informed, careful patient selection, and close collaboration between qMRI experts and oncologists will be essential. In particular, radiation oncologists should have a more advanced understanding of the underlying qMRI mechanisms so that they can be comfortable in adapting treatment.

Initial efforts toward consensus guidelines for qMRI on MRgRT systems have recently begun. However, the current focus of qMRI in RT is on the target volume, where QIBs that monitor normal tissue toxicity could be further explored. To further develop guidelines for RT QIBs, there are opportunities to learn from and work with the diagnostic qMRI community, building on pre-existing work. Such opportunities include initiatives for accurate and reproducible qMRI, learning from the Quantitative Imaging Network, and guidelines from the Quantitative Imaging Biomarker Alliance (QIBA) on DWI and DCE-MRI. In adapting diagnostic recommendations for RT, we must remember that MR-Linacs differ from conventional MRI systems. For instance, images from MRgRT systems typically exhibit a lower SNR than those obtained using diagnostic devices and sometimes have unconventional field strengths.

### 5.3 Adaptive treatment

One major opportunity for qMRI that arose with the onset of MR-Linac systems is daily tumor biology-based treatments. For instance, the availability of real-time qMRI techniques could improve RT efficacy by allowing treatment to be timed to when the tumor is at its most sensitive to irradiation, such as outside of hypoxic periods. MRI might also be used to directly enhance treatment. For instance, radiation sensitivity could be increased with drugs targeted to the tumor tissue with MRI, using a similar approach as MR targeting. Alternatively, hypoxia could be reduced by breathing hyperoxic or hyperbaric oxygen, with qMRI used to confirm normoxic status.
Another application of qMRI in RT could be real-time visualization of biological dose. Because radiation dosimetry can be assessed in vitro using Bang gels, one can envisage extension to in vivo applications. Through a deeper understanding of the short-term effects of dose on tissue, we may find an MRI contrast mechanism, such as CEST or DWI, is sensitive to the short-term biological effect of the treatment beam. Such qMRI methods could be used for validation and adaptation of the planned treatment.

5.4 Protocol optimization

The image quality of qMRI is notoriously poor when compared to conventional MRI. Because multiple images must be obtained to model and measure signal changes, image resolutions are low despite long acquisition times. Therefore, clinicians often prefer conventional MR images for tumor assessments. Technical improvements in acquisition speed and image quality will greatly aid implementation of qMRI in clinical workflows.

For state-of-the-art RT on MR-Linacs, faster qMRI is imperative. Today, qMRI measurements are acquired during the opportunity-time created by manual contouring. With the clinical adoption of autocontouring (see 2 Delineation), the time available for qMRI measurements for MRgRT will be shortened. Methods such as MR fingerprinting, model-based image reconstruction, and MR-spin tomography in time-domain could enable substantially shorter acquisitions and could yield higher resolution qMRI images with improved accuracies. However, shorter acquisition times often come with a trade-off of longer reconstruction times. For online applications, the solution may be machine learning-based methods for rapid reconstruction and modeling. Ultimately, we may measure QIBs in tumors directly from undersampled raw k-space data to meet the goal of real-time monitoring and treatment adaptation.

5.5 Unmet needs

- Established, standardized QIBs for RT derived from qMRI parameters;
- Demonstration of the predictive value of QIBs across large multi-center cohorts;
- Accelerated pipelines for acquisition, reconstruction, and interpretation of qMRI; and
- Improved image quality of qMRI parameter maps.

6 HARDWARE

6.1 MRI for RT

Standalone MRI systems may be used for simulation imaging (i.e., MR-Sim). Compared to conventional diagnostic MRI scanners, these pre-treatment imaging systems must meet additional RT-specific requirements. For instance, high spatial accuracy is important since geometrical image distortions can lead to under-exposure of the tumor site and unnecessary dose to healthy structures. Geometric fidelity depends on magnetic field homogeneity and gradient linearity, which are typically worse at higher field strengths and can be compromised by the integration of the Linac system. Unlike for diagnostic MRI, the geometric fidelity of MR images is critically important in RT applications. The implementation of MRI for RT has, therefore, largely focused on minimizing and characterizing distortions as new techniques and QA procedures were developed. Today, this issue is largely solved, but will remain an important factor to consider as the technology develops.

The installation of conventional MRI systems in RT departments can be complex and costly. Large scanner weights, the need to incorporate a quench pipe in shielding designs, and the undesirable interaction between MRI fringe fields and nearby Linacs are often challenging factors, and the need for MR-Safe immobilization devices and other devices (e.g., intravenous-contrast pumps) further adds to the cost. In addition, wide scanner bores are required to accommodate immobilization equipment.

Several recent developments can help adapt diagnostic systems for RT purposes. The industry has recently developed diagnostic MR scanners with low helium content (e.g., <8 L) that do not require a quench pipe, allowing for lower installation costs and a reduced environmental impact. Low-field scanners are another example of these developments, which can improve geometric accuracy for RT, but this trade-off must be considered with SNR and image quality losses below 1 T (particularly for qMRI). However, in the future, low-field SNR may be significantly boosted through machine learning driven reconstruction.

6.2 MR-Linac systems

Hybrid MR-Linac systems present a different set of engineering challenges. For instance, the influence of the MR system fringe field on the Linac must be minimized. In addition, the RF radiation originating from the Linac must be shielded from the MRI sub-system,
which must, in turn, be carefully designed to meet radiation attenuation requirements.

Currently, two MR-Linac solutions are commercially available, which have each taken different approaches to the integration of an MR scanner with RT beam-generation components. Both MR sub-systems are based on diagnostic designs, which have been modified to meet RT workflow and dosimetric requirements while maintaining imaging performance (e.g., spatial integrity). The Unity (Elekta AB) MR device is based on a modified 1.5 T MRI (Philips Healthcare). The magnet is optimized to create a surrounding annulus of a low magnetic field to enable its decoupling from the rotating-gantry-mounted ferromagnetic components that include the beam generation sub-systems. The magnet was also modified to create a radiation window by splitting the gradient coils. The MRIdian (ViewRay Inc) system houses a superconducting, 0.35-T, split-magnet design, using ferromagnetic shielding to isolate the Linac sub-systems on the ring gantry from the magnet. In both vendor designs, the gap between the two magnet cryostat components permits megavoltage X-ray beams to pass through with very little attenuation. Non-commercially available MR-Linac systems have focused on bi-planar rotatable MR designs and the use of a standalone magnet with a non-rotatable radiation beam.

To date, ~200 MR-Linacs have been installed, which is limited compared to the global installed base of roughly 13,000 conventional Linacs. To provide improved access to MR-Linacs, it is important that they become cheaper and simpler to use in the future. There are several challenges associated with the current designs. First, the MR magnet structure offers limited access to the patient table inside the bore. Second, the overall size, weight, and cost of the MR scanner adds complexity. The footprint of MR-Linac systems may pose significant demands on the construction space required, greatly increasing installation costs. Third, only a very limited range of coils, with low numbers of coil elements, are available. Fourth, state-of-the-art treatments, like volumetric-modulated arc therapy, are not yet available for MR-Linac systems.

As the MR-Linac market grows, optimized components may start to differ from the mainstream diagnostic solutions to become more aligned with the unique needs of RT. Future iterations of MR-Linac technology may include greater use of modeling for the MR magnet optimization problem, such that additional Linac structural and performance specifications are considered. MR-Linac designs could also put more focus on requirements for maximizing patient access and minimizing hardware size. Vendors should facilitate easy MR-Linac upgrades because these are essential to enabling rapid integration of novel treatment and imaging innovations.

Conversely, maintaining a similar blueprint could reduce overheads through the sharing of manufacturing, obsolescence, and supply costs. This could be aided further by focusing on open-source hardware. Comparable designs could facilitate fast and easy translation of MRI solutions to the MR-Linac domain. Another advantage is that when MR-designs are similar, less retraining is required for in-house radiology experts.

In addition to imaging performance, RF coil design for MRgRT applications must balance patient setup and dosimetric requirements. For instance, for some MR-Linac designs, the RF coil elevates above the patient to reduce the excess surface dose at the expense of SNR. Future designs could instead reduce excess surface (skin) dose by constructing RF coils with inbuilt foam boluses, to allow the coils to be positioned closer to the patient for improved SNR. Low-weight coils minimize body-contour deformations and ease patient setup. Another important design consideration relates to the local beam attenuation and positioning of sensitive electronics, which limit the number of coil elements and, consequently, the parallel imaging capabilities of the MR subsystem. RF coils using high impedance capacitors could enable a high number of coil elements to be used while meeting the beam attenuation requirements. Alternatively, high-density, disposable coils that allow electronics to be directly in the path of the radiation beam might be considered. Overall, many new coil designs could yet be exploited to optimize image quality for MRgRT, including wireless, flexible and disposable RF coils, as well as inbuilt bolus designs.

### 6.3 | Unmet needs

- Easier access to MR-Sim and MR-Linac systems: simpler installation, reduced costs, and reduced footprint; and
- Optimized MR-Linac components that are more aligned with the unique needs of RT e.g., improved coil designs.

### 7 | REDUCING PATIENT BURDEN

An important aspect for the success of MR in RT is minimizing patients’ treatment burden. In addition to well-being, patient burden includes the time, difficulty, and costs devoted to healthcare. Critical components of the treatment burden are the number of visits to the hospital (including travel time and costs), the duration and comfort of the treatment position.
7.1 | MR guided radiotherapy

The advent of MRgRT systems has initially increased treatment burden for patients since treatment session durations are substantially longer on MR-Linacs than conventional Linacs. The average treatment time for prostate cancer has increased from 15 to 20 min on CBCT-Linacs to 45 min on MR-Linacs. Liver treatment on the MR-Linac is particularly long, ranging from 60 to 90 min. Although 45 min is generally acceptable on diagnostic MRI scanners, MRgRT patients are set up in the treatment position, which can include a hard, flat tabletop, fixation devices (e.g., closely fitted full-head masks), and uncomfortable positions (e.g., arms above the head). Furthermore, patients treated on MR-Linacs experience increased MRI-related acoustic noise and anxiety because of limited space. With the many repeated MRgRT sessions throughout an RT course, acoustic noise has a more substantial impact on hearing than for a one-off diagnostic MRI examination.

In the future, MRgRT on MR-Linac systems presents several opportunities to reduce patient discomfort. First, adaptive planning using onboard MR imaging could allow for more comfortable treatment couches, where a hard, flat tabletop is no longer needed for consistent setup. Second, future MR-Linac models could be developed with wider bores to reduce claustrophobia and aid access to the patient. Third, uncomfortable setup devices may be rendered unnecessary with online MRI tracking. Fourth, future developments in MRI, tumor tracking, and gated deliveries could remove the need for breath-hold imaging and treatment deliveries.

7.2 | Hypofractionation

Hypofractionation, increasing the dose per fraction and reducing the number of fractions, allows a biologically similar treatment plan to be delivered in fewer hospital visits. However, a major challenge of hypofractionated approaches is that treatment becomes more sensitive to patient setup errors. MR-Linac systems promise to make setup errors smaller and overcome these limitations. Consequently, clinicians are currently attempting to increase the dose per fraction in several MRgRT protocols. By further improving image quality at planning and real-time monitoring during treatment, we could further reduce uncertainties and gain confidence in continuing the reduction of fraction numbers. However, it should be noted that spreading treatment over multiple fractions can allow for the repair of healthy tissues between sessions, reducing potential toxicity. Furthermore, while hypofractionated treatment regimens have shown dramatic improvements to treatment response for many disease sites, these results are only possible with excellent geometric precision and, in some instances, tumor dose spread must be more heterogeneous to ensure normal issues are preserved.

Next-generation RT workflows could be “one-stop-shop”, only requiring 1 to 3 patient hospital visits. In these workflows, once the patient is set up on the MR-Linac, fast MRI scans are acquired, target volumes and OARs are automatically contoured and, within seconds, an automated, single-fraction, high-dose treatment plan is developed and delivered, all during a single visit (Figure 1). Such an approach would increase clinical throughput and improve patient experience, especially for palliative patients. However, at present, “day-one” treatment planning would require substantial clinical resources and automation is strongly desired.

7.3 | Unmet needs

- Increased patient comfort via removal of uncomfortable elements from MRgRT treatment chains, such as hard tabletops, immobilization devices, and breath-holding;
- Faster imaging and treatment to reduce time in the machine; and
- Improved image quality at planning and real-time monitoring during treatment to improve confidence in hypofractionation.

8 | IMPLEMENTATION AND DISSEMINATION

The implementation of MRI in RT will be accelerated and steered by the introduction of MRgRT on MR-Linacs. Currently, however, only a few RT centers have MRI scanners installed in the RT department and MR-Linacs only account for a small fraction (∼1.5%) of all treatment machines in clinical use. We expect that developments will initially take different directions for non-academic and academic centers. In non-academic centers with MR-Sim only, the focus will be on targeting accuracy. For non-academic centers with an MR-Linac, this will be combined with fast and automated hypofractionated RT and target tracking, where hypofractionation greatly reduces the cost of RT. In academic centers, experiments will focus heavily on development for qMRI methods. As the use of MRI for RT increases, guidelines for its practical implementation should be reviewed and updated.
8.1 | Clinical burden

Operating an MR-Linac currently requires a large team of clinical and technical experts. Centers with MR-Linacs often require an on-site clinician for recontouring, two dual-trained RT-MR technologists to drive treatment, an on-call MR-RT physicist, and a large group of physicists available for quality control and maintenance. Even where the additional facilities and expertise required were minimal, the increased strain on staff resources caused by using an MR-Linac is often significant, with treatment times typically doubling those of conventional systems. The cost of developing and maintaining new support teams for MR-Linac treatments is manageable for large cancer therapy centers, but could be prohibitive for smaller (2-3 Linacs), community-based radiation therapy centers, which are typical across Europe and the United States. We, therefore, predict that over the next few years, MRgRT will predominantly be conducted at larger specialized centers.

For the dissemination of MRgRT and MR-Sim to non-academic centers and for long-term usage in academic centers to be successful, logistic, environmental, and staff burdens must be reduced. Efforts should be made to reduce treatment times and simplify the operation of the MR-Linac. Simple solutions to reducing clinical burden include transferring staff training to external parties and investing in AI-assisted workflows. Standardized MRI acquisitions for treatment planning and motion monitoring combined with AI-driven MRI-scanning methods reduce the complexity of MR knowledge needed by radiographers. Similarly, AI-driven contouring and treatment planning will greatly reduce the time and staff requirements for on-table plan adaptation of treatments. Looking further ahead, the operational burden on the physics staff could be reduced by increased automation of patient treatment and QA procedures and highly hypofractionated treatment courses. Such a workflow would resemble the ultimate minimization in operational burden and drive the uptake of MRgRT across all radiation oncology centers. It is of vital importance that clinicians are included in the introduction of these new approaches since automated workflows remove clinical decision making and hypofractionated treatments deviate from the current clinical standard.

8.2 | Unmet needs

- Standardization of MRI acquisitions for treatment planning and motion monitoring;
- Collaborative development of automated workflows by researchers and clinical teams; and
- Easy (or automated) operation of MRI and MRgRT systems.

9 | CONCLUDING REMARKS

MRI has become indispensable in modern RT pipelines and its role is expected to grow. Advances in tumor delineation, onboard image guidance, and imaging biomarkers afforded by MRI promise to transform RT over the next 25 years. With this paradigm shift, a rich spectrum of new challenges and opportunities is presented for the MR community.

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Magnetic Resonance in Medicine

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