The impact of technology on the changing practice of lung SBRT

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\begin{abstract}
Stereotactic body radiotherapy (SBRT) for lung tumours has been gaining wide acceptance in lung cancer. Here, we review the technological evolution of SBRT delivery in lung cancer, from the first treatments using the stereotactic body frame in the 1990’s to modern developments in image guidance and motion management. Finally, we discuss the impact of current technological approaches on the requirements for quality assurance as well as future technological developments.
\end{abstract}

1. Introduction

Radiotherapy has changed dramatically during the last decades, following advances in information technology and the wide progress in computer power and processing. Yet, there is a wide discrepancy in the distribution of technology not only between countries and institutions, but also among patient groups. Stereotactic body radiotherapy (SBRT) for lung tumours, also referred to as stereotactic ablative radiotherapy (SABR), is perhaps unique in the sense that it was originally targeted to patients with poor performance status (inoperable stage I lung cancer), but due to its success it is now gaining acceptance in operable patients. A large part of this success is attributable to technological advances in image guidance and motion management: increased precision in radiotherapy delivery enabled treatment of more challenging cases, which in turn meant that more patients became candidates for SBRT.

As the pool of patient candidates increases, the delivery of SBRT for lung cancer can no longer be limited to high-throughput academic centres. In this context, we observe the trend that SBRT is adopted by smaller and non-academic centres. We also observe that many technological solutions are available (see Table 1) and the investment in terms of equipment cost and human resources varies widely.

Lung SBRT is delivered in few fractions, typically 3–5. Hypofractionated treatment has benefits of preventing repopulation of neoplastic cells, a better cost effectiveness and is more convenient for patients compared to conventional fractionation \cite{Bauman2006}. Bauman et al. \cite{Bauman2006} were the first to introduce the idea of cornerstones of SBRT in 2006; in a modern context these would be: 1) target localisation, 2) treatment planning and dose calculation, 3) hypofractionation and 4) motion management during treatment delivery.

In this review, we aim to present the evolution of SBRT practice from the earliest clinical trials to today’s practice. We will review the impact of the progressive technological advances on each cornerstone in terms of clinical workflow and patient outcome, where appropriate. In addition, we provide a discussion on patient selection and which technologies are, in today’s perspective, considered a minimum standard (“must have”) and which offer incremental improvements (“nice to have”).

2. Target localisation

Target localisation encompasses patient immobilisation, imaging for SBRT treatment planning, and in-room imaging for image guidance and verification. The choices made for each of these steps will have an impact on inter- and intra-fractional accuracy of dose delivery, and should be mirrored with appropriate treatment margins and a dose prescription level adapted to the target localisation method.

2.1. Patient immobilisation

Proper immobilisation permits reproducible positioning of the patient during the course of treatment, and is of crucial importance in highly precise treatments such as SBRT. Lax et al. developed a so-called stereotactic body frame enabling the first lung SBRT treatments \cite{Lax2010}. This specially developed frame (see Fig. 1) served two purposes: it ensured the reproducible position of the patient through a head-to-
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<td>Blomgren 1995 [8]</td>
<td>Stereotactic body-frame CT (repeated if necessary)</td>
<td>Fluoroscopy AC if motion &gt; 1 cm</td>
<td>4–8 Non coplanar static beams</td>
<td>GTV-PTV 5 mm TR 10 mm CC</td>
<td>Aimed for 150%, achieved between 130 and 175% 125%</td>
<td>Varied, 10–20 Gy in 1–3 fractions (minimum dose to PTV)</td>
<td>Type A (TMS, Helax)</td>
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<td>Uematsu 2001 [22]</td>
<td>Daily verification using in room CT (slow scan) and X-ray simulator, coupled with external markers (FOCAL unit)</td>
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<td>Multiple non coplanar arcs</td>
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<td>“ITV”-like approach based on the 3 repeated CT scans + 5 mm</td>
<td>Maximum 125%</td>
<td>66 Gy x 10 (2 daily fractions) to the “border of the PTV”</td>
<td>Type A (Focus, CMS)</td>
</tr>
<tr>
<td>Baumann 2009 [63]</td>
<td>Stereotactic frame CT (repeated before each fraction if necessary) 4D CT (6–10 phases) or multiple slow CTs as alternative At treatment: online match on bony anatomy is minimum requirement KV-CT, MV-CT and orthogonal kV images are allowed</td>
<td>Fluoroscopy AC if motion &gt; 1 cm ITV or TAMP Gating, tracking, AC allowed, but not required.</td>
<td>5–9 beams</td>
<td>GTV-CTV 1–2 mm CTV-PTV 5–10 mm TR 10 mm CC ITV + 3–5 mm Or TAMP (min 3 mm, accounting for tumour motion) (larger margins if slow CT instead of 4D CT)</td>
<td>About 150%</td>
<td>15 Gy x 3 to the periphery of the PTV</td>
<td>Type A with HC</td>
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<tr>
<td>Hurkmans 2009 [69]</td>
<td>4D CT (6–10 phases) or multiple slow CTs as alternative At treatment: online match on bony anatomy is minimum requirement KV-CT, MV-CT and orthogonal kV images are allowed</td>
<td>Fluoroscopy AC if motion &gt; 1 cm ITV or TAMP Gating, tracking, AC allowed, but not required.</td>
<td>7–13 beams Usually static, but dynamic arc allowed</td>
<td>GTV-CTV 1–2 mm CTV-PTV 5–10 mm TR 10 mm CC ITV + 3–5 mm Or TAMP (min 3 mm, accounting for tumour motion) (larger margins if slow CT instead of 4D CT)</td>
<td>About 150%</td>
<td>15 Gy x 3 to the periphery of the PTV</td>
<td>Type A or B (with different dose prescriptions) HC: mandatory</td>
</tr>
<tr>
<td>RTOG 0618 [64]</td>
<td>Fiducials allowed in tumour Single 3D CT with contrast Daily orthogonal MV required as minimum IGRT</td>
<td>Gating, Breath hold, tracking, AC allowed but needs approval</td>
<td>“typically ≥ 10°” non-opposing fields, IMRT and dynamic arc allowed</td>
<td>GTV-PTV 5 mm TR 10 mm CC identical margins regardless of the technology used</td>
<td>110% &lt; hot spot &lt; 140%</td>
<td>20 Gy x 3 prescribed “at edge of PTV”</td>
<td>HC is not allowed and must be turned off if available</td>
</tr>
<tr>
<td>Lambrecht 2016 [113]</td>
<td>4D CT required along with 3D or 4D PET/CT Volumetric image guidance system (2D images allowed for gating and tracking)</td>
<td>Gating, tracking or ITV</td>
<td>No requirement other than satisfaction of target and OAR constraints</td>
<td>GTV (if no tracking/gating) Institution-specific margins (verified by CBCT before/after each fraction)</td>
<td>110% &lt; hot spot &lt; 130%</td>
<td>7.5 Gy x 8 Prescribed so that 95% of PTV gets prescription dose, 99% of PTV gets 90% of prescription dose</td>
<td>Type B required</td>
</tr>
</tbody>
</table>

Examples:
From Blomgren et al. [8], patient nr. 6 in Table 3: hotspot = $\frac{120\text{ Gy}}{90\text{ Gy}} = 133\%$

From Hurkmans et al. [69], following the prescription (i.e. no patient data): hotspot = $\frac{75\text{ Gy}}{50\% \text{ of } 120\text{ Gy}} = 139\%$

* The hotspot was calculated as followed: hotspot = $\frac{\text{maximum dose to PTV} - \text{minimum dose to PTV}}{\text{maximum dose to PTV}}$ or $\frac{\text{minimum dose to PTV}}{\text{PTV encompassing isodose}}$

‡ All doses are expressed as physical dose.
thighs vacuum pillow and provided an external coordinate system to localise the tumour inside the patient (similar to the head frames used in cranial treatment). In practice, it meant that target localisation for treatment was performed in two steps: first finding the CT-image coordinates of the tumour in the stereotactic frame and, second, aligning the isocentre of the linac with the treatment isocentre using these coordinates.

Today, several commercial versions of the stereotactic body frame are available [3], but with the advent of in-room imaging possibilities, the tumour can be visualised in one single step with the patient on the treatment couch. While some institutions retained the stereotactic frame for its immobilising properties combined with abdominal compression, others have phased it out and replaced it with devices with a lesser degree of immobilisation (such as chest boards or vacuum cushions). Frameless stereotactic treatments combined with daily image guidance [4,5] showed residual uncertainties in tumour localisation similar to that of the stereotactic body frame [6,7].

2.2. Imaging for treatment planning

SBRT also requires high quality imaging for treatment planning, with excellent visualisation of the tumour to enable precise target delineation. When the first patients receiving SBRT were treated at the Karolinska Institute in Stockholm, Sweden in 1994 [2], treatment planning based on computed tomography (CT) was used though it was not a standard procedure at the time. In addition, CT scans were acquired before each fraction in order to assess the reproducibility of the position of the tumours and baseline shifts > 5 mm in the CC direction were observed [8].

Awareness of interfraction tumour motion, combined with faster scan acquisition times, increased concerns about the reliability of conventional CT scans performed under quiet respiration: Shimizu et al. recommended the use of sequential CT scans for a more accurate estimate of the tumour localisation and appropriate treatment margins [9]. The use of slow CT scans was suggested as an alternative in the early 2000’s: images are acquired with a gantry rotation of several seconds’ duration, and the tumour is imaged throughout most of the breathing cycle, resulting in a larger (blurred) target volume and a more reproducible target visualisation than with a single conventional CT [10]. In addition, the range of tumour motion can be assessed using fluoroscopy, if the contrast between the tumour and surrounding tissue is sufficient or if fiducial markers are used (see “in-room image guidance” section below.

Respiratory correlated four dimensional CT (4DCT) was developed in the early 2000’s [11], using externally measured respiratory signal coupled to an oversampled CT acquisition. Use of 4DCT for lung SBRT was first reported by Underberg et al. [12]: they found 4DCT advantageous over sequential 3D CTs for correct evaluation of the extent of individual tumour motion. Motion-related image artefacts were also reduced compared to regular 3D CT acquisitions [13,14]. However it should be remembered that the 4DCT remains just a snapshot of the tumour motion and position: the respiratory pattern may change with time [13] and in-room image guidance is required to verify the tumour position (and possibly, the position of the organs at risk) before each treatment fraction.

In order locate and delineate the tumour with high precision, high spatial resolution imaging is also necessary. The slice thickness of the CT images used for SBRT for the very first patients was 10 mm [2]. It has since been shown using phantom studies that the size of a small lesion is increasingly overestimated by large slice thickness [15], and for modern lung SBRT a slice thickness of ≤ 2 mm is desirable [16] (Table 2). Positron emission tomography (PET) for radiotherapy treatment planning is becoming increasingly available. Though it has been suggested that 4D PET/CT images (compared to 3D PET/CT) improve the inter-observer target delineation agreement for centrally located lesions [17], the primary role of PET imaging in lung SBRT remains to confirm the absence of disease spread, prior to referral [18]. Delineation uncertainties for the gross tumour volume (GTV) in lung SBRT of the order of 2 mm have been estimated by Persson et al. [9], when combining 3D PET/CT and 4DCT information. Although small, these uncertainties should still be included in the margins to be applied for treatment, to avoid geographical miss of part of the tumour.

Lung tumours can have a complex and irregular respiratory motion pattern. As a result, respiratory motion can degrade image quality in all imaging modalities, even respiratory-related ones. This needs to be evaluated and taken account for during imaging, or during registration of multiple images. Most tumours move predominantly in the cranio-caudal direction [20] and have moderate peak-to-peak amplitude of ≤5 mm [21]. The degree and complexity of motion cannot be predicted from the anatomical tumour position, although in general, tumours in the lower lobes move more than those in the upper lobes of the lung.

2.3. In-room image guidance and management of geometrical uncertainties in the position of the tumour

In-room imaging for treatment guidance for lung SBRT was first reported using in-room CTs [22]. It became more widely available with the development of electronic portal imaging devices (EPID), enabling 2D MV imaging of either treatment fields beam’s eye view or orthogonal setup images. However, MV image quality is poor, and visualisation of lung tumours is often not possible [23,24]. Planar 2D kV imaging decreases imaging dose and improves image quality of the bony anatomy. In a selected group of patients (e.g., patients presenting with peripheral and well defined tumours), 2D kV imaging can be used to directly visualise the tumour [25]. However, using bony anatomy as a surrogate for tumour position does not match the high precision requirements of modern lung SBRT and should be avoided [26,27] (see Table 2). 2D image registration based exclusively on bony anatomy is unfortunately still applied at a range of institutions, despite the well documented inter-fractional deviations between the bony anatomy and the SBRT lung target, ranging as high as 3 cm [6,28].

Visualising the tumour position is of paramount importance for daily verification in order to avoid geographical miss [3,29]. Percutaneously implanted fiducials can be used as an adequate surrogate for

Fig. 1. The stereotactic body frame, as designed by the Karolinska institute. The coordinate system consists of a set of graduated scales, visible on CT images and is used to position the isocentre before each fraction. Abdominal compression was used if the motion of the diaphragm was estimated to be over 5 mm as assessed under quiet respiration fluoroscopy. Image courtesy of Kristin Karlsson, Karolinska institute.
the position of the tumour, since it was shown that markers with non-smooth surfaces are stable within the tumour throughout the treatment course [30–32]. However the implementation process comes with a high risk of pneumothorax [33–35]. This risk may be reduced by using endovascular coils [36] or bronchoscopy-guided implantation [37] but it remains present. In contrast, volumetric imaging is a non-invasive procedure where the tumours and the adjacent organs at risk can visualised. The use of in-room CT scanners on rails [28] and, more widely available, on-board kV cone beam CT (CBCT) resulted in improved geometrical precision of lung SBRT [5,6,38,39] and reduced interfractional uncertainties.

Typical CBCT acquisition takes around one minute and hence consists of several breathing cycles, which may impact visibility and increase interobserver variation during registration. In the presence of substantial respiratory motion, time resolved 4D CBCT has been shown to improve image quality [40] and reduce the inter-observer variability associated with manual image registration [41]. In order to reduce the dose associated with 4D CBCT, O’Brien et al. have proposed reducing the number of projections acquired by using the breathing signal as an input [42]. At the time of writing, online pre-treatment 4D CBCT is not yet commercially available from all vendors of medical linear accelerators (linacs), but it is nonetheless gaining acceptance where available. The development and utilisation of 4D CBCT should be encouraged, especially in patients with small and very mobile tumours which can “disappear” on 3D CBCT image (Fig. 2a). Further work on the clinical implementation of 4D CBCT is required in order to make this form of in-room guidance available as a routine option for lung SBRT patients. As an alternative, compliant patients with small mobile tumours can be imaged and treated in deep inspiration breath-hold (DIBH) (Fig. 2b).

Respiratory motion management strategies were used from the first implementations of lung SBRT. The most commonly reported strategy was abdominal compression [8,22,43,44], but Japanese groups introduced breath hold, gating and even tracking in the early 2000s [45–48]. The efficiency of abdominal compression has been questioned: it may result in increased inter-fractional variation [49] and, while respiratory motion is decreased in some tumours, it is actually increased in others [50]. Strategies for gating and voluntary or controlled breath hold are covered in detail in Caillet et al. (this issue) [51] but it is important to remember that none of them completely eliminates breathing-related uncertainties. In addition, respiratory management strategies may only be applicable in patients who are relatively fit and compliant, either because they can create discomfort (active breathing control, abdominal compression), require active patient cooperation (voluntary breath hold) or increase the treatment delivery time (gating). Hence, appropriate patient selection is crucial for the implementation of those techniques which should be limited to compliant patients who get a substantial benefit in terms of improved accuracy.

In patients treated in free-breathing, remaining intra-fractional uncertainty of the tumour position, evaluated from a pre- and post-treatment CBCT would result in systematic and random errors within ∼2 mm [5,6,52]. A mid-point (e.g. half way through a SBRT treatment) CBCT can be acquired to ensure minimal intra-fraction baseline shifts [53].
Surprisingly, intra-fractional uncertainties do not seem related to the degree of immobilisation [38] or extent of tumour motion [5] but it has been suggested that they are substantially increased with longer treatment duration (exceeding 34 min) [6]. However, tracking the implanted fiducials during beam-on time revealed substantial intra-fractional base-line shifts occurring at an earlier time point [54]. Intra-fraction monitoring is desirable and might reveal otherwise undetected shifts in some patients.

Acquisition of intra-fraction CBCT images for lung SBRT has also been described [55–58] by reconstructing CBCT images from planar kV images acquired during VMAT irradiation. This allows verification of the tumour position and movement during free-breathing, whilst the treatment beam is on, although there is a minimum treatment arc angular range of around 80° required in order to generate useful CBCT images. Current clinical implementation requires the kV fluoroscopy beam to be on during the entire treatment arc, so there may also be scope to reduce the imaging dose, e.g. by gating the kV fluoroscopy. These techniques are currently being introduced and commissioned by large academic centres, and some further development is required before they become widely available for routine implementation from all vendors.

2.4. Treatment margins

While GTV and clinical target volume (CTV) in lung SBRT are often considered to coincide [3], the planning target volume (PTV) should take into account all other uncertainties. The ITV concept, introduced in ICRU 62 was to account for internal organ motion separately from set-up or geometrical errors, which are then added to the PTV margins. However, it has been pointed out that the uncertainties should not be added linearly [59] and that use of ITV resulted in an “unnecessarily large volume” for random respiration variability. The ITV concept is still widely used in lung SBRT, but with slightly different approaches, resulting in a smaller treated volume: a simple fusion of GTV volumes delineated in inhale and exhale phases, delineating the GTV on the resulting PTV may be smaller than with ITV-based margins [4,66]. As the maximum dose was 100%, this means that for random respiration variability the PTV encompasses the PTV (see examples in Table 1). In RTOG 0236, 0618 and 0915 lung SBRT trials the prescription dose was three fractions of 20 Gy, prescribed to the isocentre which encompasses the PTV (see examples in Table 1). In RTOG 0236, 0618 and 0915 lung SBRT trials the prescription dose was three fractions of 20 Gy, prescribed to the PTV encompassing 65% isodose [71]. This can be presented equivalently, as the 100% isodose encompassing the PTV with a 140%-150% hot spot at the isocenter [1]. The resulting high dose fall off outside the PTV also limits the exposure of nearby organs at risk. However, there is no inter-institutional agreement or international code of practice for prescribing and reporting the dose to the target in lung SBRT. The most common practice is prescription to the isodose which encompasses the PTV for prescribing and reporting the dose to the target in lung SBRT. The most common practice is prescription to the isodose which encompasses the PTV for the prescription of 65% isodose [71]. This can be presented equivalently, as the 100% isodose encompassing the PTV with a 140%-150% hot spot at the isocenter [1]. Within the ROSEL trial, the RTOG 0618 dose prescription was specified in a similar way, with 95% of the PTV receiving nominal fraction dose of 20 Gy, and aiming for a maximum dose of 110–140% [69]. A discussion of how to achieve these

3. Dose planning and calculation

3.1. Dose prescription

Inhomogeneous dose distributions were first rationalised for cranial stereotactic radiosurgery [68], where the hot spot in the centre of the tumour could be beneficial for eradicating an increased density of clonogenic tumour cells. The same argument was used upon implementation in extra cranial targets [1]. The resulting high dose fall off outside the PTV also limits the exposure of nearby organs at risk. However, there is no inter-institutional agreement or international code of practice for prescribing and reporting the dose to the target in lung SBRT. The most common practice is prescription to the isodose which encompasses the PTV for prescribing and reporting the dose to the target in lung SBRT. The most common practice is prescription to the isodose which encompasses the PTV for the prescription of 65% isodose [71]. This can be presented equivalently, as the 100% isodose encompassing the PTV with a 140%-150% hot spot at the isocenter [1]. Within the ROSEL trial, the RTOG 0618 dose prescription was specified in a similar way, with 95% of the PTV receiving nominal fraction dose of 20 Gy, and aiming for a maximum dose of 110–140% [69]. A discussion of how to achieve these
inhomogeneous dose distributions is presented by Giglioli et al. in this issue [72].

Studies have also been published with different dose prescription approaches: to mean PTV dose, maximum dose and even with relatively homogeneous dose distribution within the target [73,74]. As discussed by Ricardi et al. in this issue [75], this is still considerable debate about the optimal dose prescription and fractionation in lung SBRT. The variety of dose prescription practices (see table 1 and the calculation example provided) makes it challenging to compare studies from a dosimetric point of view and the introduction of standard methods of dose reporting are sorely needed.

3.2. Dose calculation algorithms

The accuracy of the calculated dose distribution in lung SBRT is affected by uncertainties related to small-field dosimetry (which will impact input data used in dose calculation algorithms), and handling the lateral electron transport and charged particle disequilibrium. Knöös et al. divided dose calculation algorithms used in commercially available treatment planning systems into simple ones not taking into account changes in electron transport (type A), and more complex ones (type B), which partially account for these changes in electron transport [76]. Both type A and B algorithms are analytical or semi-analytical. The third-generation algorithms, recently referred to as type C, solve the linear Boltzmann transport equations either stochastically (Monte Carlo) or deterministically (Acuros XB). Modelling physical interactions of particles in media enables reporting absorbed dose as dose-to-media as opposed to widely used dose-to-water. There have been some controversies as to which to use [77], and AAPM TG 105 recommended indicating the material to which the dose was computed, and allowing conversion from dose-to-medium to dose-to-water, until clinical justification for either method was assessed [78]. The properties and dosimetric impact of these different algorithms are reviewed in detail in other articles within the present issue [66,79]. The choice of dose calculation algorithm in lung SBRT has been shown to have a significant impact on treatment outcomes. Recurrence rates for type A algorithm planned SBRT were higher (hazard ration 3.4; 95% CI: 1.18–8.83) [80], despite tumours treated in the type A group were smaller [80]. 2 years local control was in better agreement with dose recalculated with type B or C algorithms [81].

Recent SBRT trials require use of type B or C algorithms (RTOG 0915) or suggest differentiating the prescribed dose according to the dose calculation algorithm of choice [69,82].

The accuracy of the calculated dose distribution is also affected by the calculation grid size. AAPM TG 101 recommends using 2 mm or smaller grid size [3], although a 1 mm calculation grid size for Acuros XB has not shown statistically significant benefits for lung SBRT delivered with VMAT [83]. Optimisation of Monte Carlo approaches is discussed by Chetty et al., however without focus on SBRT [78].

3.3. Dose prediction and verification prior to delivery

Recent developments in handling 4D-CT image datasets in commercial treatment planning systems (TPS) have opened up the possibility of in silico assessment of treatment plans across all breathing phases. These capabilities have so far been used more in the research setting than in clinical routine [84] and rely heavily on the use of deformable image registration (DIR). The accuracy of DIR algorithms in commercial TPS is probably sufficient in the high-contrast thoracic region to be useful for propagation of the outline of normal tissues. A 3–5 mm variation in contour accuracy between different commercial software packages has been found [85], and variations of the same magnitude are also found when using the same software package, but different workflow. This highlights that work still remains to be done on establishing guidelines for use of DIR software, along with an evaluation of the size of registration errors for specific tumour sites [86].

The development of semi-automated tools (which allow re-calculation of the treatment plan dose on each phase of the 4D CT) now permit inspection of the dose distribution(s) throughout the breathing cycle prior to treatment. Plan assessment tools such as DVH graphs with error limits or confidence intervals to guide the user in assessing multiple plans would be helpful in future software development.

Use of DIR for dose warping or dose accumulation (to sum the dose delivered across all breathing phases onto a single reference phase) is more problematic. Errors in the deformation vector, particularly in zones with a high dose gradient could lead to very significant errors in accumulated dose [87,88]. Further work is also necessary to fully examine the interplay effect – this requires information about the duration of the breathing cycle, and timing of the treatment delivery, particularly for intensity-modulated beams. Inverse optimisation of plans using multiple phases of the 4D CT image dataset could also be implemented in the future, to create treatment plans which are ‘robust’ to breathing motion. Experimental evaluation of the accuracy of 3D versus 4D planning suggests that for each patient, uncertainties in 4D planning need to be smaller than the uncertainties in the standard 3D approach, in order for this technique to be adopted in clinical routine [89].

4. New delivery techniques in lung SBRT

The development of volumetric modulated arc therapy (VMAT), in particular when coupled with the introduction of flattening filter free (FFF) beams with dose rates of 2400 MU per minute, has allowed lung SBRT plans to be delivered much more rapidly and conveniently than multiple static beams. Delivery of lung SBRT plans using FFF arc therapy has been shown to be feasible with acceptable acute toxicity rates [90].

Some caution is required when selecting patients for these treatments, to avoid any kind of Interplay effect, where the periodic nature of tumour displacement, and the temporal pattern of dose delivery could overlap to degrade the delivered dose compared to that planned on the ‘static’ CT image used for planning. Experimental phantom studies have suggested that the interplay effect is significant for large tumour displacements (> 3 cm) or very long respiratory patterns (period > 60 s) [91].

Specialised treatment machines, such as the Cyberknife robotic radiosurgery system, allow tracking of the tumour in real-time, by constructing a patient specific correlation model to monitor the respiratory tumour movement, and reposition the treatment beam using the robotic treatment arm. Briefly: external optical markers (endpoints of optical fibres that transport the light of light-emitting diodes) are placed on the chest or abdomen of the patient and are monitored by a camera array and correlated with the tumour location derived from orthogonal kV X-ray images of the patient anatomy. The correlation model is constructed prior to each treatment fraction and verified and adapted during each treatment fraction. Fiducials can be used for target localisation, and Cyberknife treatment using fiducial tracking has been found to give comparable results for 3 year local control (91%) as for surgical resection [92]. A fiducial-less tracking system (Xsight lung tracking) is also available, and measurements on patient specific lung phantoms indicate this tracking accuracy (with mean error 0.38 ± 0.54 mm cranio-caudal) is comparable to tracking using fiducials [93]. However, not all lung tumours may be amenable to tracking using the fiducial-less system – small tumours with low density being more challenging to visualize during treatment [25]. If implanted fiducials are not available and the tumour cannot be localized directly by one of the two X-ray cameras, there is an option to use only one X-ray camera for real-time tracking. A treatment margin is then needed to encompass the motion that is blurred to the other X-ray camera and is therefore not being tracked. If implanted fiducials are not available and the tumour cannot be localized by either X-ray camera, margins need to be added that account for the full inhale-to-exhale motion of the tumour.
Efforts are also made to increase the potential of DIBH treatments: Péguret et al. have reported achieving apnea-like conditions lasting up to 16.5 min using a pneumatic ventilation system [94]. Though compliance was acceptable in the healthy volunteers, one out of five patients could not hold a long enough breath hold to be treated with this approach and further investigation is necessary to judge whether this approach is feasible in clinical practice.

5. Quality assurance

Geometric tests for verification of the alignment and coincidence of the mechanical and radiation isocentres and the image guidance system should be tighter for machines delivering SBRT treatments: for example, a maximum difference of ±1 mm compared to the indicated position, and ±0.5° in rotation (as opposed to ±2 mm/1° for conventional RT) has been recommended [3,95]. QA procedures have to be performed frequently and should include a daily check of the coincidence of treatment (MV) and imaging (kV) isocentres [27,29]. For volumetric vs planar kV in-room imaging, reported isocentre deviations were within 1 mm [96].

Although checking each component individually is required, end-to-end (E2E) testing with simple geometric phantoms [97,98] is also recommended, and developments in EPID technology mean these tests can be now be semi- or fully automated [99].

For lung SBRT, a patient specific QA program using E2E tests with a heterogeneous anthropomorphic phantom [100] is required, as this can be used to verify the mechanical, dosimetric and imaging components for SBRT delivery, and also check correct transfer and application of image-guidance positioning data (see an example in Fig. 3). Additional QA may be required when using a robotic “6 degrees of freedom” couch, to correct for rotational errors in patient position [101]. Hurkmans et al. [102] showed that differences in 4D-CT protocols (due to, e.g. type of scanner, breathing surrogate or image reconstruction strategy) could introduce systematic errors in the treatment planning process: regular QA checks using a moving phantom can help optimise 4D acquisition protocols and should be included in the commissioning program. Consideration should be given to the quality of images to be used during real-time imaging for each patient, as image contrast or target visibility may be degraded compared to studies using phantoms due to patient size or other artefacts. This is especially important if tracking or gating is applied, and specific QA procedures relevant to the exact configuration of each treatment and imaging system will be necessary [103].

The challenges associated with small field dosimetry have been described [27], and use of a detector with an active area of 1 mm² or less is recommended for SBRT dose measurements [3]. A new formalism for small-field reference dosimetry has been proposed, which is also applicable to treatment machines such as CyberKnife or Tomotherapy where the reference geometry outlined in the standard code of practice is not achievable [104]. In addition, the development of flattening-filter free (FFF) linacs (e.g. Varian Truebeam and Elekta Agility HD) which do not deliver ‘flat’ beam profiles also means that standard parameters for field homogeneity and flatness are no longer applicable. Novel parameters for beam profile consistency checks have recently been proposed and analysed [105,106].

Frequent use of non-co-planar fields means that in addition to collision verification, the dosimetric effects of the treatment couch and immobilisation devices on beam attenuation or increased skin dose should be considered in the delivery of high-dose per fraction treatments with standard or FFF beams [107,108].

A list of guidelines and protocols specific to SBRT is given in the Supplementary Table material. These also give guidelines on staffing and resources needed to implement a successful SBRT program. Reports for commissioning and benchmarking QA procedures for specialised machines such as CyberKnife [103], Novalis [109], Vero [110] are also available. In addition, a review of in vivo dosimetry methods is given by McCurdy et al. [111] (this issue).

6. Discussion & conclusion

SBRT has led the development of technologies that are today widely used for other treatment schemes and diagnostic sites. The improvements in local control and overall survival in patients treated with SBRT cannot be ascribed to technology alone, but it has certainly played a role: improved SBRT technology facilitated safely escalating the dose to the tumour, which in turn seems to correlate with improved outcomes [112]. The potential of future technological developments may be classified in three categories: 1) workflow improvements (such as FFF or automated planning and contouring), 2) improvements likely to show a clinical benefit in a small number of patients, and 3) improvements likely to benefit the group of SBRT patients as a whole. Innovations in intra fraction management arguably belong to the second category and will be welcome for a subgroup of patients with very mobile tumours. Improved imaging for contouring of tumours and organs at risk is also desirable especially in central lung tumours [113], but we are presently missing data about which structures drive toxicity and how much they should be spared. When this knowledge becomes available, advanced delivery techniques (such as proton therapy or MRI-guided RT (Menten et al. [114] this issue)) may prove valuable for a larger group of patients.

The next technological priority belonging to the third category and benefiting the largest group of patients should be on standardisation (for example, in reporting heterogeneous dose distributions) and/or a better use of the technology available today. This means more thorough commissioning and rigorous QA so that the present technology can be used close to its potential accuracy. It is worth noting that even in these optimal conditions, uncertainties will be present and zero-margins are unlikely to be achieved.

Optimal use of the technology currently available will both benefit clinical practice and ensure the highest quality in clinical trials.

Fig. 3. Motion phantom used for end-to-end testing of tracking at Erasmus MC.
Unfortunately, this is not yet the case and a few current trials still allow the use of 2D portal imaging for peripheral lung tumors despite the wide availability of CBCT (or other methods which enable to visualize the position of the tumor) and the well-documented uncertainty of portal imaging.

Along with developing new tools and making the most of the ones we already have, the ultimate goal is to achieve high quality and high precision lung SBRT in both clinical trials and “daily” treatments, in both academic and non-academic institutions, benefiting all SBRT patients.

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Appendix A. Supplementary data

Supplementary data associated with this article can be found in the online version, at http://dx.doi.org/10.1016/j.ejmp.2017.12.020.

References


