Developments in radiation techniques for thoracic malignancies

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Shareable abstract (@ERSpublications)
Technical advances have led to a changing perception of the role of radiation therapy in multidisciplinary care of lung cancer. This article provides an overview of recent developments in radiation therapy as a cornerstone of modern lung cancer treatment.

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Abstract
Radiation therapy is a cornerstone of modern lung cancer treatment alongside surgery, chemotherapy, immunotherapy and targeted therapies. Advances in radiotherapy techniques have enhanced the accuracy of radiation delivery, which has contributed to the evolution of radiation therapy into a guideline-recommended treatment in both early-stage and locally advanced nonsmall cell lung cancer. Furthermore, although radiotherapy has long been used for palliation of disease in advanced lung cancer, it is increasingly having a role as a locally ablative treatment in patients with oligometastatic disease. This review provides an overview of recent developments in radiation techniques, particularly for non-radiotherapy oncologists who are involved in the care of lung cancer patients. Technical advances are discussed, and findings of recent clinical trials are highlighted, all of which have led to a changing perception of the role of radiation therapy in multidisciplinary care.

Introduction
Lung cancer accounted for more deaths than breast, prostate and colorectal cancers combined in 2020 [1]. However, data from the Surveillance, Epidemiology and End Results (SEER) programme revealed that declines in mortality had accelerated for lung cancer in the period from 2008 to 2017, a finding which may be related to the rapidly evolving treatment landscape for lung cancer, especially for nonsmall cell lung cancer (NSCLC). In the past decade, molecular targeted therapies, immune checkpoint inhibitors and improved high-precision radiation delivery have joined the established modalities of chemotherapy, large-field radiotherapy and surgery in thoracic disease management. With the availability of new treatment options, the role of multidisciplinary tumour boards has become crucial in selecting and tailoring treatment strategies, and for managing toxicity and survivorship issues.

Updated guidelines have incorporated many of the incremental improvements in radiotherapy planning and delivery [2–4]. There are now more standardised definitions of target volumes, improvements in radiotherapy plan quality (including daily on-table treatment plan revisions) and reductions in organ at risk doses, all of which have increased clinician confidence to deliver ablative doses of radiation. Changes in thoracic radiotherapy guidelines [5–7] have partly been driven by results of studies showing improvements in population outcomes following the implementation of new radiotherapy techniques [8–12]. Recent prospective trials in early-stage lung cancer [13], locally advanced NSCLC [14, 15] and oligometastatic (lung) cancer [16–20] have also contributed to a changing perception of the role of radiation in multidisciplinary care.
This review aims to provide an overview of recent developments in radiation techniques for non-radiation oncologists involved in the care of patients presenting with thoracic malignancies.

**Radiation therapy for early-stage lung cancer**

Nearly 25 years ago, the first clinical experience with so-called stereotactic radiation therapy for lung tumours was reported [21]. This technique, now termed stereotactic ablative radiotherapy (SABR) or stereotactic body radiation therapy, has since become the preferred treatment for early-stage NSCLC in patients who are medically inoperable or unwilling to undergo surgery [2, 3, 5]. Briefly, SABR is an approach used to deliver high radiation doses to tumours outside the brain, by using precise techniques to ensure that the tumour position is reproducible, while sparing critical surrounding normal tissues from toxicity. Curative treatments can then be delivered in fewer fractions than with use of conventionally fractionated radiotherapy, as on-board imaging can verify the tumour position before each session (figure 1).

The superior tumour control rates with lung SABR compared to conventionally fractionated radiotherapy have been demonstrated in large institutional series, as well as two randomised trials, one of which also showed improved overall survival with SABR [13, 22]. Lung SABR is typically delivered in between one and eight fractions during a period of up to 2 weeks in an outpatient setting. For selected peripherally located lung tumours, SABR can be delivered in a single session, based on the findings of two randomised studies in early-stage NSCLC showing outcomes comparable to SABR delivered in three or four fractions [23, 24]. Local control rates with SABR can exceed 90% for early-stage NSCLC and are generally considered comparable to those achieved with sublobar resection [25–27]. Regular computed tomography (CT) imaging is recommended after SABR in order to distinguish common patterns of focal fibrosis (figure 1) from so-called high-risk features that are suspicious for recurrence [28, 29]. Multidisciplinary assessment of radiological changes that evolve with time may aid in reassuring patients and also avoid unnecessary invasive procedures. Furthermore, multidisciplinary review is necessary for coordinating salvage treatment options in patients with a confirmed post-SABR recurrence [30].

Prospective data comparing SABR to lobectomy, the current standard of care in stage I NSCLC, are limited [31], but randomised trials comparing SABR to surgical resection are ongoing [32]. With the population of elderly patients with lung cancer increasing rapidly, it is relevant to note that SABR is well tolerated in unfit patients with multiple comorbidities, including severe COPD [2]. Elderly patients presenting with an early-stage NSCLC show an improved survival with SABR compared with observation alone [8]. A Dutch population study demonstrated increases in survival of elderly patients with stage I NSCLC following the introduction of SABR, which was also associated with a decrease in the numbers of patients who were untreated [9]. Following the introduction of SABR, significant changes in patterns of care for the elderly were observed in the Dutch population-based study, with 84% of patients aged ≥80 years now undergoing SABR for stage I NSCLC [33]. However, patient groups who are at increased risk for SABR-related toxicity have also been identified, such as those with tumours abutting central

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**FIGURE 1** Serial diagnostic images of a peripheral stage I nonsmall cell lung cancer treated using a) stereotactic ablative radiotherapy (SABR) to a dose of 55 Gy, delivered in five fractions, on a linear accelerator. An on-board cone-beam computed tomography (CT) scan is used for the daily on-couch verification of tumour position. b) The planned dose distribution is illustrated using so-called isodose lines, representing 50% (yellow), 75% (purple) and 100% (green) of the prescribed radiation dose. High-precision dose delivery to the tumour minimises irradiation of surrounding normal tissues. c) Follow-up images at 14 months reveal ground-glass changes and focal fibrosis in the treated region. d) At 36 months, residual scarring is present, with a positron emission tomography-CT showing no residual 18F-fluorodeoxyglucose uptake in the lesion.
airways [34–37], with large tumours [38], or patients with coexisting interstitial lung disease (ILD) [39, 40]. For patients who are at a higher risk of surgical complications, SABR can offer a lower risk alternative, both for primary and recurrent NSCLC [41]. Patients undergoing surgery for a second NSCLC were found in a meta-analysis to have a pooled operative mortality of 7% for the second resection and a pooled 5-year overall survival of only 46% [42]. Life-long follow-up after any initial treatment for lung cancer is supported by population studies showing that the post-surgical cumulative risk of second primary lung cancer is 20% at 10 years post-surgery and 25% at 14 years post-surgery [43]. In patients undergoing SABR for early-stage NSCLC, distant (or out-of-field) recurrences remain the most common pattern of failure [44]. In order to reduce the risk of distant recurrences, combinations of immune checkpoint inhibitors and SABR in the early-stage or locally recurrent setting are currently being evaluated in phase II (e.g. I-SABR) and phase III (e.g. PACIFIC-4 and KEYNOTE-867) clinical trials [45–47]. The rationale supporting immune-SABR involves preclinical studies showing that the immunomodulatory effects of short-course radiotherapy can enhance both local and systemic antitumour immune response [48].

Further improvements in SABR delivery are desirable, for example by using techniques such as intensity-modulated radiation therapy (IMRT), an approach which allows for reductions in dose to critical organs such as the heart, lungs and oesophagus [49–51]. Volumetric modulated arc therapy (VMAT) is an approach which allows for the faster delivery of IMRT, which is now recommended in clinical practice guidelines for SABR [2]. SABR can also be delivered using proton beams, a technique where charged particles are used to deposit their energy more precisely in the tumour, thereby potentially reducing radiation doses to normal tissues. However, proton-based lung SABR is technically challenging as respiration and cardiac-induced tumour motion can degrade the quality of proton treatment plans even more than with conventional beams [52, 53]. Proton-based lung SABR is currently undergoing evaluation in clinical trials [54–56].

Better tumour targeting in SABR can improve tumour control rates, and different approaches have been endorsed in clinical practice guidelines [2, 4]. A recommended strategy for lung tumours involves the use of a 4-dimensional (4D) CT scan for treatment planning, and all observed motion is used to generate a so-called internal target volume (ITV). However, in cases where tumours exhibit significant motion, the ITV approach can expose the healthy lung to a higher radiation dose [57]. Daily pre-treatment verification of the tumour position is mandatory [2], and this is commonly performed using integrated cone-beam CT technology on a conventional linear accelerator. However, as tumour motion during treatment delivery can exceed that observed on pre-treatment 4D-CT imaging [58, 59], motion monitoring during SABR delivery is desirable. Respiration-gated radiotherapy refers to radiation delivery that is restricted to a predefined tumour position and/or phase of the patient’s respiratory cycle. Another approach in clinical use is tumour tracking, where the treatment beam continuously follows or is shaped to the tumour position [57]. Such so-called active motion management approaches can permit use of smaller treatment volumes [60–63]. However, motion management involving implantation of fiducial markers [64] or electromagnetic transponders [65] carries risks, especially in the elderly and frail patients [64, 66, 67]. Noninvasive monitoring of respiratory movements is possible using spirometric devices [68] or infrared or optical cameras to monitor surface breathing motion [68, 69], although these approaches can fail to accurately capture variations in tumour position [70].

Recent advances in radiotherapy technology aim to circumvent the need for implanted fiducials, or other surrogate markers, to capture tumour motion in real-time. Since 2014, direct tumour visualisation during radiation delivery has become possible with use of so-called magnetic resonance-guided radiotherapy devices. These hybrid machines incorporate both magnetic resonance imaging and radiotherapy technology into a single treatment system, which allows for continuous acquisition of magnetic resonance images during treatment [71, 72]. On-board magnetic resonance technology facilitates tumour gating without additional radiation exposure, and enables daily on-table plan adaptation, which improves treatment plans if doses to the tumour and/or critical organs are suboptimal due to changing anatomy. Magnetic resonance-guided lung SABR can be delivered with high precision [73], to significantly smaller target volumes than with a traditional motion-encompassing ITV approach (figure 2) [60]. In addition, the safety of SABR delivered to tumours in the proximity of critical organs at risk is improved using magnetic resonance guidance (figure 3) [74–77].

Developments such as magnetic resonance-guided radiation therapy, or proton therapy, serve to illustrate more complex techniques that are required to further improve treatment accuracy and reduce doses to healthy tissue. The latter becomes more important when performing so-called hypofractionation, which refers to the delivery of large radiation doses in a shorter overall treatment time. Hypofractionation, including single-fraction treatment, is being encouraged for reasons including better resource utilisation,
patient comfort and easier scheduling of SABR treatments in patients with oligometastases in whom systemic therapies cannot be interrupted for long [78]. In the coronavirus disease 2019 (COVID-19) era, the European Society for Radiotherapy and Oncology (ESTRO) and the American Society for Radiation Oncology (ASTRO) have also recommended performing single-fraction lung SABR for stage I...
peripherally located NSCLC [7], and developments such as magnetic resonance guidance can increase clinician confidence in doing so [79]. However, such resource-intensive procedures are best suited for high-risk tumours, such as those in the hilar region, tumours arising after prior lung radiotherapy and when coexisting ILD is identified [80]. This technology is new, and technical challenges remain as magnetic resonance resolution is limited by respiratory and cardiac motion, and by the low signal-to-noise ratio of lung parenchyma [81].

Locally advanced lung cancer
Locally advanced NSCLC comprises a heterogeneous group of patients, including tumours with advanced local infiltration (T4), or ipsilateral (N2) and contralateral (N3) mediastinal lymph node metastases, among other criteria. These stage III tumours make up ≈35% of all NSCLC cases at diagnosis [82]. The management of stage III NSCLC is complex and requires assessment by a multidisciplinary team. Many patients with stage III NSCLC are considered to be unresectable due to the extent of disease, even after induction therapy. However, in potentially resectable stage IIIA NSCLC, the standard of care should include consideration of surgical resection after multidisciplinary review [82, 83]. The overall prognosis of locally advanced NSCLC patients is poor even in resectable patients, for whom a 5-year overall survival of only 29.0% and 13.0% was observed for stage IIIA and IIIB disease, respectively [84].

Treatment paradigms for patients with inoperable stage III NSCLC have evolved in recent decades, from sequential to concurrent chemotherapy and radiotherapy (chemoradiotherapy; CRT), with the recent addition of consolidation immunotherapy as a standard of care in non-progressing patients [85]. Concurrent platinum-based CRT is the recommended treatment backbone for fit patients with unresectable stage IIIA and IIIB disease, as it improves 5-year overall survival compared to sequential CRT [86]. In the recent PACIFIC trial, 12 months of immune checkpoint inhibitor therapy with durvalumab, a monoclonal antibody targeting programmed death-ligand 1, improved overall survival compared to placebo in patients who did not progress following CRT. Updated follow-up at a median of 34.2 months revealed a median overall survival of 47.5 months in the durvalumab arm, as compared to 29.1 months in the placebo arm [14, 15, 87].

Guidelines by the European Society for Medical Oncology for concurrent CRT recommend the delivery of 60–66 Gy in 30–33 daily fractions of 2 Gy, with the maximum overall treatment time not exceeding 7 weeks [5]. The dose of 60 Gy is considered a standard based on the Radiation Therapy Oncology Group (RTOG) 0617 study, in which a higher dose of 74 Gy was associated with worse overall survival [88]. Although modern radiotherapy techniques may allow for safer dose escalation to, for example, 66 Gy, this is generally not recommended in patients with extensive nodal disease due to increased oesophageal toxicity. In both concurrent and sequential CRT, the choice of chemotherapy regimen is based on histology [82]. Concurrent CRT is associated with higher acute oesophageal toxicity rates (grade 3–4) compared to sequential CRT (18% versus 4%) [86]. Patients who are considered unsuitable for concurrent CRT, for example due to patient frailty or extent of disease, are candidates for sequential CRT, which is a valid alternative according to European Society for Medical Oncology guidelines [5, 85]. For sequential CRT, the radiotherapy scheme is preferably delivered in a shorter overall treatment time using hypofractionated regimens. Such shorter treatments, e.g. delivery of 55 Gy in 20 fractions of 2.75 Gy [89], appear desirable in the COVID-19 era, during which the treatment of patients with stage III NSCLC can be challenging, but should not be delayed [90]. However, the ESTRO/ASTRO consensus recommended that hypofractionated radiotherapy should only be used with radiotherapy alone or sequential CRT, whereas this should be avoided when large volumes are treated using concurrent CRT [7].

Toxicity can be reduced by limiting radiation fields to sites of known disease. For operable lung tumours, use of a systematic endobronchial ultrasound (EBUS) followed by an oesophageal investigation using the same EBUS-scope increases sensitivity for the detection of N2/N3 disease by 9% compared to use of positron emission tomography (PET)-CT targeted EBUS alone [91]. However, the role of systematic endoscopic nodal staging is unclear in patients undergoing CRT for inoperable, multi-level N2 disease. Similarly to treatment planning for early-stage disease, a respiration-correlated 4D-CT scan is desirable for radiotherapy planning in locally advanced NSCLC, as it allows for target and normal organ motion to be considered [4]. In addition, information from co-registered diagnostic PET-CT scans can improve target delineation, and the safety and efficacy of PET-based target definition has been demonstrated in a recent prospective study [92]. Limiting the target volume to the primary tumour and involved nodes, or suspected lymph node metastases, is referred to as involved-field radiotherapy. Involved-field radiotherapy represents the current standard of care in both European [4] and North American guidelines [93]. This underlines the importance of incorporating all information gathered from endoscopic nodal staging and imaging studies into the radiotherapy planning process. Use of larger, so-called “elective” nodal fields has been associated
with worse survival [93], a finding which may be due to the risk of radiation-induced cardiac mortality [94] and immunosuppression [95].

In patients with a stage III NSCLC, CRT results in a 5–15% likelihood of symptomatic radiation pneumonitis (table 1) [88, 96, 98, 101]. In the PACIFIC trial, any grade of (radiation) pneumonitis occurred in 33.9% and 24.8%, with grade 3–4 pneumonitis reported in 3.4% and 2.6% of the durvalumab and placebo groups, respectively [14, 15]. This emphasises the need for careful attention to identifying high-risk patients, for whom optimal radiotherapy delivery techniques [4] and supportive care during and after combined modality treatment are essential [102]. Optimising radiotherapy planning parameters, such as the mean lung dose or the lung volume receiving a certain threshold dose (e.g. 20 Gy), can minimise the risk of radiation pneumonitis [4]. Doses to the heart and the oesophagus should be kept as low as possible, although no clear safe dose thresholds have been identified. Importantly, the introduction of new systemic treatments may potentially change the spectrum and frequency of adverse events observed during CRT (figure 4). Although the PACIFIC trial reported no significant increase in high-grade pneumonitis with durvalumab, a 14% incidence of grade 3 radiation pneumonitis was reported in real-world data from a Korean centre, where more than half of the patients receiving durvalumab consolidation did not meet the inclusion criteria of the PACIFIC study [103]. When nivolumab, a programmed cell death protein 1 inhibitor, was administered during CRT in stage III NSCLC (NICOLAS trial), any grade pneumonitis was reported in 42.5% of patients, with 10.4% experiencing grade 3 pneumonitis [100]. Further improvements in radiation delivery, and improved patient selection, are areas of investigation in patients undergoing CRT with immune checkpoint inhibitors.

The severity of radiation-induced oesophagitis can be decreased by using appropriate dose fractionation schemes, and with IMRT techniques, which can allow for oesophagus-sparing delivery [104]. Limiting oesophagitis, and other acute toxicities, may also allow for the timely initiation of durvalumab consolidation, as an unplanned subset analysis of the PACIFIC trial suggested an improvement in progression-free survival (PFS) if durvalumab was initiated within 2 weeks after completion of CRT [14, 15].

Despite the advances in treatment of stage III NSCLC, improvements in population survival rates are less apparent for this subgroup of patients. Large variations exist in real-world treatment patterns for patients with stage III NSCLC, especially for the elderly. For example, only 11% of stage III NSCLC patients in England received chemotherapy and radiotherapy, with sequential CRT delivered almost twice as often as concurrent CRT. Furthermore, only 33% of all patients with stage III disease, of whom one-third did not receive active treatment, survived for >1 year [105]. The complexity is further increased by a variability in expert opinions on best practices, particularly in N2 disease [106]. A population-based study in the Netherlands and Belgium showed that older age and higher N-stage were associated with the choice for sequential CRT [107]. A National Cancer Database (NCDB) study reported that sequential CRT was superior to concurrent CRT in elderly patients, suggesting a need for better tailoring of treatments in the elderly [108].

### TABLE 1 Incidence of oesophagitis and pneumonitis observed in selected recent trials

| Toxicity: CTCAE | Pneumonitis % | Oesophagitis % |
|-----------------|---------------|---------------|
|                 | Any grade     | Grade 3–4     | Any grade | Grade 3–4 |
| PROCLAIM: standard therapy [96] | 10.7 | 2.6 | 50.7 | 20.6 |
| RTOG 0617: 60 Gy [97] | Acute 11.1; late 9.8 | Acute 5.5; late 1.2 | 45.1 | 6.9 |
| RTOG 0617: 74 Gy [97] | Acute 14.5; late 3.9 | Acute 3.4; late 1.1 | 54.1 | 17 |
| KCSG-LU05-04: CRT phase [98] | 2.4 | 0 | 79.3 | 9.5 |
| CONVERT: LD-SCLC, once daily [99] | Acute 22.4; late 32.6 | Acute 1.6; late 2.6 | Acute 74; late 18.5 | Acute 19.1; late 1.7 |
| CONVERT: LD-SCLC, twice daily [99] | Acute 22; late 31 | Acute 1.6; late 2.4 | Acute 81.1; late 11.7 | Acute 18.5; late 0 |
| PACIFIC: durvalumab [15] | 33.9 | 3.4 | 27.5 | 6.5 |
| PACIFIC: placebo [15] | 24.8 | 2.6 |
| NICOLAS: nivolumab [100] | 42.5 | 10.4 |

CTCAE: Common Terminology Criteria for Adverse Events; CRT: chemoradiotherapy; LD-SCLC: limited disease small cell lung cancer.
The growing use of improved radiation treatment planning and delivery techniques, such as IMRT and VMAT, has not been shown to adversely impact survival in population-based studies of stage III NSCLC, suggesting that use of these techniques should be encouraged due to their ability to spare normal organs [109]. Improved radiation delivery is also relevant because survivors remain at risk for developing second primary lung cancer, which occurs at a rate of 2.9% per patient per year after CRT [110]. Although proton delivery may potentially reduce low-dose exposure of uninvolved lung tissue, a prospective trial using Bayesian adaptive randomisation of proton therapy versus conventional IMRT did not show any improvements in the rates of radiation pneumonitis or local failure with proton therapy [111]. Ongoing studies, such as the RTOG 1308 trial, are using improved proton delivery approaches to explore the role of this approach in decreasing doses to organs at risk, such as the heart.

The role of post-operative radiotherapy (PORT) in patients with a resected stage III NSCLC had been controversial, as a meta-analysis in the late 1990s reported that PORT was detrimental to survival in patients with resected NSCLC pN0 and pN1 [112]. There had been uncertainty regarding the utility of PORT in patients with pN2 disease, especially as subsequent analyses from the SEER and NCDB databases had suggested that PORT could improve overall survival in patients with N2 disease [113–115]. LungART was a well-powered, prospective trial in this patient group which randomised 501 patients with completely resected stage IIIA N2 NSCLC to either PORT (54 Gy in 27–30 fractions) or no PORT [116]. The study reported no significant differences in the primary end-point of disease-free survival. In addition, the rates of grade 3–4 cardiopulmonary toxicity were more than doubled with PORT. It can therefore be concluded that PORT is not beneficial for most patients with a completely resected stage IIIA N2 NSCLC, and it is doubtful if improvements in radiotherapy delivery will be sufficient as to improve on these outcomes.

In the treatment of limited disease small cell lung cancer (SCLC), concurrent CRT is the standard of care, and radiotherapy planning principles are similar to those discussed for stage III NSCLC. Concurrent CRT
with twice-daily radiotherapy to a dose of 45 Gy was established as a standard treatment based on a randomised trial that revealed superior outcomes compared to once-daily radiotherapy to 45 Gy in limited disease-SCLC [117]. However, not all centres perform twice-daily radiotherapy, as this can be a logistical challenge for departments and patients. The recent CONVERT trial compared twice-daily radiotherapy to 45 Gy as a standard of care, to once-daily radiotherapy to 66 Gy, in a randomised superiority trial. No major differences were seen in survival or toxicity between twice-daily and once-daily radiotherapy [99]. Although twice-daily radiotherapy to 45 Gy is a current standard of care, a recent randomised phase II trial suggested that survival was superior with twice-daily thoracic radiotherapy to 60 Gy, as opposed to 45 Gy [118]. However, the limitations of the phase II trial design mean that a confirmatory phase III trial must be performed to establish the superiority of a higher dose twice-daily radiotherapy scheme [119]. The ongoing ADRIATIC trial is evaluating the use of consolidation immunotherapy in patients with limited disease-SCLC who do not progress after concurrent CRT [120], although a similar study, STIMULI, recently reported no improvement of either PFS or overall survival with such an approach [121].

Chemotherapy is the treatment backbone for patients presenting with extensive disease SCLC, and two recent trials have revealed that the addition of immune checkpoint inhibitors to chemotherapy improves overall survival in these patients [122, 123]. The potential role of consolidative radiotherapy following first-line chemotherapy was previously investigated [124–126]. The CREST trial randomised patients with extensive disease-SCLC who responded to chemotherapy to either thoracic radiotherapy (30 Gy in 10 fractions) or no thoracic radiotherapy. CREST failed to meet its primary end-point of improved overall survival at 1 year, although thoracic radiotherapy increased 2-year overall survival rates from 3% to 13% [124]. As the trials evaluating immunotherapy with first-line chemotherapy did not permit consolidative thoracic radiotherapy [122, 123], the role of thoracic radiotherapy in the era of chemoimmunotherapy is the subject of an ongoing study [127].

**SABR for oligometastatic disease**

SABR is increasingly used for the treatment of patients presenting with up to five metastases, or so-called oligometastatic disease. In patients with limited metastatic NSCLC who do not progress after first-line systemic therapy, local consolidative therapy can improve PFS compared to standard treatment [16, 17]. The randomised SABR-COMET trial demonstrated that SABR delivered to one to five metastatic lesions can improve long-term survival compared to standard of care in a mixed cohort of patients presenting with different primary tumours [18, 20]. Nearly half of all metastases treated in the SABR-COMET trial were located in the lung, and lung SABR accounted for two of three fatal toxicities observed in the trial [18, 20]. Additional drivers of the need for improving SABR delivery are the suboptimal local control rates observed for some patients, with an overall long-term local control rate of only 63% for metastases in the SABR-COMET trial [20].

SABR has also been used for the treatment of isolated thoracic nodal recurrences [128, 129]. The results of ongoing trials, such as SABR-COMET-10 [78], may lead to a broadening of the indication for patients to undergo SABR for multiple lesions. Improved SABR delivery techniques will be required in such patients in order to reduce lung doses. Scheduling SABR treatments to multiple sites between cycles of systemic therapy will be challenging, and this may be facilitated by delivering shorter treatments. The safety, efficacy and cost-effectiveness of single-fraction SABR for oligometastatic patients with one to three lung metastases are being compared to a four-fraction SABR regimen in a randomised phase II trial, which has completed accrual [130].

Brain metastases are a major cause of morbidity and mortality in lung cancer. The management of these patients has become complex due to differences in prognosis based on factors such as the patient’s performance status, number of brain metastases, extracranial disease and the presence of targetable (e.g., EGFR or ALK) gene alterations [131]. There is a role for whole brain radiotherapy (WBRT) to provide symptom relief in patients with brain metastases, although WBRT provides little additional benefit for quality of life or survival in patients with brain metastases from NSCLC who are unsuitable for surgical resection or stereotactic radiotherapy [132]. In contrast to WBRT, single-session radiosurgery has become established as an effective local treatment for patients with up to 10 brain metastases [133, 134]. Furthermore, as modern magnetic resonance imaging allows for the detection of very small brain metastases, regular imaging allows for repeated stereotactic irradiation of new lesions, a concept which is currently being evaluated in a prospective trial [135]. For patients with SCLC, administration of prophylactic cranial irradiation following chemoradiation in limited disease-SCLC is considered standard due to the high risk of central nervous system (CNS) relapse [136]. Due to the risk of disseminated CNS progression, WBRT has traditionally been recommended in patients with brain metastases from SCLC, although this paradigm has been challenged by a recent cohort study suggesting that similar overall survival outcomes can be achieved with first-line...
radiosurgery [137]. First-line brain radiosurgery for patients with SCLC and up to 10 brain metastases is now being evaluated in a prospective randomised trial [138].

In a broader context, the role of radiation therapy in metastatic disease is being transformed by rapid changes in systemic treatments. In particular, the synergy of radiotherapy and immunotherapy remains an active area of research [139]. Both preclinical and clinical studies are aiming to better understand the complex immunomodulatory effects of radiation, and how to best use its potential to induce a systemic antitumour immune response [48]. In a prospective clinical trial in patients with progressive NSCLC where SABR was delivered to a single metastatic lesion prior to pembrolizumab (200 mg·kg\(^{-1}\) every 3 weeks) versus pembrolizumab alone, a doubling of overall response rates was observed, although the results did not meet the study’s prespecified end-point criteria for meaningful clinical benefit [19].

**Conclusion**

Recent advances in radiation therapy have contributed to the improvements observed in the survival of patients presenting with a lung cancer. Clear examples of this are the decreasing rates of non-treatment in early-stage lung cancer in population studies and the survival gains observed in trials incorporating immunotherapy following CRT in locally advanced NSCLC. In many parts of the world, the lack of patient access to these newer techniques remains an impediment. Similarly, as treatment options become more complex, such as for oligometastatic lung cancer, the role of the multidisciplinary tumour board in selecting appropriate strategies will be paramount.

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