



A single institutional experience with central lung stereotactic body radiation therapy demonstrating encouraging results with increased inter-fraction time

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Background: Stereotactic body radiation therapy (SBRT) is an effective treatment modality for non-small cell lung cancer (NSCLC); however, there are concerns regarding potential toxicity for centrally located tumors.

Methods: This retrospective study considered patients with SBRT for central lung NSCLC (defined as a tumor within 2 cm of any mediastinal critical structure). The institutional protocol was that patients with central tumors received SBRT less frequently than daily—generally once or twice weekly.

Results: A total of 115 patients with 148 lesions were treated with SBRT to a median 45 [5–60] Gy in 4 [1–5] fractions over a median 5.3 [0–18] days. Many patients treated with this method presented with advanced disease: 58 treatments involved nodal targets, and 42 had stage 3 disease. 52% of patients had chronic obstructive pulmonary disease (COPD), and only 49% had a biopsy, often due to concerns regarding other medical comorbidities. Rates of prior chemotherapy, thoracic surgery, and thoracic radiotherapy were 32%, 21%, and 49%, respectively. Via the Kaplan-Meier method, 2-year overall survival was 65%, and 2-year local control was 77%. Two-year local-progression free survival was 53%, and 2-year progression-survival was 48%. Treatments for stage 3 disease had an impressive 82% 2-year local control that was comparable to early stage treatments. Patients with stage 3 disease had a 2-year overall survival of 59%, which trended towards decreased overall survival compared to early stage patients. There were 13 grade 1 (9%) and 14 grade 2 (9%) toxicities. There were no reported grade ≥ 3 acute or late toxicities and only 3 cases of pneumonitis.

Conclusions: Our series demonstrates encouraging local control with low rates of toxicity for central lung SBRT, including many stage 3 patients. This may be the result of the relatively large inter-fraction interval. This interval may allow for greater tumor effects (such as reoxygenation) and improved tolerance from normal tissues.

Keywords: Dose fractionation; non-small cell lung cancer (NSCLC); patient selection; stereotactic body radiation therapy (SBRT); treatment outcome

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Introduction

Stereotactic body radiation therapy (SBRT), also known as stereotactic ablative radiotherapy (SABR), delivers high doses of radiation in relatively few treatment fractions. Studies in non-small cell lung cancer (NSCLC) have demonstrated high rates of local control and low toxicity, allowing for its use in patients with significant medical comorbidities (1). As the refinement of patient selection continues, patients who are surgical candidates have also been considered, and population studies have associated the increasing utility of SBRT for elderly patients and improved overall survival (2,3). Though SBRT has been favorably compared to conventional fractionation in early stage lung cancer, the dose-limiting toxicity, radiation pneumonitis, continues to affect patients after 9–28% of cases (4,5).

Centrally located NSCLC carries the potential for increased rates of toxicity, largely secondary to the proximity to key central structures. Though varied definitions exist, most analyses define a central tumor as a lesion within 2 cm in all directions of any mediastinal critical structure (6). RTOG 0813 found a 7.2% rate of dose-limiting toxicity (grade ≥ 3) with a maximum tolerated dose of 12.0 Gy/fraction, but other studies have demonstrated lower rates of toxicity with central lung SBRT (7,8). Two papers, including the prospective phase II trial from Bral *et al.*, noted a correlation between local control and tumor size, which could subsequently affect dose delivery and toxicity (9,10). These results have prompted some providers to increase the number of fractions for dose delivery up to 8 treatments (11). Notably, survival outcomes appear to be comparable between early stage NSCLC, regardless of central or peripheral location (12).

Further questions have arisen regarding “ultra-central” and hilar SBRT. “Ultra-central” tumors are generally defined as those with gross tumor volume (GTV) directly abutting the central airway. While some studies have shown encouraging outcomes, careful patient selection is recommended until large prospective trials are completed (13). A key concern emerges regarding prioritization of target coverage or organs at risk (14). Treatments for paramediastinal and hilar targets similarly challenge

physicians (15). Hilar radiotherapy has been discussed in only a few studies, the largest of which considered 40 patients with hilar/mediastinal SBRT, finding any grade ≥ 3 toxicity in 4 patients (10%) (16–18).

This study seeks to analyze a large patient cohort treated with SBRT for central lung NSCLC and present outcomes of overall survival, progression-free survival, and local-progression free survival. This group consists of many treatments with uncommon and potentially high-risk features, such as node-positive disease, unbiopsied tumors, recurrent cancers, and stage 3 malignancies, and the authors seek to demonstrate that increased inter-fraction time may reduce the risk of side effects, like the PATRIOT trial in prostate cancer SBRT (19,20). We present the following article in accordance with the STROBE reporting checklist (available at <http://dx.doi.org/10.21037/jtd-20-2659>).

Methods

Patients

Patients were considered if they received SBRT for a central lung tumor diagnosed as or empirically treated as NSCLC. Patients were treated from 2007–March 2018. A central lung cancer was defined as a tumor within 2 cm in all directions of any mediastinal critical structure. Exclusion criteria included a lack of clinical or imaging follow-up, a different primary tumor, or a peripheral location. Patients were not excluded from the study based on clinical stage, hilar or mediastinal location (grouped as “nodal” for the analysis), or a lack of biopsy. Patients were also not excluded from the study if they had a documented death but a lack of clinical or imaging follow-up. In such a case, the patient would be included in the survival analysis but not in the assessment of local control. Patient characteristics and outcomes were recorded via retrospective chart review, and this approach was exempt by the institutional review board due to the retrospective nature of the study and the establishment of a previous protocol for conducting retrospective studies. The initial patient set was developed to include all patients treated from 2007 through 2014 who met the above criteria. This resulted in an initial set of 105 lesions treated in 85 patients. A separate review of all

treated SBRT cases from 2007 through 2018 was performed to additionally include all patients with treatments targeting the hilar or mediastinal regions. This resulted in the addition of 30 patients with 43 treated lesions to the dataset. This study was conducted in accordance with the Declaration of Helsinki (as revised in 2013). The study was exempt by the institutional review board and the informed consent, due to the retrospective nature of the study and the establishment of a previous protocol for conducting retrospective studies.

Radiation therapy

For treatment, patients were immobilized using a full-body vacuum bag system for position stabilization and consistency. Serial computed tomography (CT) scans were taken (free-breathing, inhale, exhale) for treatment planning purposes to assess the motion of the target during the breath cycle and generate an internal-target-volume (ITV) margin around the GTV. For patients with previous radiation therapy, treatment planning files were obtained and imported into the treatment planning system for evaluation of prior dose to organs-at-risk. Dose was prescribed to the planning target volume (PTV), which was defined as the ITV plus 3–5 millimeters of margin to account for uncertainties in imaging and localization. In general, a 3D conformal treatment planning approach with non-coplanar gantry angles was used to minimize dosimetric overlap of entrance and exit portals. SBRT was delivered using a 6 MV photon beam on a linear accelerator with a 2.5–4 mm width multi-leaf collimator for custom shaping of portals. On-board cone-beam CT (CBCT) with 4D capabilities was used prior to treatment and would be repeated during treatment as needed to correct any intra-fraction motion of the patient or target. Treatment was generally provided once or twice weekly, per institutional policy.

For early stage tumors, the most common treatment regimens included 60 Gy in 4 fractions [n=17, biologically effective dose (BED) =150 Gy], 50 Gy in 4 fractions (n=28, BED =112.5 Gy), and 50 Gy in 5 fractions (n=13, BED =100 Gy). 40 Gy in 4 fractions (n=12, BED =80 Gy) was commonly chosen in the setting of prior radiation therapy. Nodal targets often received lower doses, due to concern for potential toxicity. These most commonly included 28 Gy in 4 fractions (n=8, BED =47.6 Gy), 24 Gy in 4 fractions (n=8, BED =38.4), or 20 Gy in 4 fractions (n=7, BED =30 Gy). Other dose-fractionation schemes for nodal targets included

36 Gy in 4 fractions or 45 Gy in 3 fractions. In 11 cases, prescription doses of at least 50 Gy were delivered. This was generally done in the setting of minimal prior treatment and treatments for the hilar lymph nodes (as opposed to other mediastinal targets). Lower dose regimens, such as 15–21 Gy in 3 fractions (n=6), were sometimes utilized, especially in the setting of reirradiation. Most treatments involved 3–5 fractions of therapy, with only two treatments involving one or two fractions. The one case in which only one treatment was offered was included because it contributed to the analysis of treatments involving prior thoracic surgery. Biologically effective dose (BED) was calculated using the formula $\text{dose} \times (1 + \text{dose per fraction}/\alpha \text{ beta ratio})$, and an alpha beta ratio of 10 Gy was used.

SBRT is not considered the standard of care for advanced or recurrent disease. Patients treated with such disease with SBRT in this study generally were considered to be high-risk candidates for whom concurrent chemoradiation to 60 Gy was considered to be a suboptimal choice. Many of these individuals had received prior treatment or declined these more standard treatment approaches, due to concerns regarding toxicity. Due to these constraints, relatively lower treatment doses were often used, as well. In the setting of these nodal treatments, dose constraints varied, as dictated by the impact of prior radiation therapy fields. Generally, however, a maximum point dose of 105% of the prescription dose was allowed to the esophagus and proximal tracheobronchial tree. Whenever possible, these maximal doses were kept to less than 30 Gy.

Key dosimetric characteristics of the treatment were recorded, including dose, fractions, days between fractions, and PTV. The use of concurrent chemotherapy was noted, where applicable. The reason for SBRT was recorded, as were any previous therapies: chemotherapy, thoracic surgery, and thoracic radiotherapy. Synchronous lesions were defined as two or more lesions present in a patient's lungs. This included two lesions within the same lobe, different lobes, or different lungs. The presence of synchronous lesions, as well as the target (nodal *vs.* non-nodal), were considered as potential predictive factors for local control in the multivariate analysis. Since many of these patients were treated prior to the emergence of data concerning the benefits of immunotherapy, this data was not recorded.

Patient outcomes

The principal outcomes of the study were local control,

overall survival, progression-free survival, and local-progression free survival. These were assessed via clinical and imaging follow-up, the latter of which was generally conducted 2–3 months after completion of SBRT. The imaging modality used for this follow-up was recorded, generally consisting of CT and positron emission tomography (PET) scans. Tumor recurrence was recorded as the date of the imaging which noted the recurrence, and it was designated as local, regional, or distant failure. Local failure was defined as tumor recurrence within the same lobe as the primary tumor or in the ipsilateral hilum. Regional failure was defined as tumor recurrence in the contralateral hilum or ipsilateral lung but a different lobe than the primary tumor. Other cases of recurrence were deemed to be distant failure. Local control was still evaluated even after the detection of regional or distant metastases. Similarly, regional control was still assessed even after the detection of distant metastases, as opposed to censoring at the time of distant failure. Overall survival was defined as the time between the last day of treatment and either the date of death (if known) or the last date of interaction with the medical system. Toxicity was recorded and graded according to the Common Terminology Criteria for Adverse Events v4.0. Toxicity was most commonly reported on subsequent follow-up visits with radiation oncology, but pulmonology follow-up also contributed to the toxicity outcomes.

Statistical methods

Statistical analysis involved univariate analysis, Cox proportional-hazards analysis, and the Kaplan-Meier method. A P value of 0.05 was considered to be the threshold for statistical significance. Univariate analyses involved exclusively two-tailed *t*-tests and served to compare binary variables. Cox proportional-hazards analysis demonstrated hazard ratios for the potential predictive factors. Though all independent variables were initially assessed, very low variance was detected for non-surgical candidacy for the overall survival analysis, so the reason for SBRT was eliminated from the subsequent hazard analysis for overall survival. The Kaplan-Meier method was used to portray time-dependent variables, and stratifications of the data were conducted according to the results of the initial statistical analyses. Treatments for stage 3 disease were analyzed as an important subgroup of the analysis, and analysis was conducted using the statistical methods previously described. Any patients or treatments with

missing data for a particular statistical analysis were not included in that analysis.

Results

Patient characteristics

A total of 115 unique patients with 148 treated lesions were included (*Table 1*). Patients started treatment at a median 78 (53.1–92.2) years of age, and 60 of the 115 patients had chronic obstructive pulmonary disease (COPD). Patients presented at a median 9.8 (0.4–233.0) months after primary diagnosis, and only 56 (49%) obtained biopsy prior to SBRT. 93% of treated lesions had controlled systemic disease, with a median longest dimension of 2 (1–6.7) cm. Lesions were treated at clinical stage 1 (46%), 2 (20%), 3 (28%), 4 (4%), and unknown (1%). 58 treatments involved nodal targets. Synchronous lesions were present in 31% of cases, and 75% had a pre-treatment PET scan, demonstrating a median pre-treatment standardized uptake value (SUV) of 4.5 (0–17.1).

Radiotherapy and prior treatments

Patients were treated with a median 45 [5–60] Gy in 4 [1–5] fractions. The median days between fractions was 5.3 [0–18], and the median overall treatment time was 21 days. Concurrent chemotherapy was given with only 3% of treatments. The median PTV of treatments was 25.7 (1.3–269.6) cc. SBRT was offered for primary cancer (39%), recurrence (57%), and medically inoperable status (5%). Many treatments involved lesions that were previously treated with chemotherapy (32%), thoracic surgery (21%), and thoracic radiotherapy (49%). Of these prior radiation treatments, the median radiation dose was 57 Gy in 5 fractions, and these treatments occurred at a median 15.6 months prior to SBRT.

Patient outcomes

Median overall survival, progression-free survival, and local-progression free survival were 20.0, 14.0, and 16.1 months, respectively. There were 28 instances of local failure (19%) at a median imaging follow-up of 13.2 months. Median clinical follow-up was 17.5 months. There were 20 cases of regional failure (14%), which occurred at a median 6.16 months. Of these, 11 occurred in patients with stage 3 disease, as opposed to only 3 and

Table 1 Patient demographics and treatment characteristics are considered

Variable	Number
Patient demographics	
Patients	115
Lesions	148
Median age (years)	78 (53.0–92)
Patients with COPD	60 (52%)
Median time from primary diagnosis to SBRT (months)	9.8 (0.4–233)
Patients with biopsy obtained	56 (49%)
Patients without biopsy obtained	59 (51%)
Squamous cell carcinoma	25 (17%)
Adenocarcinoma	32 (22%)
Other or unknown histology	91 (61%)
Controlled systemic disease	137 (93%)
Lesions with pre-treatment PET	111 (75%)
Median pre-treatment SUV	4.5 (0–17.1)
*Median longest dimension (cm)	2 (1–6.7)
Stage 1	68 (46%)
Stage 2	30 (20%)
Stage 3	42 (28%)
Stage 4	6 (4%)
Treatment history or characteristic	
Median dose (Gy)	45 (5.0–60)
Median fractions	4 (1.0–5)
Median BED (Gy)	100 (7.5–180)
Median days between fractions	5 (0–18)
Concurrent chemotherapy	4 (3%)
Median PTV (cc)	25.7 (1.28–69.6)
Reason for SBRT: primary cancer	57 (39%)
Reason for SBRT: recurrence	84 (57%)
Reason for SBRT: not surgical candidate	7 (5%)
Treatments with prior chemotherapy	48 (32%)
Treatments with prior surgery	31 (21%)
Treatments with prior thoracic radiotherapy	72 (49%)
Median prior radiotherapy dose (Gy)	57
Median prior radiotherapy fractions	5

Table 1 (continued)**Table 1** (continued)

Variable	Number
Median time from prior radiotherapy to SBRT (months)	15.6 (0.2–281)
Synchronous lesions	46 (31%)
Nodal target	58 (39%)
Central, non-nodal target	90 (61%)

*This includes nodes and primary disease. COPD, chronic obstructive pulmonary disease; SBRT, stereotactic body radiation therapy; PET, positron emission tomography; SUV, standardized uptake value; BED, biologically effective dose; PTV, planning target volume.

6 cases for stage 2 and stage 1 patients, respectively. 15 distant failures (10%) were noted at a median 9.3 months. Fifty lesions (34%) demonstrated some disease progression (local, regional, or distant). Using the Kaplan-Meier method, the 2-year overall survival was 65%. Two-year progression free survival and local-progression free survival were 48% and 53%, respectively. Two-year local control was 77% (Table 2).

Predictive factors

On univariate analysis, BED >100 Gy demonstrated a trend towards improved local control (P=0.10). Higher clinical stage trended towards increased disease progression (P=0.051), and prior chemotherapy was predictive of this outcome (P=0.017).

Cox proportional-hazards analysis was performed, showing that prior chemotherapy trended towards increased local failure (P=0.05), with a hazard ratio of 2.49 (0.98–6.31). Disease progression had multiple predictive factors (Figure 1). Synchronous lesions (P=0.03) and prior radiotherapy (P=0.03) were protective, with hazard ratios of 0.35 (0.13–0.92) and 0.25 (0.07–0.89), respectively. Prior chemotherapy (P=0.04) and prior thoracic surgery (P=0.05) demonstrated increased disease progression, with hazard ratios of 2.44 (1.02–5.82) and 2.36 (1.00–5.58). Finally, prior thoracic surgery predicted for decreased overall survival (P<0.005), with a hazard ratio of 4.39 (2.10–9.20). There was also a trend towards improved overall survival with obtaining a biopsy (P=0.08) while prior chemotherapy trended towards decreased overall survival (P=0.11).

Kaplan-Meier analysis was utilized to demonstrate local

Table 2 Key patient outcomes are tabulated

Variable	Number	Rate
2-year overall survival		65%
2-year progression-free survival		48%
2-year local-progression free survival		53%
2-year local control		77%
Local failures	28	19%
Median time to local failure (months)	15.2	
Regional failure	20	14%
Median time to regional failure (months)	6.2	
Distant failure	15	10%
Median time to distant failure (months)	9.3	
Any disease progression	50	34%
Grade 1 toxicity	13	9%
Grade 2 toxicity	14	9%
Grade ≥ 3 toxicity	0	0%
Median imaging follow-up (months)	13.2	
Median clinical follow-up (months)	17.5	

control, local-progression free survival, progression-free survival, and overall survival for the entire cohort (Figure 2). It was demonstrated that nodal targets had worse local-progression free survival than non-nodal targets ($P=0.05$). Patients without prior chemotherapy had improved progression-free survival ($P=0.11$) and trended towards improved overall survival ($P=0.05$). Similarly, patients without prior thoracic surgery had improved progression-free survival ($P<0.005$) and overall survival ($P<0.005$).

Stage 3 disease

Patients with stage 3 disease were also considered separately as part of the outcomes analysis to compare their outcomes to patients with early stage disease. Two patients had a clinical stage unavailable, and two patients (six treated lesions) were also not considered because they had stage 4 disease. A total of 111 patients with 140 treatments were, therefore, used for this subgroup analysis. The stage 3 group consisted of 27 patients and 42 treated lesions. Of these, only 1 treatment involved concurrent chemotherapy, but most treatments involved prior chemotherapy (55%), thoracic surgery (31%), or radiation (76%). Forty-three

percent of treatments involved patients with COPD. The 2-year local control for stage 3 disease was 82%, and the 2-year overall survival was 59%. There was no difference between stage 3 disease and early stage disease for 2-year local control (82% *vs.* 77%, respectively); however, stage 3 disease trended towards decreased overall survival and progression-free survival ($P=0.18$ and $P=0.32$, respectively) (Figure 3).

Toxicity

There were 13 total acute and late grade 1 toxicities reported (9%), in addition to 14 total grade 2 toxicities (9%). There were no reported grade ≥ 3 morbidities and no grade 5 toxicities. The most common grade 1 side effect was fatigue (7 cases); others included dyspnea, cough, shortness of breath, nausea, and esophagitis. The most common grade 2 toxicities were pneumonitis and shortness of breath (3 cases each). The other grade 2 morbidities involved fatigue, dyspnea, cough, pleural effusion, pain, and one rib fracture. Though attempts were made to risk stratify the patients for increased rates of toxicity based on various disease characteristics, the relatively low number of overall side effects prevented a rigorous analysis of this kind.

Discussion

Overall, this study demonstrated encouraging treatment outcomes in a moderately sized cohort of centrally-located NSCLC treated with SBRT. Despite a high rate of adverse risk factors in this data set, an impressive 2-year local control of 77% was reported. Patients were treated with an increased inter-fraction time, likely contributing to the 0% rate of grade ≥ 3 toxicity.

A six-year analysis of 102 primary lung cancers treated with once-weekly SBRT from Salazar *et al.* is perhaps the most comparable study to the present analysis (21). The study included 30 stage 3 cancers, and the authors documented 12%, 7%, and 0% rates of grade 1, 2, and ≥ 3 toxicity, respectively. These results are similar to those of the present analysis and point towards the potential utility of SBRT in stage 3 disease, especially in elderly patients or those who are poor candidates for combined chemotherapy and radiation. Further, it adds support to the hypothesis that increased inter-fraction time can help to minimize treatment toxicity. Rates of symptomatic radiation pneumonitis with SBRT have ranged from 9% to 28%, but here only 3 cases (2%) were found, despite high rates

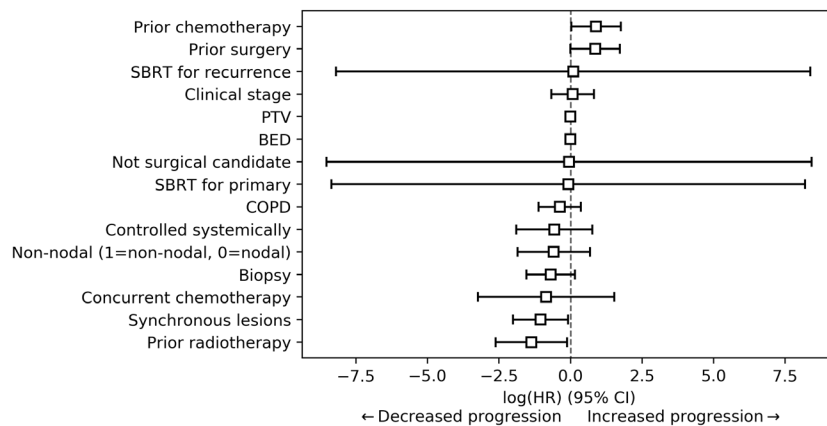


Figure 1 Cox proportional-hazards analysis demonstrates predictive factors for disease progression.

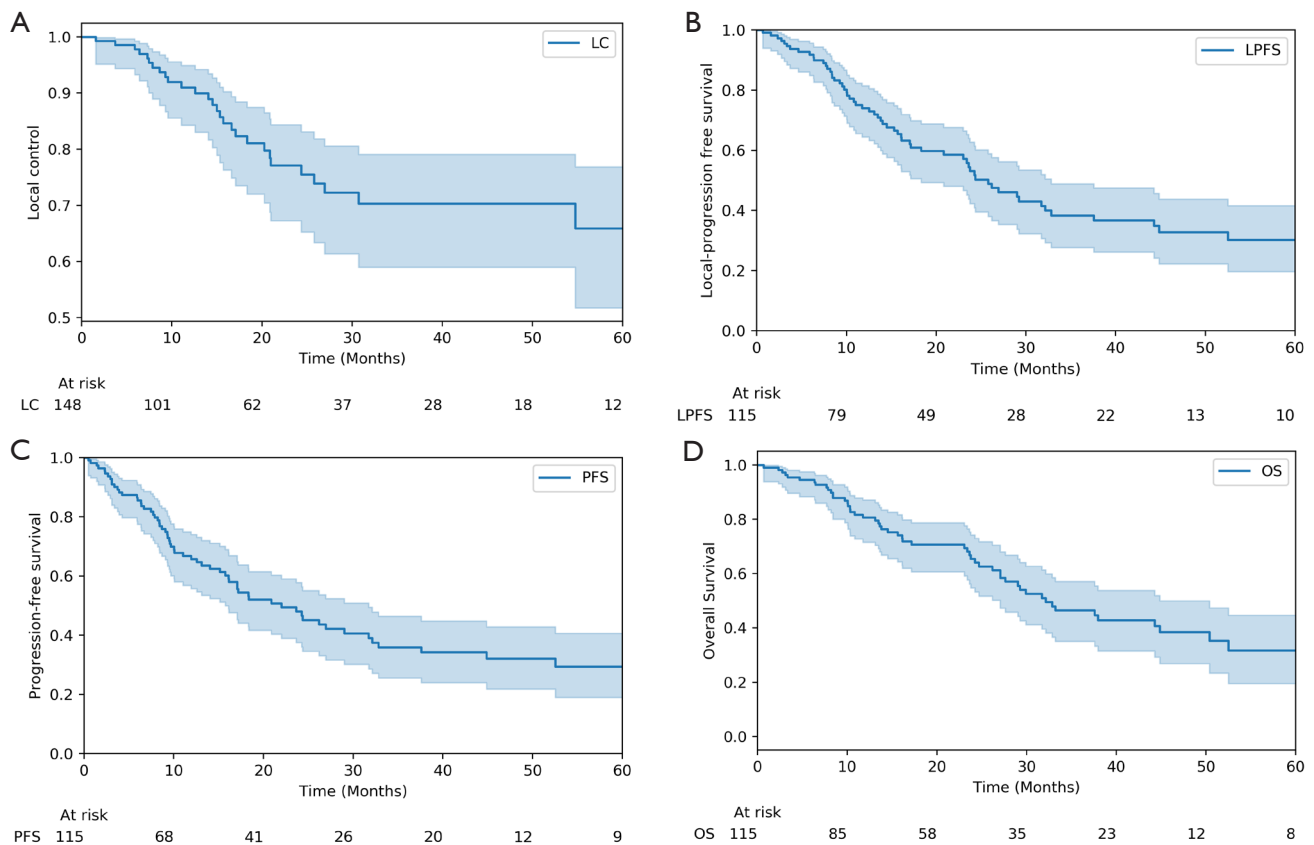


Figure 2 Kaplan-Meier analysis was utilized to demonstrate (A) local control, (B) local-progression free survival (LPFS), (C) progression-free survival (PFS), and (D) overall survival (OS).

of adverse risk factors (5,22). In addition, patients have commented on the flexibility of once-weekly treatments because weekly activities can be continued relatively uninterrupted by radiation therapy. This treatment strategy

may be particularly optimal for patients living relatively near the treatment facility while patients who traveled a long distance for treatment may prefer to complete therapy sooner.

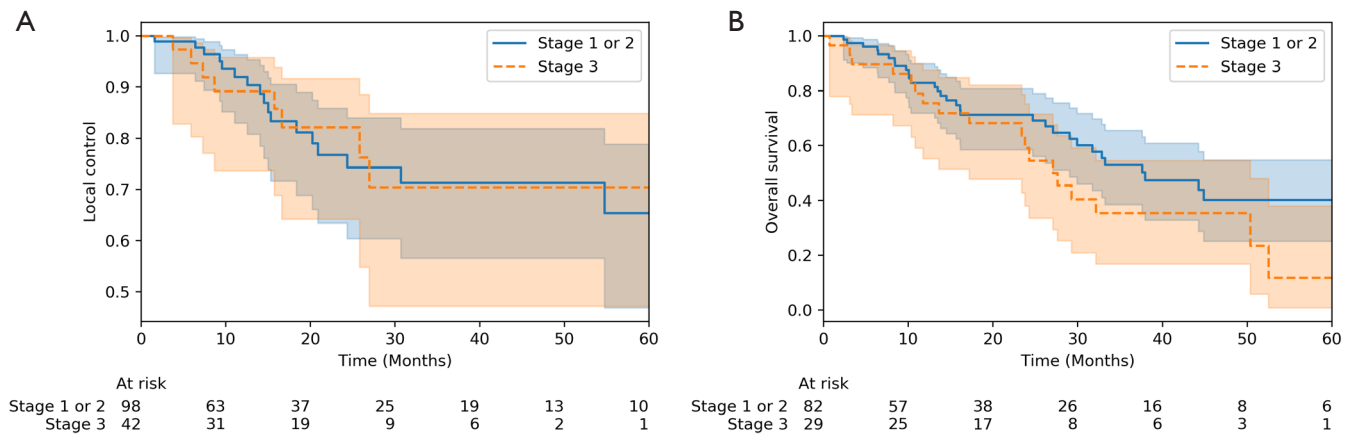


Figure 3 SBRT for stage 3 disease demonstrated comparable local control (LC) to treatments for stage 1 or 2 disease (A, $P=0.94$), but patients with stage 3 disease trended towards decreased overall survival (OS) (B, $P=0.18$). SBRT, stereotactic body radiation therapy.

This analysis consisted of a unique patient cohort with more advanced disease and adverse risk factors than most lung SBRT studies. Generally, most lung SBRT has focused on early stage NSCLC or oligometastatic disease, but this study was quite novel in that 42 treatments for stage ≥ 3 NSCLC were included (1,23,24). SBRT has been used as a method to boost residual disease in locally advanced NSCLC, but rarely has it been considered for definitive therapy (19). Despite trends towards decreased overall and progression-free survival for stage 3 disease, this study demonstrated comparable local control rates for stage 3 disease compared to early stage NSCLC. Such a result is hypothesis-generating and presents the possibility of treating stage 3 disease with SBRT for palliation or local control.

Most treatments for stage 3 disease were the result of N2 or N3 disease, rather than T3N1 or T4 tumors, and most dose-fractionation schemes involved the delivery of 20–30 Gy in 4 fractions. These treatments resulted in an encouraging side effect profile that was not significantly different than that for other targets. It is likely that the increased inter-fraction time (and potentially the lower treatment doses) used in this analysis contributed to the 0% rate of grade ≥ 3 toxicity. These outcomes for locally-advanced NSCLC indicate that SBRT may prove beneficial for carefully selected patients (particularly with N2 or N3 disease, rather than large tumor size). Further study is recommended to assess for the role of increased inter-fraction time in this setting as a potential tool to reduce toxicity in high-risk patients.

The underlying radiobiology of SBRT involves key differences from that of conventional radiotherapy, and it remains unclear whether classic radiobiologic modeling and the linear-quadratic model appropriately describe cell death in this clinical scenario (25,26). Preclinical and modeling studies have indicated that hypoxia may have a particularly negative impact on tumor control in SBRT, and the time between fractions in conventional radiotherapy allows for improved reoxygenation (26). Another study of SBRT in murine tumors indicated that the surviving hypoxic cells decreased over several days post irradiation, likely secondary to reoxygenation (27). The authors hypothesized that SBRT may be significantly improved by allowing an inter-fraction time for reoxygenation while avoiding repopulation, which occurred after reoxygenation. Finally, high doses of radiation therapy may serve to slow cell cycle progression, and it is possible that increasing inter-fraction time may serve to allow for increased redistribution within the cell cycle and therefore additional tumor cell killing (28). It appears to be biologically plausible for an increased inter-fraction time to reduce toxicity, but prospective studies would be needed to confirm this hypothesis-generating result.

Several important risk stratifications were identified for central lung SBRT. Though BED >100 Gy trended towards improved local control in support of prior literature, lower dose treatments were also effective (1). Patients who obtained a biopsy trended towards improved overall survival. This may merely be the result of patients with fewer medical comorbidities having an increasing likelihood

of obtaining a biopsy, and these patients have improved overall survival, irrespective of their lung cancer. Intuitively, clinical stage, prior chemotherapy, and prior thoracic surgery trended towards increased disease progression, as these factors related to advanced or recurrent disease (21). In the case of prior thoracic surgery, patients presenting with recurrent disease may represent a different clinical scenario for SBRT than patients presenting with primary disease (29). While nodal targets trended towards decreased local-progression free survival, the fact that this variable was not predictive of overall survival or progression-free survival suggests that therapy offered clinical benefit (16). Prior radiotherapy was interestingly predictive of decreased disease progression. This result supports offering repeat courses of SBRT after recurrence. 57% of the treatments in this cohort involved recurrent cancer, and 49% of treatments had prior thoracic radiotherapy, offering a fairly large patient group to support such a hypothesis. Parks *et al.* studied a set of 27 patients treated with prior thoracic radiation for stage IIB-IIIB NSCLC and demonstrated encouraging outcomes but high rates of toxicity (63% rate of symptomatic pneumonitis) (30). While it is possible that incorporating increased inter-fraction time could serve to improve toxicity outcomes in cases of reirradiation, further study is needed. The prior courses of radiotherapy included in this cohort were heterogenous, including other courses of SBRT as well as concurrent chemoradiation to 60 Gy in 30 fractions. These significant differences indicate that these findings concerning reirradiation should be considered with caution.

The key limitation of this study is the variation in dose-fractionation schemes used for the different tumors. While this can partly be explained by the range of clinical stages included in the analysis, comparing these treatments presents inherent difficulties. Prospective analysis could select more specific dose-fractionation schemes for the different stages. Additional information to more completely describe the nodal distribution of the nodal targets in this study would have also proven beneficial. A prospective study could expand upon the hypothesis-generating results presented here and consider a cohort of patient treated with a more homogeneous inter-fraction time (e.g., exactly one week). Further, some of the variables considered may be interdependent, such as increasing stage with prior chemotherapy and nodal targets. Finally, multi-institutional study may serve to determine patients at higher risk for toxicity with the once-weekly fractionation. Because the rates of side effects were so low using this treatment

regimen, a large patient cohort may be required for such an analysis.

Conclusions

Our series demonstrated encouraging local control with low rates of toxicity for central lung SBRT, including many patients with stage 3 disease. This may be the result of the relatively large inter-fraction interval. This interval may allow for greater tumor effects (such as reoxygenation) and improved tolerance from normal tissues.

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Footnote

Reporting Checklist: The authors have completed the STROBE reporting checklist. Available at <http://dx.doi.org/10.21037/jtd-20-2659>

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Ethical Statement: The authors are accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved. This study was conducted in accordance with the Declaration of Helsinki (as revised in 2013). The study was exempt by the institutional review board and the informed consent, due to the retrospective nature of the study and the establishment of a previous protocol for conducting retrospective studies.

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