Stereotactic body radiotherapy (SBRT) for central and ultracentral node-negative lung tumors

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Abstract: Advancements in imaging and radiotherapy (RT) techniques have allowed for remarkably precise delivery of high radiation dose per treatment fraction to intrathoracic targets. As a non-invasive therapeutic modality (compared to surgery), stereotactic body radiotherapy (SBRT) is an attractive option for patients with early-stage non-small cell lung cancers and oligometastases, especially for older patients with significant comorbidities and pre-existing pulmonary dysfunction. However, the outcomes and side effect profile of SBRT are highly dependent on tumor location, especially if the tumor is located centrally (within 2 cm of the proximal bronchial tree [PBT]) or ultracentrally (touching or within 1 cm of the mediastinum, esophagus, and PBT). In this focused review, we will examine the contemporary practice and principles of using hypofractionated RT or SBRT for central and ultracentral thoracic tumors. We will identify future directions on how this practice may be incorporated into the increasingly complicated modern paradigm of lung cancer treatments which now include immunotherapy along with proton beam radiotherapy.

Keywords: Stereotactic body radiotherapy (SBRT); SABR; lung cancer; central; ultracentral; local control; toxicities; review

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Introduction

In the modern era, stereotactic body radiotherapy (SBRT) is a powerful tool that can deliver ablative, tumoricidal radiation dose over a convenient schedule of typically 3 to 5 treatments. SBRT is commonly applied for thoracic malignancies, and is the standard of care for patients with early-stage, inoperable non-small cell lung cancer (1,2). The delivery of SBRT requires significant input from medical physics and dosimetry: it necessitates accurate tumoral and organ structure mapping with tight margins (stereotaxy), sophisticated intensity modulated radiotherapy techniques to spare normal organs and to target the tumor, the ability to deliver a higher dose rate of radiation by a linear accelerator, and also sophisticated imaging (image-guided radiotherapy, IGRT) and respiratory motion management techniques (3).

For thoracic targets, including primary lung cancers and metastases, the anticipated side effects (or toxicities) of SBRT are most dependent on the location of the tumor, namely peripheral versus central location (4) (Figure 1). In the radiation oncology literature, we are now separating lesions into central and ultracentral locations, as an ultracentral location may confer a higher risk of toxicity. In this focused review, we will examine the current understanding and clinical pearls of management of central and ultracentral thoracic tumors. We will also point out the existing challenges and controversies while highlighting...
potential improvements in radiation techniques (such as proton beam radiotherapy) and their relation to systemic therapy advances including immunotherapy for lung cancer.

**Definition of central and ultracentral tumors**

The radiation oncology definition of a central tumor is a tumor that lies within or is touching the zone of a uniform, 2-cm expansion around the proximal bronchial tree (PBT) or immediately adjacent to the pleura covering the mediastinum or pericardium (4,5) (Figures 1,2). The nuance of an ultracentral tumor is that the margin is narrowed to a 1 cm expansion of the PBT, and the lesion immediately touches or invades one of the organs at-risk such as the mediastinum, trachea, bronchus, or esophagus (6) (Figure 3).

In practice, the PBT structure is contoured on axial CT slices averaged from a 10-phase 4D-CT scan, and it includes the distal 2 cm of the trachea, the carina, the bilateral (left and right) mainstem bronchi, the bilateral upper lobe bronchi, the bronchus intermedius leading to the right middle lobe bronchus, the bilateral lower lobe bronchi, and the lingular bronchus (a left lung structure). This structure is then expanded isometrically by 1 and 2 cm, to be visually compared and verified with the tumor volumes that are drawn by the treating radiation oncologist. It is important
to note that, any part of the tumoral volume overlapping or touching this expanded PBT structure (or mediastinum) would qualify as a central or ultracentral tumor, according to the definitions noted above. This is of particular importance during SBRT dosimetric planning, as it impacts radiation dosing. For example, a peripheral tumor (i.e., not central or ultracentral) can be treated with much higher doses per fraction and over a shorter radiation course, e.g., 54 Gray (Gy) in 3 fractions or 48 Gy in 4 fractions (2,7) (Figure 4). Additional details on dose and fractionation for central and ultracentral lesions are discussed below.

**Workup and treatment options**

As with any primary lung neoplasm, staging is vital for determining the paradigm of treatment. By definition, central and ultracentral lesions would likely be T2NX and above disease given that T2 disease already encompasses lesions that invade the main bronchus and extend to the hilum (8,9).

As per the National Comprehensive Cancer Network (NCCN) guidelines, one would begin with computerized tomography (CT) chest with and without IV contrast, biopsy, positron emission tomography-computed tomography (PET-CT), surgical mediastinal staging (mediastinoscopy or endobronchial ultrasound/EBUS), and pulmonary function tests. Mediastinal node staging is critical as there can be occult metastases to N1 and N2 nodes in central and ultracentral lesions. Endobronchial ultrasound with biopsy of suspicious/enlarged nodes should be strongly considered (10-12). Brain magnetic resonance imaging (MRI) or head CT with contrast should be considered as well; especially for stage II and above non-small cell histologies, all histologically confirmed small cell lung cancer (13,14) or any patient who may be symptomatic.

The nuance of central and ultracentral lesions lies in determining whether definitive surgery or radiotherapy/chemoradiation are potential options. The standard of care for resection is lobectomy of the involved lobe and mediastinal node dissection/sampling. The decision for resection must occur in a multidisciplinary setting; particularly with ultracentral lesions, as surgery may require a pneumonectomy which the patient may not tolerate depending on performance status and pulmonary function at diagnosis, or the patient may refuse.

For lesions whose radiation volume will overlap esophagus, main bronchus, great vessels, or trachea, part of the workup includes ruling out any direct invasion of these organs. A CT angiogram or MR angiogram can be very useful for identifying direct vascular invasion. Direct visualization of the trachea and bronchial tree via endoscopy is also recommended to identify any endoluminal studding which can affect both the delineation of the gross tumor volume as well as the clinical target volume. Upper endoscopy can also determine if there is any direct invasion of the esophagus. For cases where the pericardium-tumor interface is compromised, MR of the heart can be very informative for early cardiac muscle invasion. Knowing the extent of disease in advance informs risk, especially if one is considering stereotactic radiation as the risk of fistula or fatal hemorrhage would be very high in the scenario where there is direct tumor invasion (15-18).

There is an increasing body of literature now examining late cardiac effects of radiation and this is especially pertinent to central and ultracentral lesions where the heart dose is particularly high (19-24). Most of the literature comes from recent publications on conventionally delivered radiation in locally advanced lung cancer in which excess mean heart dose (>11–20 Gy) resulted in higher grade 3 cardiac events defined as: acute coronary syndrome, new congestive heart failure, cardiac arrest, new onset arrhythmia, valvular disease, and new pericardial effusion (19-24). In the randomized dose escalation radiation trial for locally advanced lung cancer, RTOG 0617, there was worse overall survival in the dose escalation group that was attributable to higher mean heart dose from radiotherapy.
(25,26). In the era of immunotherapy where survival from Stage III lung cancer has doubled and in patients who are receiving SBRT for localized node-negative disease, it may be a consideration to optimize cardiac status prior to and after delivery of radiation. The long-term cardiac effects of SBRT is less well characterized as there has been no dose-volume correlate that is predictive of late cardiac events. This is likely due to shorter follow up data on SBRT patients and treatment of much smaller tumors and smaller cardiac/pericardial volumes in the SBRT setting (27).

**Current practice for central tumors**

The final results of the NRG Oncology/RTOG 0813 dose-escalation trial (5) were recently published. The pivotal phase II trial tested the maximally tolerated dose (MTD) of SBRT for centrally located tumors, over a 5-fraction schedule with radiation delivered every other day (dose per fraction ranged from 10.0 Gy per day to the MTD of 12.0 Gy). The grade 3 and above dose-limiting toxicities (DLT), defined as radiation pneumonitis, fatal pulmonary hemorrhage, hypoxia, new pleural effusion and death, were found to be 7.2%. The higher-dose arms (11.5 and 12.0 Gy per fractions) achieved local control rates of 89% at 2 years. However, among patients who were evaluable for DLT, zero patients developed a severe toxicity at the 10.0 Gy per fraction level, compared to the higher dose groups at 10.5 Gy per fraction or higher. These higher dose cohorts developed at least 1 DLT which was grade 3 or above. As a result, most radiation oncologists would recommend that the central lung tumoral dose be kept at 10.0 Gy per fraction, for 5 fractions (50 Gy total) (1,28). Other acceptable dose regimens include, 9.0 Gy x5 fractions (45 Gy total), 7.5 Gy x8 fractions (60 Gy total), and 12.5 Gy x4 fractions (50 Gy total) (29-33).

Several studies, including RTOG 0813 which is discussed above, have evaluated patient outcomes after SBRT for central lung tumors. The 2-year local control after central and ultracentral lung SBRT are reported to range from 79% to 90% with 2- and 3-year overall survival reported as 43% to 80% respectively (5,29,31,32,34-38). These are excellent outcomes despite the patients’ significant pre-existing pulmonary comorbidities and smoking history (5,29,31,32,34-38). However, there still exists a small percentage (5% to 7.5% or less) of possible treatment-related deaths (grade 5 toxicities) associated with SBRT for central lung tumors (5,29,31,32,34-38). Most of these series included ultracentral lesions, and two studies focused particularly on ultracentral lesions (35,38). Potentially fatal side effects may include radiation-related pneumonitis, central airway necrosis (39), tracheoesophageal fistula, or bronchopulmonary hemorrhage (5,29-33).

**Current practice for ultracentral lung tumors**

Most ultracentral lung tumors are not resectable and primary therapy usually involves radiotherapy. For unresectable locally advanced non-small cell lung cancer and limited stage small cell lung cancer, the standard of care would be definitive chemoradiation. There are practice considerations that could limit the dose to the heart, lung, and esophagus to try to mitigate acute and late toxicities from definitive chemoradiation.

For conventionally fractionated (1.8–2.0 Gy/fraction) radiotherapy, one consideration is proton delivery for bulky tumors where the mean lung dose or V20 are exceeded by an intensity modulated radiation therapy (IMRT) photon plan (40,41). For tumors that do not about the esophagus, one could try to exclude dose to the esophagus to minimize mean esophageal dose thereby reducing the risk of both acute esophagitis and late stricture. Similarly, there have been multiple recently published data on limiting the mean heart dose using either an IMRT or proton technique (20,22). In the end, while the radiation plan attempts to minimize toxicities, it is also important to balance the implications of recurrence with undercoverage of the target.

For small lesions (T1 and small T2) that are ultracentral, SBRT has been utilized safely and effectively (38,42-44). For patients with node negative disease but T3 and above tumors who are not candidates for chemoradiation, some investigators have treated these with hypofractionated radiotherapy in an attempt to achieve local control or SBRT type doses (6,35,45). However, the standard of care for the treatment of ultracentral tumors continues to be a subject of heated debate among radiation oncologists.

**Controversies in central and ultracentral lung treatments**

The greatest controversy for both central and ultracentral lung radiation is the incidence of toxicity and the safest dose regimen for radiotherapy delivery. There have been multiple single institution studies examining the outcomes of SBRT delivered in 8 fractions or less to ultracentral lesions (29-33). Memorial Sloan Kettering noted that tumors that were located closest to the proximal bronchial tree had
the highest risk of treatment-related death attributable to radiation-associated airway necrosis (18). The VU Center in Europe has also published their ultracentral SBRT series with a very high rate (15%) of fatal pulmonary hemorrhage using 60 Gy in 12 fractions, a relatively modest hypofractionated regimen (35). Similarly, a Chinese study in which doses as low as 35 Gy in 5 fractions delivered for ultracentral lesions still showed a 3.9% incidence of treatment-related death in 51 patients and a 9.8% risk of grade 3 or higher toxicity (38).

There is limited prospective data as the original definition of central lung lesions were those within 2 cm of the proximal bronchial tree. As discussed above, NRG/RTOG 0813 was a prospective central lung SBRT dose escalation trial that showed no maximum tolerated toxicity even to a dose of 60 Gy/5 fractions although analysis of ultracentral vs central location is still pending (5). Ongoing radiation studies are attempting to examine the optimal dose schema including the Canadian SUNSET trial that starts at 60 Gy in 8 fractions (44) and the EORTC LungTech study which also employs the 60 Gy in 8 fraction regimen (46). There are also trials examining multimodality therapy for ultracentral lesions. One such trial was published by UCLA in which thermal ablation and SBRT were combined for lesions abutting esophagus and other critical structures (47).

**Future directions**

SBRT will remain an important tool for patients with central or ultracentral lung tumors. Often, it may be the only locoregional treatment option if the patients are moderately to severely compromised from a cardiopulmonary prospective. The 5-year survival for localized non-small cell lung cancer (NSCLC) patients with untreated tumors is 7%, with a 11-month median survival (48). While the benefits of giving SBRT treatment frequently outweighs the risks in most clinical scenarios, the consistently reported grade 5 toxicities in the 2–7% range are a real concern and further investigation is needed to minimize fatalities attributable to SBRT in this setting.

Two particular areas of future research are worth noting: First, there has been an increase in the utilization of protons in lung cancer, mainly in locally advanced cases when chemoradiation is required (41,49,50). Due to a physical phenomenon called the Bragg peak effect, a proton-based radiotherapy plan can deliver full radiation dose to an intrathoracic target with radiation “stopping” and dissipating entirely beyond the target’s range, resulting in virtually zero radiation dose exiting the body. This often translates to lower lung and cardiac doses. As proton technology continues to advance (mainly needing to provide a higher dose rate, and also IGRT improvement such as daily cone-beam CT’s), stereotactic body proton beam therapy (SBPT) will become a reality and may offer an advantage beyond what typical photon SBRT can achieve today, especially in the setting of ultracentral and central tumors.

Secondly, in the immunotherapy era, more investigators are looking at strategies in combining checkpoint inhibitors and SBRT safely, as the ablative dose effect of SBRT can also release tumoral antigens directly and augment tumor response to immunotherapy. Data from initial clinical trials combining radiotherapy and immunotherapy show a higher rate of overall response to immunotherapy compared to immunotherapy alone (50% vs. 20%, P<0.1) in patients with advanced NSCLC (51). More clinical trials are currently ongoing.

**Conclusions**

In the current review, we have summarized the contemporary practice and principles of using hypofractionated RT (more than 5 fractions) or SBRT for central and ultracentral thoracic tumors, and also discussed the relevant clinical pearls and controversies associated with these techniques. As more studies combining SBRT with systemic therapy are published, the use of SBRT should see a surge; ultimately improving care for our patients in the next 5 to 10 years.

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