Case Report

Treatment of T3N0 non-small cell lung cancer with chest wall invasion using stereotactic body radiotherapy

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A B S T R A C T

Objectives: Chest wall invasion (CWI) is observed in 5% of localized non-small cell lung cancer (NSCLC). The role of stereotactic body radiotherapy (SBRT) in these patients is unknown. We investigate the safety and efficacy of SBRT in patients with T3N0 NSCLC due to CWI.

Methods: Patients with T3N0 NSCLC due to CWI were identified using a prospective registry. CWI was defined as radiographic evidence of soft tissue invasion or bony destruction. We excluded patients with recurrent or metastatic disease. All patients were treated with definitive SBRT. Prescribed dose was 50 Gy in 5 fractions for most patients. Kaplan-Meier analysis was used to estimate survival outcomes.

Results: We identified 12 patients treated between 2006 and 2017. Median age was 70 (range, 58–85). Median tumor diameter was 3.0 cm (range, 0.9–7.2). Median survival was 12.0 months (range, 2.4–63). At a median follow-up of 8.9 months (range, 2.1–63), 1-year primary tumor control was 89%, involved lobar control was 89%, local–regional control was 82%, distant control was 91%, and survival was 63%. Of the 4 patients with pre-treatment chest wall pain, 3 reported improvement after SBRT. Two patients reported new grade 1–2 chest wall pain. No grade 3+ toxicity was reported, with 1 patient experiencing grade 1 skin toxicity and 3 patients experiencing grade 1–2 radiation pneumonitis.

Conclusions: SBRT for CWI NSCLC is safe, with high early tumor control and low treatment-related toxicity. Most patients with pre-treatment chest wall pain experienced relief after SBRT, with no grade 3+ toxicity observed.

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1. Introduction

Non-small cell lung cancer (NSCLC) is the leading cause of cancer-related death in both the United States and worldwide [1]. Surgical resection and radiation therapy (RT) are considered standards of care for patients with early-stage NSCLC. For patients with medically-inoperable, early-stage disease, stereotactic body radiotherapy (SBRT) remains standard. Multiple prospective series have shown SBRT achieves high rates of tumor control with acceptably low treatment-related toxicity in both the medically operable and inoperable settings [2–4]. Of patients with localized NSCLC, approximately 5% have T3N0 staged disease by evidence of chest wall invasion (CWI) at diagnosis [5]. While the Radiation Therapy Oncology Group (RTOG) 0236 trial evaluating SBRT in medically inoperable early NSCLC did allow patients with T3N0 disease, no patients with T3N0 disease were enrolled.

With no prospective data to inform us of the role of SBRT in this select population, questions regarding effectiveness and safety are still yet to be answered. Of particular importance is chest wall toxicity, including skin toxicity, chest wall pain, and rib fractures. While tumors more than 2 cm from the chest wall or 5 cm from the posterior skin are at a very low risk, chest wall toxicity is relatively common in tumors of closer proximity [6]. With chest wall toxicity rates ranging from approximately 5% to 25%, multiple studies have identified additional factors predictive of developing chest wall toxicity, with the volume of irradiated chest wall, tumor size, prescription dose and fractionation all being significant variables [7–9]. However, T3N0 NSCLC with CWI commonly presents with pre-treatment chest wall pain and these models did not include this cohort. The purpose of our study is to identify tumor control and tolerability of SBRT in the management of patients with T3N0 NSCLC by virtue of CWI.
2. Materials and methods

An institutional review board-approved, prospectively-maintained database in accordance with the Health Insurance Portability and Accountability Act for patients with newly-diagnosed, previously untreated stage T3N0 NSCLC with CWI receiving SBRT with curative intent from 2006 to 2017 was reviewed. All patients signed an informed consent document for inclusion in this database prior to treatment. Chest wall invasion was defined as clinical or radiographic evidence of soft tissue and/or bony involvement, accordingly T3N0 per American Joint Committee on Cancer staging, 8th edition [10]. Patients with distant metastatic disease or recurrent disease at time of SBRT were excluded from this analysis. Patients with parietal pleural-based T3N0 lesions without evidence of chest wall soft tissue or bony destruction were also excluded.

All patients were evaluated by a multidisciplinary team, with medical inoperability determined by a cardiothoracic surgeon and/or pulmonologist. Pretreatment evaluation included history and physical examination as well as pulmonary function tests in most patients. All patients were staged via computerized tomography (CT) of the chest, positron emission tomography (PET), and/or mediastinal staging with endobronchial ultrasound (EBUS) or mediastinoscopy for equivocal PET findings. Our institutional SBRT technique has been described in a previous manuscript [11]. Tumor motion was controlled with abdominal compression to minimize the range of tumor motion. Patient setup and target verification was confirmed using daily cone beam CT. For SBRT treatment planning, although hotspots are typically avoided in the chest wall, if possible. For our five-fraction scheme, plan optimization was set to recapitulate similar hotspots as the 3DCRT plans. No specific chest wall constraints were utilized in plan optimization, but the planning target volume (PTV) received at least 95% of the prescribed dose covers the PTV. For IMRT/VMAT treatments, delivery was predominantly with non-coplanar intensity-modulated radiation therapy (IMRT), and coplanar volumetric modulated arc therapy (VMAT). For 3DCRT, prescription was typically to the 75% to 85% isodose line so that 95% of the prescribed dose covers the PTV. For IMRT/VMAT treatments, plan optimization was set to recapitulate similar hotspots as the 3DCRT plans. No specific chest wall constraints were utilized in the treatment planning process, although hotspots are typically avoided in the chest wall, if possible. For our five-fraction scheme, normal tissue constraints were respected for the spinal cord (maximum dose (Dmax) < 20 Gy), esophagus (Dmax < 30 Gy), heart (Dmax < 30 Gy), brachial plexus (Dmax < 25 Gy), and the volume of lung minus internal target volume (ITV) receiving at least 20 Gy (V20Gy) < 10%.

Patients were followed with serial CT scans at 6 weeks’ post-treatment, then typically every 3 months for 2 years, followed by every 6 months for an additional 3 years. Survival was calculated from time of treatment until death or last follow-up. Primary tumor failure was defined as radiographic evidence of local enlargement of the treated primary tumor. Hilar and mediastinal nodal recurrences were included in local–regional failures. The Kaplan-Meier method was used to estimate control and survival probabilities and Cox analyses were performed. Significance was considered at p < 0.05 and all significance levels were 2-sided. IBM® SPSS® Statistics, version 23 was applied for all statistical analyses.

3. Results

3.1. Patients and treatment characteristics

We identified 12 patients who met inclusion and exclusion criteria. Patient, tumor, and treatment-related characteristics are summarized in Table 1. Median age at diagnosis was 70 years (range, 58–85). No patients had received prior in-field RT or previous chemotherapy. Prescription dose was 50 Gy in 5 fractions (biological effective dose, BED10 = 100 Gy) for all but 3 patients: two received 54 Gy in 3 fractions (BED10 = 151 Gy) and one other patient received 62.5 Gy in 10 fractions (BED10 = 102 Gy). Patients received treatment every other day (10 patients) with the remaining two receiving fractions on consecutive week days. For the 9 patients requiring motion control, abdominal compression was used in all but 1 patient, who was managed with BodyFix (Bodyfix; Medical Intelligence, Inc., Quebec, QC, Canada) vacuum restriction bag. The remaining 3 patients had minimal tumor motion so compression was not used. Delivery was predominantly with non-coplanar 3D CRT (9 patients), with 2 patients being treated with coplanar IMRT and 1 other patient with noncoplanar VMAT (Fig. 1). No patients received adjuvant chemotherapy. Pretreatment pulmonary function tests were obtained in 8 patients (66%). The median FEV1 was 1.3 L and 44% (range, 0.4–2.5 L and 22–72%, respectively), and median DLCO was 46% (range, 33–69%). Only 2 patients had post-treatment testing, however, with minimal changes observed in both.

3.2. Outcomes

The median follow-up for all patients was 8.9 months (range, 2.1–62.5 months). For surviving patients, the estimated 1-year primary tumor control was 89%, involved lobar control was 89%, local–regional control was 82%, distant control was 91%, and over-

### Table 1

| Characteristic | Value (% or range) |
|---------------|--------------------|
| Age (years), median | 70 (58–85) |
| Sex | Male 5 (42%), Female 7 (58%) |
| BMI, median | 24.6 (17.3–40) |
| Smoking status | Never 0 (0%), Former 6 (50%), Current 6 (50%) |
| Smoking pack-years | 65 (15–108) |
| Age-adjusted CCI, median | 5 (3–10) |
| KPS, median | 60 (50–80) |
| Pretreatment hemoglobin (g/dl), mean | 12.7 (9.8–15.7) |
| Histology | Squamous cell carcinoma 6 (50%), Adenocarcinoma 2 (17%), NSCLC NOS 2 (17%) |
| Biopsy unable to be obtained | 2 (17%) |
| Tumor diameter (cm), median | 2.7 (0.9–7.2) |
| PET SUV, median | 13.8 (4.4–25.9) |
| Mediastinal staging | PET alone 8 (66%), EBUS 2 (17%), Mediastinoscopy 2 (17%) |
| Dose and fractionation | 50 Gy in 5 fractions 9 (75%), 54 Gy in 3 fractions 2 (17%), 62.5 Gy in 10 fractions 1 (8%) |
| Treatment schedule | Consecutive days 2 (17%), Nonconsecutive days 10 (83%) |
| Delivery Type | Noncoplanar 3D conformal 9 (75%), Coplanar IMRT 2 (17%), Noncoplanar VMAT 1 (8%) |
| PTV volume (cm³), mean | 72.2 (10.8–210.6) |
| BMI, body mass index; CCI, Charlson comorbidity index; KPS, Karnofsky performance status, NSCLC NOS, non-small cell lung cancer not otherwise specific; PET SUV, positron emission tomography standardized uptake value; EBUS, endobronchial ultrasound; IMRT, intensity-modulated radiation therapy; VMAT, volumetric modulated arc therapy; PTV, planning target volume
all survival was 63% (Fig. 2). Table 2 shows the patterns of failure among study patients. Nodal and distant failures were the most common sites of treatment failure. No isolated primary tumor failures were observed. Of the two patients who experienced local failures, both had either nodal or distant involvement at time of local failure as well.
Of the 4 patients who presented with pre-treatment chest wall pain, 3 (75%) of them reported either improvement or resolution of their symptoms, with one patient reporting improvement but not resolution of their symptoms with no further need for narcotic use, and the other two patients reporting complete resolution of their symptoms. The other patient reported no change in the symptoms post-treatment. Of the 8 patients who did not have pre-treatment chest wall pain, 2 patients developed chest wall pain requiring over-the-counter medication. The first patient developed chest wall pain 2 months after SBRT, correlating with development of pleural-based metastases outside the treatment field. This patient’s symptoms were improved with over-the-counter non-steroidal anti-inflammatory (NSAID) medication, and the patient passed away from metastatic disease before resolution of their symptoms was documented. The other patient developed mild rib pain 13 months after treatment that did not require narcotics and resolved with a short course of NSAIDs. No patients developed treatment-related chest wall pain requiring narcotics after SBRT and no patients had clinical or radiographic evidence of rib fractures on follow-up CT imaging. Grade 1 skin toxicity was observed in 1 patient after treatment which resolved spontaneously. Radiation pneumonitis was observed in 3 patients, grade 1 in two cases and only one case of symptomatic grade 2 pneumonitis managed with steroids. No grade 3+ pneumonitis or other toxicity was observed in this cohort.

4. Discussion

We report that SBRT can be safely and effectively delivered in patients with primary T3N0 NSCLC invading the chest wall. The frequency of treatment-associated toxicity was low, with no grade 3+ events and symptoms typically being mild and/or self-limited in the few patients experiencing side effects. Local and lobar control were high at 89% each in our cohort, although duration of follow-up was limited, with nodal and distant spread being the most common patterns of failure in this medically high-risk patient population. Importantly, in the subset of patients presenting with pre-treatment chest wall pain, most patients experienced relief of their symptoms, indicating that SBRT may be an effective tool to provide pain relief by virtue of local control.

Historically, patients with lung cancer invading the chest wall have presented unique challenges to definitive management. The standard of care for these patients has long been en bloc surgical resection, often followed by adjuvant chemotherapy. Generally, acceptable margins are 1 cm in all directions, although some argue for removing 1 rib above and below the tumor as well as achieving lateral margins of 3 to 4 cm, given that survival is highly dependent on completeness of resection [12]. The necessity of complete resection is balanced by the fact that these are often extensive surgeries requiring reconstruction of large chest wall defects, although chest wall defects less than 5 cm can typically be managed without reconstruction [5]. Even at high-volume centers with skilled thoracic surgeons, perioperative mortality is relatively high, with experiences from Memorial Sloan-Kettering and multi-center French series reporting mortality rates of up to 8% [13–15]. Survival after surgical resection varies significantly in the literature but is generally poor, as 5-year overall survival is approximately 40%, with margin status, extent of invasion, and pathologic nodal status all significantly influencing this rate [16–18].

Given that surgical resection presents unique, significant challenges and that many patients are poor operative candidates due to comorbid cardiopulmonary disease, nonoperative approaches to management are often necessary. Conventionally-fractionated radiotherapy has long been described in medically-inoperable NSCLC, with 2-year disease-free survival only 25% for T3N0 patients treated to doses of at least 60 Gy in 2 Gy fractions [19]. Given historically poor disease control rates with conventionally fractionated definitive radiotherapy alone for early stage NSCLC as well as the medical frailty of this demographic, it is unsurprising that survival at 2- and 5-years has been 35% and 15%, respectively, in multiple large series [20,21].

Chest wall pain is a common presenting symptom in these patients, causing significant detriment to quality of life and further underscoring the importance of local control as an important metric in this cohort. Accordingly, the local control in our series was 89% at 1 year, comparable to reported rates in large SBRT series of T1-2N0 lesions not invading the chest wall [22,23]. The need for maximizing local control is balanced with minimizing treatment-related chest wall toxicity, as large SBRT series have shown rates of acute and chronic chest wall pain of approximately 5% and 17%, respectively with chest wall V30 and obesity predicting for both pain and skin toxicity [24]. In our subset of patients that did present with pre-treatment chest wall pain, 75% achieved pain relief with 50% experiencing complete pain relief. These findings corroborate the Cleveland Clinic experience with treating 10 patients with primary T3N0 NSCLC and an additional 3 patients with recurrent NSCLC invading the chest wall, which showed pain relief of 80%, with 50% achieving complete resolution of symptoms [25]. To our knowledge, there are no other published studies to-date characterizing outcomes other than the above work by Cleveland Clinic and our current series. While two of our patients did experience new chest wall pain after treatment, both cases were grade 1–2 with one of these likely due to development of pleural-based metastases and the other relieved with over-the-counter medication. We observed no grade 3+ chest wall toxicity, including no rib fractures, although these events may be observed over 1-year post-treatment. Furthermore, although post-SBRT pulmonary function testing was only obtained in 2 of our patients, no clinically-significant changes from baseline were observed, consistent with other series [26].

Nodal and distant relapse were much more common than local failures in our series, which is consistent with the greater SBRT experience for early-stage NSCLC [27]. This observation has led some to advocate for adjuvant chemotherapy after SBRT, extrapolating from the survival advantage seen in the Cancer and Leukemia Group B (CALGB) 9633 trial for node-negative tumors 4 cm or greater who had undergone adjuvant chemotherapy after surgery, although none of our patients with larger tumors received adjuvant systemic therapy [28]. Following a similar logic, multiple retrospective series have shown improved survival with adjuvant chemotherapy in surgically-rected pT3N0 NSCLC with chest wall invasion, independent of size [29–31]. In our series, one third of patients developed regional nodal metastases as a component of failure, which might be decreased by systemic therapy. Despite these observations, the role of adjuvant systemic therapy after SBRT for NSCLC is still poorly-defined and may warrant prospective investigation.

### Table 2
Patterns of failure among study patients (n = 12).

| Failure type                      | Number |
|----------------------------------|--------|
| Local                            | 2      |
| Local only                       | 0      |
| Local and nodal                  | 1      |
| Local, lobar, nodal and distant  | 1      |
| Nodal                            | 4      |
| Nodal only                       | 1      |
| Nodal and distant                | 1      |
| Distant                          | 4      |
| Distant only                     | 2      |
| Metachronous lung                | 0      |
Our study is affected by several limitations inherent to its design. We investigated a small population of patients with multiple medical comorbidities, with follow-up duration limited by the high number of intercurrent deaths in the study interval. Consequently, estimates of tumor control are short and further follow-up with larger sample size may provide more accurate assessment of these measures. Rates of treatment-related toxicity may be underestimated by retrospective review, and the addition of patient-reported quality of life data would further delineate the analgesic effect of SBRT in this cohort. Furthermore, SBRT techniques have evolved over the study duration at our institution. Currently, patients are typically treated on consecutive days while most of the patients in our study were treated when treatment on nonconsecutive days was routine at our institution, although previous studies have shown no impact of treatment schedule on outcome [32]. Furthermore, while most patients on our study were treated with noncoplanar 3D conformal techniques, we are increasingly treating patients with VMAT, although its impact on control and toxicity, if any, are unknown. Despite these limitations, SBRT appears to be a promising treatment modality for patients with chest wall-invading tumors.

5. Conclusion

SBRT for patients with NSCLC invading the chest wall appears to be a feasible, safe, and well-tolerated treatment modality associated with high rates of early tumor control. For patients experiencing local symptoms, SBRT is effective in providing at least partial pain relief. For medically-inoperable patients with T3N0 NSCLC due to CWI, SBRT should be considered as a potential management option, and warrants further investigation.

Declarations

Ethics approval and consent: Approved by the institutional review board in accordance with the Health Insurance Portability and Accountability Act.

Consent for publication: Not applicable.

Availability of data and materials: Please contact author for data requests.

Authors’ contributions: WK was responsible for study conception and design, acquisition of data, analysis and interpretation of data, and drafting of the manuscript. PG was responsible for acquisition of data, analysis and interpretation of data. JB was responsible for study conception and design, interpretation of data, and critical revision.

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Conflict of interest

We have no conflict of interest to disclose for this work.

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Not applicable.

Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.ctro.2019.02.004.

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