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Review paper

MRI-guided lung SBRT: Present and future developments



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ABSTRACT

Stereotactic body radiotherapy (SBRT) is rapidly becoming an alternative to surgery for the treatment of early-stage non-small cell lung cancer patients. Lung SBRT is administered in a hypo-fractionated, conformal manner, delivering high doses to the target. To avoid normal-tissue toxicity, it is crucial to limit the exposure of nearby healthy organs-at-risk (OAR).

Current image-guided radiotherapy strategies for lung SBRT are mostly based on X-ray imaging modalities. Although still in its infancy, magnetic resonance imaging (MRI) guidance for lung SBRT is not exposure-limited and MRI promises to improve crucial soft-tissue contrast. Looking beyond anatomical imaging, functional MRI is expected to inform treatment decisions and adaptations in the future.

This review summarises and discusses how MRI could be advantageous to the different links of the radiotherapy treatment chain for lung SBRT: diagnosis and staging, tumour and OAR delineation, treatment planning, and inter- or intrafractional motion management. Special emphasis is placed on a new generation of hybrid MRI treatment devices and their potential for real-time adaptive radiotherapy.

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Contents

1.	Introduction			
	1.1.	The clinical rationale for lung SBRT	139	
	1.2.	Current clinical workflow for lung SBRT	140	
		MRI-guided lung SBRT		
2.	Techn	nical implementations of MRI scanners into the radiotherapy workflow	140	
	2.1.	Offline MRI.	140	
	2.2.	Online MRI.	141	
3.	Oppor	rtunities for MRI to enhance the lung SBRT workflow	142	
	3.1.	Diagnosis and staging	142	
	3.2.	Delineation, dose calculation and treatment planning	143	
	3.3.	Patient setup and interfractional adaptations	144	
	3.4.	Intrafractional motion management	144	
4.		usion and outlook		
		owledgements		
	ences	146		

1. Introduction

1.1. The clinical rationale for lung SBRT

Lung cancer is the most common type of cancer, with non-small cell lung cancer (NSCLC) representing the majority of cases [1,2].

Within this patient cohort, roughly 15–20% present with an early and localised form of the disease (stage I), indicating a lack of nodal involvement and metastatic spread. The recent roll-out of widescale, low-dose computed tomography (CT)-assisted screening might further increase the number of patients diagnosed at an early disease stage. Traditionally, surgery is the treatment method of choice for these patients [3]. However, some patients are classified as *medically inoperable* due to comorbidities, or they decline surgery for other reasons.

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In the past, the alternative treatment option of prescribing conventionally fractioned radiotherapy resulted in worse outcome compared to surgery, mainly due to the lack of local control [4]. Taking heed from cranial radiosurgery, stereotactic body radiotherapy (SBRT) aims at delivering fewer, but more intense radiotherapy fractions (hypo-fractionation). Several studies have shown that increasing the biologically effective dose above 100 Gy decreases the number of patients with local failures and increases overall survival in early-stage NSCLC patients compared to conventionally fractionated radiotherapy [5], leading to SBRT being the new standard of care for inoperable early-stage NSCLC patients.

Traditionally, only patients with tumours located at least 2 cm away from the proximal bronchial tree are considered for SBRT [6]. This pre-selection of eligible patients intends to minimise bronchial toxicities, but it also severely limits treatment options for inoperable early-stage NSCLC patients with more centrally located tumours [7]. More recently, trials were set up to identify a slightly milder form of hypo-fractionation suitable for these patients [8]. Furthermore, SBRT is being investigated for the treatment of oligometastatic lung cancer [9,10] and small cell lung cancer [11].

1.2. Current clinical workflow for lung SBRT

While SBRT aims at delivering high doses to the target, it is crucial to limit the exposure of nearby healthy organs-at-risk (OAR), including surrounding lung tissue, the brachial plexus, spinal cord, trachea, proximal bronchial tree, oesophagus, heart and great aortic vessels, liver and ribs [12]. An additional challenge for the radiotherapy of lung tumours are inter- and intrafractional anatomical changes [13].

After diagnosis, staging and treatment decision, the clinical radiotherapy workflow begins with the acquisition of a planning CT scan. Accurate delineation of the tumour and nearby OAR in this image is crucial, as a treatment plan is created based on these contours. Today, many institutions utilise modern delivery techniques for SBRT, such as intensity-modulated radiotherapy and volumetric-modulated arc radiotherapy [14–16]. In conjunction with inverse treatment plan optimization it is possible to design highly conformal dose distributions, even in inhomogeneous anatomies [17,18].

In order to ensure correct delivery of the treatment plan, utmost attention is paid to precise patient positioning for each fraction, for example by using body frames for fixation [19]. About 10–15 years ago, volumetric X-ray imaging started being integrated with modern linear accelerators (linacs) [20–22]. Utilising the provided imaging information, the patient can be rigidly shifted to position the target inside the treatment beam [23,24]. Implanted fiducials often support this process. In addition to rigid shifts, the patient's body may deform between fractions due to weight loss, tumour growth or shrinkage, or volume changes of healthy organs. The volumetric images acquired prior to each fraction can be used to evaluate whether the original treatment plan is still suitable after interfractional motion. If not, a substitute plan can be created, either based on a newly acquired simulation CT scan or using the in-room images themselves [25,26].

Intrafractional motion during treatment delivery, such as respiratory and cardiac movement, may cause the anatomy to shift or deform up to several centimetres [13]. Using a 4D CT scan for treatment planning allows the consideration of breathing motion by deploying either an internal target volume (ITV), mid-ventilation or mid-position approach [27–29]. However, all these methods are based on extended treatment margins and the characteristics of the target motion can change over the course of the treatment [30]. Breathing protocols attempt to reproduce a certain anatomical state during treatment delivery by having the patient hold its

breath, either voluntarily [31] or aided by an active breathing control device [32]. Gating turns off the treatment beam if the target leaves a previously defined volume [33]. Both these strategies result in an increased treatment time and reduced patient comfort. More advanced techniques aim at adapting the treatment in realtime by either moving the entire linac [34], tilting the treatment head [35], repositioning the patient using a robotic treatment couch [36,37], or by changing the treatment beam's position and shape by moving the leaves of the linac's multi-leaf collimator (MLC) [38–40]. Most of these techniques have been used to deliver real-time adapted lung SBRT [41-43]. They all require real-time information about the patient's anatomy. This can be obtained by monitoring either internal or external surrogates and correlating them with the tumour motion [44]. The tumour can also be localised using MV portal imaging [45,46] or kV fluoroscopy [47,48]. Automated localisation is often aided by fiducial markers, surgically inserted in or near the target. Another marker-based method detects implanted resonant circuits using an electromagnetic array

1.3. MRI-guided lung SBRT

Despite these advances in lung SBRT delivery technique, there are still residual uncertainties. MRI offers better soft-tissue contrast than CT or cone-beam CT (CBCT), does not expose the patient to additional imaging dose, and offers a larger variety of functional imaging capabilities. For this reason, there has been an increased interest in harnessing MRI for radiotherapy of different cancer sites, including lung. This has the potential to enhance several aspects of the lung SBRT workflow, including:

- diagnosis and staging (Section 3.1),
- delineation of the tumour and OAR for treatment planning (Section 3.2),
- accurate patient setup and adaptation of the treatment to interfractional anatomy changes (Section 3.3),
- monitoring of intrafractional motion and real-time adaptation of the delivery (Section 3.4).

Decreasing the uncertainties in these areas allows shrinking the treatment margins while maintaining dose coverage of the target. Consequently, reduced toxicities in normal-tissue for lung SBRT, and, ultimately, the expansion of the SBRT to more central lung tumours can be surmised.

This review first introduces the various technical implementations of MRI scanners into the radiotherapy workflow (Section 2). Afterwards, it discusses the different areas where MRI is already enhancing lung SBRT treatments or might do so in the future (Section 3). It is important to keep in mind that MRI-guided lung SBRT is still in its infancy and that the number of directly related publications is therefore limited.

2. Technical implementations of MRI scanners into the radiotherapy workflow

2.1. Offline MRI

To employ MRI for diagnostic, staging and treatment simulation purposes, having a stand-alone MRI scanner at the same institution can suffice. However, when using MRI for patient setup and interfractional treatment adaptation, identical patient positioning during imaging and treatment becomes crucial. One way to ensure this relies on the transport of the patient between linac and MRI scanner, located in different rooms, on a trolley system using a shared table top and immobilisation device (see Fig. 1) [50,51].

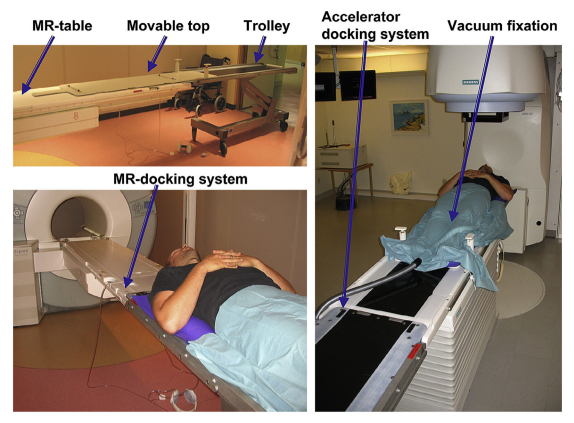


Fig. 1. Three photographs of the shuttle-based MRI-guided radiotherapy solution at Umeå University, Sweden. A movable treatment table and patient fixation system can be attached to either imaging or treatment device. Reprinted from International Journal of Radiation Oncology*Biology*Physics, 74(2), M. Karlsson, M.G. Karlsson, T. Nyholm, C. Amies, B. Zackrisson, Dedicated Magnetic Resonance Imaging in the Radiotherapy Clinic, pp. 644–51, Copyright (2009), with permission from Elsevier.

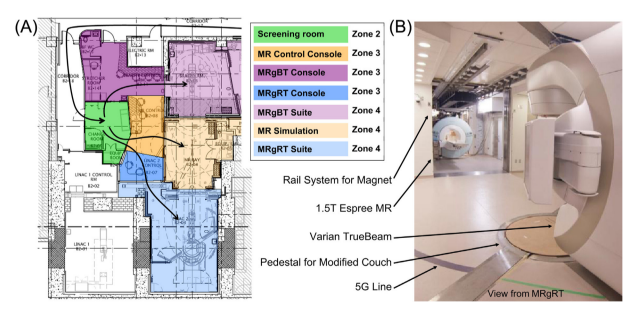
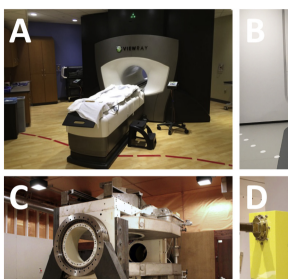


Fig. 2. (A) Floor plan of the MRI-guided facility at Princess Margaret Hospital, Toronto, Canada. (B) The MRI-scanner is attached to rails on the ceiling and can be moved either into the brachytherapy or radiotherapy suite prior to treatment. Reprinted from Seminars in Radiation Oncology, 24(3), D.A. Jaffray, M.C. Carlone, M.F. Milosevic, S.L. Breen, T. Stanescu, A. Rink, H. Alasti, A. Simeonov, M.C. Sweitzer, J.D. Winter, A Facility for Magnetic Resonance–Guided Radiation Therapy, pp. 193–95, Copyright (2014), with permission from Elsevier.

Another option is to shuttle the MRI scanner on ceiling-mounted rails into and out of the treatment room (see Fig. 2) [52]. Both system designs ensure that the magnitude of electromagnetic cross-interference is reduced to a minimum by using spatial separation.

2.2. Online MRI

Offline MRI solutions cannot monitor intrafractional anatomical changes and the patient might move between imaging and



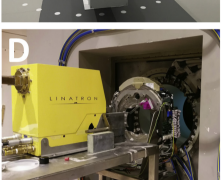


Fig. 3. MRI-guided treatment units developed by different research groups/vendors. (A) The ViewRay MRIdian, the first clinical system at Washington University, St. Louis, MO, USA (image courtesy of P.J. Parikh), (B) the 1.5 T MR-linac developed by Elekta AB installed at NKI-AvL, Amsterdam, The Netherlands, (C) the linac-MR prototype (v1) at Cross Cancer Institute, Edmonton, Canada (image courtesy of B.G. Fallone, www.linacMR.ca), and (D) the MRI-linac prototype developed by the Australian MR-linac program, located at the Ingham Institute, Liverpool, Australia (image courtesy of G.P. Liney on behalf of the Australian MRI-Linac Program).

treatment. For this reason, there are considerable efforts to integrate MRI directly with the treatment unit.

The first commercially available MRI-guided treatment system, based on irradiation with three ⁶⁰Co sources, has been developed by ViewRay Inc., Cleveland, OH, USA [53]. The system has been used to treat patients since early 2014, including lung cancer patients with SBRT [54]. Several research groups/vendors have been working on combining a linac with MRI (see Fig. 3) [55–58].

Machine design has to warrant correct functioning of both imaging and treatment system in the presence of electromagnetic cross-interference. The choice of beam source and main magnet as well as the orientation and movement of all the components relative to each other directly influences the dose delivery and imaging capabilities. There are considerable differences between systems with regard to treatment beam properties, such as beam energy, penumbra, field-of-view and MLC leaf width at isocenter, and the strength and orientation of the main magnetic field (see Table 1). In addition to beam properties, the dose delivered to the patient is affected by the B₀ magnetic field, which is present at all times. While the primary photon beam is not influenced, the Lorentz force deflects secondary electrons onto a curved trajectory. This electron return effect (ERE) is more pronounced at higher B₀ field strengths and distorts the dose especially at air-tissue-interfaces, which present themselves manifold in thoracic anatomy [59]. Signal- and contrast-to-noise ratio as well as spatial fidelity of the images are affected by the strength and homogeneity of the

B0 magnetic field as these govern the relaxation times of the MRI signal. Due to lung tissue's inherently low T2* relaxation time, lower field strengths can be beneficial when attempting to image thoracic anatomies [60].

3. Opportunities for MRI to enhance the lung SBRT workflow

Table 2 provides an overview of the most commonly employed sequences for lung MRI. The table is condensed from the recommendations in [63,64] and annotated with radiotherapy application in mind. Fast spin echo (FSE) sequences are typically acquired using respiratory gating, which is prone to artefacts caused by irregular breathing or cardiac motion. Balanced steady-state free precession (bSSFP) sequences exhibit a T2/T1-weighted contrast and a high signal-to-noise ratio, but can suffer from banding artefacts in peripheral areas of the image, which is problematic if body contours are needed for treatment planning. The traditional limitations of diffusion-weighted (DW) MRI could be overcome by moving away from the echoplanar imaging readout to a segmented readout scheme [65].

3.1. Diagnosis and staging

In order to be eligible for lung SBRT, patients must present at an early disease stage, meaning that only small, localised lesions are

Table 1 System characteristics of the most common hybrid MRI radiotherapy machines. B_0 orientation is specified relative to the treatment beam direction.

Treatment unit Treatme	ent beam(s) Magnet design	$B_0[T]$	B ₀ orientation	Additional comments
Elekta MR-linac [55] 7 MV li Canadian linac-MR (v2) [61] 6 or 10	OCO sources Split nac Closed MV linac Split MV linac Split	0.35 1.5 0.5 1.0*	Perpendicular Perpendicular Inline Inline or perpendicular	Replacement of 60 Co sources with a linac under development Beam traverses cryostat and magnet Magnet rotates with gantry Both B_0 orientations under investigation

^{*} The current research version uses a 1.5 T closed magnet [62].

Table 2MRI sequences for lung cancer imaging (adapted from [63,64]).

Sequence	Strength	Acquisition type	Spatial resolution	Distortion risk	Variants	Comments
Volumetric T1w GRE	Nodules	Breath-hold	High	Low	Dixon, 4D	Low contrast
T2w single-shot FSE	Infiltrative disease	Breath-hold	Low – moderate	Low	Cine	-
T2w gated FSE	Mediastinal structures, infiltrative disease	Respiratory gated	Moderate – high	Moderate	Dixon, volumetric	Long acquisitions
T2/T1w bSSFP	Lung function	Free-breathing	Moderate – high	Moderate	Cine, 4D	Banding artefacts
T1w FSE	Mediastinal lymph nodes	Multiple breath- hold	Moderate	Moderate	DCE	_
DWI	Dose painting	Multiple breath- hold	Low	High	-	Distortion risk reduction possible

Abbreviations: GRE = gradient echo; DCE = dynamic contrast-enhanced; FSE = fast spin echo; T1w/ T2w = T1/ T2-weighted; bSSFP = balanced steady-state free precession; DWI = diffusion-weighted imaging.

existant without any nodal or metastatic spread. Currently, combined multi-slice CT and ¹⁸Fludeoxyglucose (FDG) positron emission tomography (PET), in addition to biopsies, are considered the gold standard for staging [66]. However, biopsy procedures pose an additional burden to the patient and their execution is challenging for small, peripheral tumours [3]. Although the use of PET-CT is well established, the relatively low spatial resolution and motion artefacts, resulting from long acquisition times, are prevalent issues [67,68]. Furthermore, inflammatory areas in the lung can lead to increased FDG uptake and the diagnosis of false positives. MRI could play a role in staging [69,70] and MRI-derived motion information in hybrid PET-MRI devices could be used to mitigate PET motion artefacts [71,72].

In order to determine size and extent of tumour growth with MRI, a range of acquisition techniques has been deployed, for example standard T1- or T2-weighted acquisitions, 2D cine MRI, or single-shot FSE MRI [73,74]. T2-weighted, dynamic contrastenhanced (DCE), or DW MRI have proven promising in their ability to distinguish between benign and malignant tumours [75–78].

Identifying disease spread to lymph nodes is difficult using PET-CT due to its limited spatial resolution [68]. Several MRI sequences have been investigated to stage mediastinal and hilar lymph node metastases [79–81]. A recent meta-analysis of twelve studies has concluded that MRI offers an improved sensitivity and accuracy over PET-CT [82].

It remains to be seen whether MRI can complement the ubiquitous FDG-PET for diagnosis and staging of lesions. The higher spatial resolution of MRI could prove beneficial for staging and detection of smaller nodes and metastases that could go undetected in PET-CT.

3.2. Delineation, dose calculation and treatment planning

Currently, treatment simulation for lung SBRT is mostly based on CT images. The introduction of PET-CT for target definition has led to a significant decline in delineation uncertainties compared to CT-only contouring [83]. It remains to be seen whether the introduction of MRI for target delineation purposes will result in any additional benefit. Delineation uncertainties are larger for OAR as CT features relatively poor soft-tissue contrast (see Fig. 4), resulting in uncertainties of up to a few centimetres for the oesophagus and spinal cord [84]. The improved soft-tissue contrast of MRI may lead to a reduction in contouring variability for these structures.

When using MRI for treatment planning purposes, it is crucial to ensure spatial fidelity of the image data [85]. One potentially critical issue is the local differences in tissue susceptibility that can induce geometric distortions. For lung tissue, this effect can result in maximum distortions of up to 4 mm, depending on main magnetic field strength, orientation and gradient field strength [86]. However, mean geometric distortions were found to be substantially lower (\$\leq 1 mm). It should be noted that these errors are in addition to other distortions induced by non-linearities in the main magnetic and gradient field. Recently, the composite 4D distortion effect induced by scanner and susceptibility distortions was investigated for mobile lung tumours of different sizes [87]. Composite distortions were considered manageable (\$1 mm), even for highfield MRI scanners as long as these were operated in combination with high readout bandwidth at the cost of increased noise [88]. While not available on all vendor platforms, non-cartesian kspace trajectories, such as a radial or PROPELLER acquisition, can be employed for robustness against motion artefacts [89,90].

Similarly to 4D CT and 4D CBCT – albeit not as established – 4D MRI can be used to evaluate periodic anatomical changes due to breathing. 4D MRI can be either obtained based on repeated acquisition of slices covering the whole volume and respiratory cycle [91–93] or using a continuous 3D acquisition with sophisticated reconstruction techniques [94–96]. While 4D MRI based on 2D acquisitions is more easily available, the 3D-based techniques offer higher resolution and allow for a full 3D distortion correction.

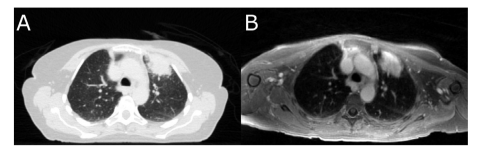


Fig. 4. (A) CT image and (B) T1-weighted MR image acquired with an ultra-fast gradient echo sequence of a NSCLC patient with a tumour located in the left upper lobe. While the bones and lung tissue are better recognizable on the CT image, the soft-tissue contrast of the MR image is superior, especially with respect to the mediastinum.

A conceptual limitation of 4D scans is that they average over several respiratory cycles and thus cannot accurately depict baseline shifts of breathing motion [97,30]. MRI could be utilised to acquire 2D cine MR images that provide a better visualisation of the patient's breathing, especially with regard to irregularities. Combining 4D MRI with 2D cine MRI using a PCA-based approach allows generation of 3D deformable vector fields with a temporal resolution of 476 ms [93]. In the future, this temporally resolved observation of the tumour motion could be used to define adequate treatment margins.

One obstacle for the widespread introduction of MRI for treatment planning is the absence of electron density and attenuation coefficient information in the MR images. Commonly, this lack of information is circumvented by either generating synthetic (referred to as pseudo from here on) CT images from MR images employing Dixon methods, or by (deformably or rigidly) registering the MR image to a treatment planning CT. The accuracy of the latter approach depends on the quality of the registration. While the generation of pseudo-CT images works well for relatively homogeneous treatment sites, such as prostate or brain [98], it is more problematic in the thoracic region, where both lung and bone feature very little MR signal. Correctly accounting for bone and lung tissue during treatment planning is key for achieving the desired dose distribution in the patient. With very limited published data available for thoracic anatomies - one exception being a study on ten patients using a segmentation/bulk density overwrite method [99] - further investigations are necessary to establish the feasibility of pseudo-CT images for treatment planning in lung cancer. Useful experiences may be gained from research on PET-MRI scanners, where electron density information is needed to correctly account for attenuation [100].

Several groups have investigated the influence of the electron return effect on MRI-guided lung SBRT treatments [101–106]. Although machine specifications, magnetic field orientations and strengths differ in these studies, all have found that the dose is distorted by the magnetic field, especially at air-tissue-interfaces in lung and skin. However, dedicated dose calculation algorithms, which are based either on Monte Carlo methods or on solving of the linear Boltzmann transport equation, are able to account for the presence of magnetic fields [107–109]. Several studies have shown that using such dose calculation algorithms to account for the ERE during treatment plan optimization allows for the design of clinically acceptable lung SBRT treatments [104–106].

In hybrid MRI treatment machines the magnetic field also affects machine and treatment plan quality assurance (QA) procedures as the ERE influences the readings of ionization chambers, solid state detectors and radiosensitive films [110–112]. Awareness of the dependence of the detector readout on its orientation relative to the magnetic field, as well as the use of dedicated detectors and phantoms are necessary for dosimetry inside a magnetic field [113–116]. Furthermore, gel-based dosimeters and a shift towards software-based QA procedures are being investigated on MRI-guided treatment units [117,118]. An electronic portal imaging device integrated with the hybrid MRI treatment unit could also be used for treatment beam QA [119].

Functional MRI offers the promise of identifying regions of the tumour that would benefit from additional dose (*boosting* or *dose painting*). While DCE MRI is expected to correlate with the oxygenation level of the tumour, DW MRI could indicate regions of increased tumour cell density [120]. Functional MRI can also identify critical OAR substructures and help minimizing dose to these. For example, for pancoast tumours DW MRI can be used to localise the brachial plexus [121]. Lung ventilation can be assessed with a variety of gaseous contrast agents [122], among which hyperpolarised ³He was employed to identify well-ventilated, healthy parts of the lung for OAR sparing in advanced NSCLC [123]. ¹²⁹Xe has also been used to

assess lung function in NSCLC patients [124] and is more widely available than ³He, but maximal hyperpolarisation is lower [122]. Due to its solubility in blood, ¹²⁹Xe could enable assessment of gas exchange [125]. A competing method that does not use hyperpolarized gases has emerged in Fourier decomposition MRI, which promises simultaneous characterisation of lung perfusion and ventilation [126]. It can be employed for functional lung avoidance mapping [127], but is free from the infrastructural requirements associated with the application of hyperpolarised noble gases.

MRI has the potential to improve OAR delineation accuracy and may allow functional evaluation of healthy tissue. It remains to be seen whether target delineation in early-stage NSCLC patients treated with SBRT will benefit from MRI, where the tumour is typically very localised with only limited involvement in adjacent soft-tissue. Issues affecting treatment planning arise from MR images not inherently containing electron density information and, on hybrid MRI treatment devices, the B₀ magnetic field distorting the dose distribution and thereby affecting treatment planning and QA procedures. While the latter issue can be accounted for with dedicated dose calculation algorithms and QA protocols, the generation of pseudo CTs for the thoracic region is still under investigation.

3.3. Patient setup and interfractional adaptations

On conventional linacs accurate patient setup is often achieved by shifting the treatment couch based on the position of the lung tumour as derived from an average or 4D CBCT [128]. Using MRI-guided delivery systems, this workflow could be replicated using 3D or 4D MR images. Potential advantages of MRI are the improved soft-tissue contrast and a larger field-of-view resulting in superior discrimination of soft-tissue structures [129].

It should also be noted that on hybrid MRI treatment machines, the ability to apply couch corrections is usually limited to the superior-inferior direction due to the restricted bore dimensions [130]. To correct for shifts in other directions or rotations, the treatment plan segments are morphed according to the position of the target immediately prior to treatment. To validate the safety of this relatively simple plan adaptation, the dose needs to be (re-) calculated on the anatomy of the day. For flattening-filter-free beams, large target shifts can introduce systematic dose deviations compared to the original treatment plan due to the sloped nature of the beam profile [131]. Compensation for this may require more complex adaptations of segment shapes and weights.

A more advanced method aims at generating a treatment plan based on the observed anatomy of the day using the initial simulation treatment plan as a starting point (warm-start optimization) [131,132]. This requires a physician, or otherwise trained member of the treatment team to check the contours, which were automatedly deformed onto the daily MR image from the treatment planning CT/MR. Re-delineation may be necessary following visual assessment [10]. This approach minimises computational costs and is expected to mitigate some of the issues related to quality control and plan approval. In the future, full re-planning based on the daily MR image (cold-start optimization) might result in even better plan quality.

The use of MRI in order to account for interfractional anatomical changes has just recently gained traction with the emergence of systems allowing MRI near or inside the treatment room. The development and automation of workflows that adapt the treatment plan to the anatomy of the day is increasingly being investigated with clinical outcome results not being available yet.

3.4. Intrafractional motion management

In the past, different methods to monitor intrafractional lung tumour motion have been developed, each with their own individual shortcomings. Internal or external surrogates do not necessarily correlate with the target motion and their relation may change over the course of a fraction [133,134,97]. Invasive marker-based localisation methods pose an additional burden to the patient. Additionally, marker migration as well as induced pneumonia have been reported [42]. Neither surrogates, nor marker-based methods are able to monitor nearby OAR or deformations of the target. Markerless tumour or OAR localisation with X-ray imaging is limited by poor soft-tissue contrast [135,136]. All X-ray based imaging methods result in an additional dose to the patient.

The promise of continuously monitoring the tumour as well as nearby healthy organs with high soft-tissue contrast MR images was a strong driving force behind the development of MRIguided treatment units. For real-time adaptive radiotherapy systems, which adjust the delivery after detecting a change in tumour position, the AAPM Task Group report 76 recommends a total system latency of less than 500 ms [13]. Currently, it is not feasible to acquire, reconstruct and post-process 3D MR images with sufficient resolution and signal-to-noise-ratio in this time frame. MRI allows acquisition of navigator echoes and use of them as a motion surrogate [137]. Visualisation of fast moving lung tumours is therefore usually based on 2D cine acquisition techniques, especially bSSFP sequences. Strategies to further increase the imaging rate have been investigated. Several techniques are based on an undersampled acquisition of the k-space data and subsequent reconstruction of the missing data using compressed sensing or viewsharing [138,139]. Others have deployed motion prediction to create a new image through extrapolation at a rate higher then the imaging frequency [140]. While coils with more than 30 individual channels are commercially available on standard clinical MRI scanners, equivalent hardware still needs to be developed for MRI-guided delivery systems. These enable advanced parallel



Fig. 5. An MR image acquired on the ViewRay system of a patient undergoing SBRT for oligometastatic cancer in the thorax at Siteman Cancer Center, St. Louis, USA. Continuous sagital cine imaging allows automated localization of the gross tumour volume (inner contour) and gating of the treatment beam if the planning target volume leaves the gating boundary (outer contour). Image courtesy of P.J. Parikh.

imaging techniques [141,142] and simultaneous multi-slice imaging [143].

Several groups have shown that it is feasible to localise lung tumours in these images using template matching [144,97,145]. More advanced algorithms deploy artificial neural networks [144,146], scale-invariant feature transforms [147,148], or particle filtering [149] to quickly delineate the tumour in each image. The orientation of these 2D imaging planes with respect to the tumour motion can be freely set and even altered during image acquisition. Studies have looked at different strategies to optimise the positioning of these planes to determine kidney or liver tumour motion [150,151], but are yet to be extended to monitoring of lung tumours. Additionally, 4D patient models built based on 4D pretreatment scans may be updated using MR images acquired during delivery [152].

The acquired real-time imaging information can support automated gating of the treatment beam on MRI-guided treatment units (see Fig. 5) [153,154]. Proof-of-concept MLC tracking has been implemented on MRI-linac prototypes [137,155]. Besides not prolonging the treatment time like gating, MLC tracking could adapt the beam aperture to tumour deformations monitored with MRI [156]. It has been shown that the presence of a magnetic field does not hinder the effectiveness of MLC tracking, which is able to reduce exposure of healthy lung tissue when compared to an ITV approach [104].

A stream of volumetric imaging information may also be used for (real-time) dose reconstruction for online quality assurance purposes [157]. Ultimately, real-time images and dosimetric information might be used to adapt the treatment plan during delivery itself, for example after the completion of each treatment beam, treatment segment or even in real-time [158]. In addition to an up-to-date 3D patient dataset, this requires ultra-fast dose calculation algorithms [107,159,160] and treatment plan optimisation strategies [161–163].

Intrafractional tumour and OAR monitoring using MRI might be able to overcome some of the shortcomings of systems based on X-ray imaging, electromagnetic markers or surrogates. However, hybrid MRI treatment devices allowing real-time MRI of the patient undergoing treatment are just becoming available. As of today, intrafractional treatment monitoring and adaptation – MRI-based or not – are only being put into practice by few institutions world-wide and most technologies described in this section are still in a research phase.

4. Conclusion and outlook

This review has highlighted elements of the clinical workflow where the integration of MRI may enhance the quality of lung SBRT. It is expected that MRI will contribute most in the areas of OAR delineation, patient setup, online motion monitoring and plan adaptations. As for most disease sites, the exploitation of MRI for lung radiotherapy is rather new. Therefore, it remains to be seen how far the technical realisation, refinement and integration into the clinical workflow can progress.

Ultimately, it has to be proven whether these conceptual advantages translate into any measurable increase in patient survival or reduction of treatment toxicities, which warrant the additional financial cost of MRI. As it stands, the 5-year absence of local recurrences for early-stage NSCLC patients being treated with lung SBRT is approximately 90% – an already very high value that rivals that of surgery, the current treatment method of choice.

If the integration of MRI is able to substantially increase delivery accuracy and, consequently, allow for higher dose conformality due to treatment margin reduction, it may be possible to deliver higher biologically effective doses to more centrally located lung

tumours. Even though there has been an interest to deliver SBRT to patients affected by these tumours, dose escalation has often been limited by normal-tissue toxicities.

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