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Abstract: Radiation therapy is the foundation for treatment of locally advanced non-small cell lung cancer (NSCLC), a disease that is often inoperable and has limited long term survival. Local control of disease is strongly linked to patient survival and continues to be problematic despite continued attempts at changing the dose and fractionation of radiation delivered. Technological advancements such as 4-dimensional computed tomography (CT) based planning, positron emission tomography (PET) based target delineation, and daily image guidance have allowed for ever more accurate and conformal treatments. A limit to dose escalation with conventional fractions of 2 Gy once per day appears to have been reached at 60 Gy in the randomized trial Radiation Therapy Oncology Group (RTOG) 0617. Higher doses were surprisingly associated with worse overall survival. Approaches other than conventional dose escalation have been explored to better control disease including accelerating treatment to limit tumor repopulation both with hyperfractionation and its multiple small (<2 Gy) fractions each day and with hypofractionation and its single larger (>2 Gy) fraction each day. These accelerated regimens are increasingly being used with concurrent chemotherapy, and multiple institutions have reported it as tolerable. Tailoring treatment to individual patient disease and normal anatomic characteristics has been explored with isotoxic dose escalation up to the tolerance of organs at risk, with both hyperfractionation and hypofractionation. Metabolic imaging during and after treatment is increasingly being used to boost doses to residual disease. Boost doses have included moderate hypofractionation of 2–4 Gy, and more recently extreme hypofractionation with stereotactic body radiation therapy (SBRT). In spite of all these changes in dose and fractionation, lung and cardiovascular toxicity remain obstacles that limit disease control and patient survival.

Keywords: Radiotherapy; carcinoma; non-small cell lung; dose fractionation

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Introduction

Lung cancer has long been the most common cause of cancer death worldwide, claiming 1.7 million lives annually (1). Non-small cell is the most common type of lung cancer diagnosed, and approximately 25% of patients will present with locally advanced disease (2). Locally advanced disease is typically defined as American Joint Committee on Cancer stage III, with unresectable primary tumors (T4) or involvement of lymph nodes in the mediastinum (N2) or supraclavicular fossa (N3) (3). While locally advanced disease is curable, patient outcomes unfortunately remain poor despite both radiation therapy, chemotherapy, and now immunotherapy having improved over time.

The foundation of treatment for locally advanced disease is radiation therapy. Radiation is usually combined with
Conventional fractionation

Radiation therapy for locally advanced non-small cell lung cancer (NSCLC) is most commonly given in fractions of 1.8 to 2.0 Gy once a day for five days a week with external beams. Radiation Therapy Oncology Group (RTOG) 7301 established a dose of 60 Gy as the standard, after escalating the total dose from 40 Gy in a split course, then to continuous delivery from 40, to 50, and finally to 60 Gy, with the higher doses having higher intrathoracic tumor control rates (8). It should be noted that this dose escalation was done in the two dimensional era when plans were made on based upon X-rays. This was before computed tomography (CT) scanners were used in the clinic. Since then, advances in radiation treatment have included three dimensional CT-based conformal treatment planning, the use of positron emission tomography (PET) for more accurate staging and targeting, and accounting for tumor motion during treatment.

Traditionally, external beam radiation therapy for locally advanced NSCLC targeted the primary tumor as well as the ipsilateral hilar and mediastinal nodal stations, and sometimes even the supraclavicular fossa. This was done even without evidence of involvement in these regions and is known as elective nodal irradiation. This was done to control any microscopic disease in areas felt to be high risk for regional metastases. The concern with elective nodal irradiation, however, is that it will limit dose escalation as the larger volume irradiated will lead to otherwise avoidable pulmonary, cardiovascular, and esophageal toxicity. Three randomized trials on its use and several cohorts have been reported. A meta-analysis of these studies shows the overall incidence of elective nodal failure was 5.5% without elective nodal irradiation and 3.4% with it (P=0.64 assuming a fixed-effects mode) (9). Given this lack of a significant difference, radiation is now most often prescribed only to gross disease that is enlarged on CT or hypermetabolic on PET.

Conventional dose escalation

In attempts to improve local control, radiation dose escalation with conventional fractionation and concurrent chemotherapy was performed by several groups in phase I and II trials. RTOG 0117 escalated to 75.25 Gy in 2.15 Gy daily fractions prescribed to the isocenter. Three of the eight patients who received 75.25 Gy developed dose-limiting pulmonary toxicity, one of which was fatal. The maximum tolerated radiation dose was therefore determined to be 74 Gy in 2 Gy fractions (10). North Central Cancer Treatment Group (NCCTG) 0028 escalated from 70 Gy up in 4 Gy increments. Similar to RTOG 0117, the maximum tolerated dose was also determined to be 74 Gy. Two of four patients experienced dose limiting pneumonitis at 78 Gy (11). The University of North Carolina first escalated from 60 to 74 Gy both with limited elective nodal radiation and with induction and concurrent chemotherapy, then later from 74 to 90 Gy showing grade 3 or greater late complications in 22% of patients (12). Cancer and Leukemia Group B (CALGB) 30105 was a phase II trial treating at 74 Gy and randomizing between gemcitabine and paclitaxel used in both induction and concurrent treatment (13). Aside from the gemcitabine arm of this trial which closed early due to high toxicity, most of these trials showed a median survival of around 2 years.

These trials then led to the multi-institutional randomized controlled phase III trial RTOG 0617. In this 2x2 factorial design trial, patients with stage III NSCLC all received weekly carboplatin and paclitaxel chemotherapy with concurrent radiation in 2 Gy per once daily fraction followed by two cycles of consolidative chemotherapy after the completion of radiation. Patients were randomized to receive either 60 or 74 Gy and to receive this with or without cetuximab. At the first interim analysis, the monitoring committee established that the trial had crossed the futility boundary with respect to the 74 Gy arm, and this high dose arm was closed. The trial continued accruing only at the 60 Gy arm. At the third interim analysis, it was established that the cetuximab arm had also crossed the futility boundary.
The published results of 0617 showed significant increased rates of death in the 74 Gy arm, challenging the assumption established in the earlier phase I/II studies that radiotherapy dose escalation with conventional fractionation and concurrent chemotheraphy will improve outcomes (14). The median survival was 28.7 months in the 60 Gy arm versus 19.5 months in the 74 Gy arm (P=0.0007; HR 1.56, 95% CI: 1.19–2.06) (5). There was a 37% increased risk of local failure in the high dose arms (P=0.0319, HR 1.37, 95% CI: 0.99–1.89). There was no difference in severe (Common Terminology Criteria for Adverse Events grade ≥ 3) toxic effects overall, in severe pulmonary events specifically, or in severe radiation pneumonitis between the radiation therapy dose groups. Severe esophagitis was more common in the high-dose group (21% vs. 16%, P<0.0001). On multivariate analyses, factors predicting worse overall survival were maximum esophagitis grade, planning target volume size, heart dose, and radiation dose.

The poor survival with treatment to 74 Gy has been attributed to several causes. Treatment-related deaths were more common in the high-dose group than in the low-dose group (10 vs. 2), but this comparison did not reach statistical significance. Concurrent chemotherapy was more difficult to complete in the high-dose group than in the low-dose group. Rates of protocol non-compliance were greater in the high-dose arm, 26% vs. 17% (P=0.02), as were treatment delays. Radiation therapy planning was more likely to be non-compliant in the high-dose group, and planning target volume coverage by the 95% isodose line was poorer in the high-dose group. Concerns that non-compliance in the high-dose groups produced these results led to analysis of overall survival only in those patients with radiation plans compliant with the protocol; nevertheless, overall survival was still better in the standard-dose groups than in the high-dose groups.

The fact that heart dose was a significant predictor of overall survival on the multivariate analysis of RTOG 0617 strongly suggests that it is not only dose to tumor that should be considered in future studies. While further analysis of RTOG 0617 is pending, three retrospective studies also suggest heart dose can predict overall survival and cardiac events. The largest with 322 patients identified higher doses as important for overall survival and generated a new and more conservative heart constraint of V50 <25%, or letting no more than 25% of the heart exceed 50 Gy (15). Two smaller series of 125 and 112 patients focused on cardiac events and showed mean heart dose was important (16,17). A secondary analysis of RTOG 0617 showed that patients treated at centers with high trial accrual, a potential surrogate for number of NSCLC patients treated annually, had better survival, lower esophageal and heart doses, and lower lethal events (18). Radiation associated cardiac toxicity after treatment of locally advanced NSCLC may occur earlier than historically understood, and thus heart doses should be minimized with any future attempts at dose escalation.

Likely due to PET staging and modern radiation therapy techniques, the 28.7-month median survival in the 60 Gy arm was longer than that seen in previous studies, was better than anticipated, and set a new benchmark for patients with locally advanced NSCLC receiving concurrent chemotheraphy and radiation therapy. Consequently, current trials in the United States such as RTOG 1306 and National Research Group (NRG) L001 have adopted 60 Gy as the standard, as well as the American Society for Radiation Oncology (ASTRO) in its guidelines (19). The National Comprehensive Cancer Network (NCCN) guidelines in the United States now suggest definitive radiation should be 60 to 70 Gy (20). The European Society for Medical Oncology guidelines state that “dose in excess of 66 Gy is not recommended outside trials” (21). The Cancer Council of Australia (CCA) says that radiation dose “should be at least 60 Gy assuming that dose-volume constraints on organs at risk are met” and that “74 Gy is not better than 60 Gy and may be potentially harmful” (22). Many other national guidelines, such as China’s from 2015 (23), do not make explicit dose recommendations yet.

Hyperfractionation
Not all patients will be felt fit enough to tolerate concurrent treatment with chemotherapy, however, and many of those same national guidelines recommend accelerated hyperfractionation in such patients. The National Institute for Health and Care Excellence (NICE) and CCA both recommend continuous hyperfractionated accelerated radiation therapy (CHART) when only radiation therapy will be used as treatment. Acceleration refers to finishing treatment more rapidly than is done conventionally. Hyperfractionation refers to dividing treatment into smaller doses more than once a day. The CHART trial compared 36 fractions of 1.5 Gy given three times per day to a total dose of 54 Gy in 12 consecutive days with 30 fractions of 2 Gy once a day to a total dose of 60 Gy in 6 weeks. CHART improved 2-year overall survival from...
20% to 29% (P=0.008). The majority of patients (81%) had squamous cell cancer, and in such patients, 2-year overall survival improved from 20% to 33% (P=0.0007) (24). This regimen has never been directly compared to conventional chemoradiation in a randomized trial, however. An individual patient data meta-analysis showed that hyperfractionation improved overall survival by 2.5% at 5 years without improving progression free survival when compared to conventionally fractionated radiation alone. This came at the risk of worse acute esophageal toxicity (odds ratio of 2.44 in NSCLC) (25). The theory is that prolongation of treatment results in increased tumor cell replication (26), and has been proposed as one reason why the high dose arm of RTOG 0617 failed.

However, the CHART regimen is inconvenient as many centers are closed on weekends, and it often requires that the patients be hospitalized. The randomized phase III CHARTWEL (continuous hyperfractionated accelerated radiotherapy weekend less) looked at 60 Gy in 40 fractions over 2.5 weeks versus 66 Gy in 33 fractions over 6.5 weeks. In contrast to CHART, CHARTWEL showed no difference in overall survival and local tumor control. Exploratory analysis suggested trends for improved local control with hyperfractionation for higher stages of disease and after neoadjuvant chemotherapy. Histology did not affect control, and it should be noted that 58% of patients had squamous cell cancers, in contrast to the 81% in CHART where hyperfractionation showed the most promise in this histology. Another potential explanation for the lack of difference in overall survival was the fact that total radiation doses were 10% higher in both arms of CHARTWEL when compared to CHART (27).

Hyperfractionated radiation with concurrent chemotherapy has also been compared to the combination with conventional radiation. A 2×2 trial randomizing 60 Gy in 30 fractions in either 6 weeks once a day or 3 weeks twice a day with or without concurrent carboplatin showed no statistically significant difference in local recurrence or survival in all arms (28). RTOG 9410 showed that concurrent chemotheraphy with hyperfractionated radiation to 69.6 Gy had statistically similar 5-year overall survival when compared to conventional chemoradiation to 63 Gy, 13% vs. 16% (P=0.46). Grade 3 esophagitis was statistically worse at 45% with hyperfractionation, double that of 22% with conventional fractionation (P<0.001) (29). When coupled with the logistical issues of delivering multiple fractions per day and patient inconvenience, for patients felt fit enough for concurrent treatment, conventional fractionation became standard in future trials.

Hypofractionation

Most recent studies have investigated accelerating treatment with hypofractionation with larger radiation doses delivered once per day, rather than with hyperfractionation and its smaller doses delivered more than once per day. Concerns about serious toxicity delayed exploration of hypofractionation until technology such as CT based planning, daily image guidance, and gating became widespread. Most reports have been for modest hypofractionation of 2–3 Gy per fraction. In patients receiving radiation alone, three studies comparing standard fractionation to hypofractionation did not report any significant differences in toxicity or disease outcomes (30–32). In a review of 22 studies of hypofractionated radiation alone, Kaster et al. found the weighted mean acute toxicity in the esophagus and lung being 1.9% and 1.2% respectively. Late toxicity was also low at 1.4% and 6.9%. Two-year overall survival ranged from 18% to 42%. There was a moderate linear relationship between biologically effective lesional dose (BED10): for every 1 Gy increase in BED10, there was an absolute overall survival benefit ranging from 0.36% to 0.70% (33). This is similar to the results found by Machtay et al. with conventional radiation where a 1 Gy increase in BED resulted in a 4% relative improvement in survival (14), with in the context of 15% long-term survival, is an absolute improvement of 0.6%.

Hypofractionation may allow for better outcomes by increasing BED without lengthening treatment time and thereby preventing cancer cell repopulation. Studies of modest hypofractionation with concurrent chemotherapy are fewer and mostly single arm, single institution studies. In the same systemic review by Kaster et al., 15 studies of hypofractionated radiation therapy with concurrent chemotherapy were found. The weighted mean acute toxicity in the esophagus and lung was 14.9% and 7.9% respectively, and for weighted mean late toxicity in the esophagus and lung, 6.6% and 12.2%. In comparison, RTOG 0617 showed acute grade 3 esophagitis in 7% and grade 3 pneumonitis in 4% with late toxicity rates <1% for both. Two-year overall survival with concurrent hypofractionation ranged from 24% to 58%. The 2-year overall survival in RTOG 0617’s standard arm was 58% (5). In the United Kingdom, 55 Gy in 20 fractions of 2.75 Gy is the most commonly used schedule, both with and without concurrent systemic therapy (34).
More aggressive hypofractionation to 4 Gy per fraction has been also been explored in patients not felt fit for concurrent chemotherapy. MD Anderson used protons to escalate to 60 Gy cobalt-equivalent in 15 fractions in a phase 1 trial. At a median follow-up of 13 months, two of 25 patients experienced dose-limiting toxicities. One treated to 52.5 Gy cobalt-equivalent in 15 fractions developed a tracheoesophageal fistula after bevacizumab was delivered for recurrent disease. The second developed “possible” grade 3 radiation pneumonitis (35). This regimen with protons is currently being explored with concurrent chemotherapy in a phase 1 study at our institution (NCT02172846).

As protons are not available at most centers, a phase 1 dose escalation study was later reported using photons at University of Texas Southwestern in patients who were not chemotherapy candidates. It enrolled 55 patients divided between 50, 55, and 60 Gy, all in 15 fractions. One patient developed grade 3 esophagitis, and 2 cases of grade 3 dyspnea were felt related to therapy. There was no association between fraction size and toxicity (P=0.24), and the median overall survival was 6 months at all dose levels (P=0.59) (36). This same group has presented in abstract form an interim analysis from a randomized phase III comparison of 60 Gy in 15 vs. 30 fractions of image-guided photon radiation therapy in patients with a Zubrod performance status of 2 or greater. Median overall survival for the 48 patients evaluable was 11.5 months with no statistical difference between conventional and hypofractionated radiation treatment arms. Two deaths from hypoxia with conventional radiation and 1 death with hypofractionated radiation were possibly related to treatment (37). Final results are pending, but the study authors feel the results could potentially change the paradigm of treatment for patients with locally advanced disease receiving radiation alone due to poor performance status.

**Adapted therapy**

Rather than prescribing the same fixed dose to all patients with locally advanced disease, isotoxic radiation therapy is a novel approach which allows for personalized treatment planning based on individual tumor and patient characteristics. This tailored approach is heavily based upon predefined organ at risk dose constraints. Treatment plans are designed to give the maximum BED achievable to the tumor target until the predefined dose constraints are reached. The increasing use of computer-based inverse-planned intensity modulated radiation therapy (IMRT) makes this approach especially feasible.

One such approach with hyperfractionation, individualized isotoxic accelerated radiotherapy (INDAR) has been used for over a decade in the Netherlands. In the first three weeks, 30 twice daily fractions of 1.5 Gy are delivered. Next, 2 Gy fractions once a day are delivered until a mean lung dose of 19 Gy is reached, with a total ranging between 54 to 69 Gy in 5.5 weeks. The equivalent conventional dose would be 72 Gy over 36 fractions. Long term results showed that with sequential chemotherapy and INDAR, the median survival was 23.6 months. For comparison, the use of sequential chemotherapy and conventional fractionation with the same group resulted in a lower median survival of only 17.5 months (38). The use of concurrent chemotherapy in a phase 1 study with INDAR showed no dose limiting toxicity (39).

Instead of hyperfractionation, other institutions have used isotoxic planning with hypofractionation. A single institution phase I trial used 25 fractions ranging from 2.28 to 3.42 Gy (a total of 57 to 85.5 Gy) with IMRT. Total dose was escalated according to each patient’s individual stratified risk for radiation pneumonitis. The maximum tolerated dose was predefined as the dose that theoretically would result in ≤20% risk of severe toxicity. Grade 4 to 5 toxicity was reached late in 6 of the 79 patients, and the maximum tolerated dose was defined as 63.25 Gy in 25 fractions of 2.53 Gy each. These severe toxicities were due to damage to the central and perihilar structures and corresponded to dose to the proximal bronchial tree (40).

RTOG 1106, which recently completed accrual, coupled the use of isotoxic hypofractionation with metabolic imaging to adapt chemoradiation treatment for locally advanced disease. The control arm was standard fractionation of 60 Gy in 30 fractions of 2 Gy each. In the experimental arm, the first 21 fractions were 2.2 Gy each. The final nine fractions were delivered only to residual disease seen on a PET taken after fractions 18 or 19. Residual disease was defined as any sites with metabolic activity at least 150% of the aortic arch. These final nine fractions could range from 2.2–3.8 Gy per fraction, corresponding to a range of 19.8 to 34.2 Gy. The highest achievable dose was given while still respecting a mean lung dose of 20 Gy. The theoretical maximum tumor dose was 80.4 Gy. Patients were randomized 2:1 into the experimental arm with stratification by stage, primary tumor size, and histology. The primary objective was local control, and results are...
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Pending.

RTOG 1106 was based on studies from University of Michigan which demonstrated that tumors decrease more in metabolic activity than size during treatment. Recently, this group reported on the results of a phase II study of conformal radiation individualized to a fixed risk of radiation-induced lung toxicity of 17.2% for grade 2 or greater pneumonitis. Dose to residual tumor was adaptively escalated to residual tumor on a mid-treatment PET up to a total dose of 86 Gy in 30 fractions. The median tumor dose delivered was 83 Gy over 30 fractions. The initial doses per fraction were 2.1 to 2.85 Gy initially over 18–24 fractions, then 2.85 to 5.0 Gy for the adaptive phase after the mid-treatment PET. Most patients (93%) received concurrent carboplatin and paclitaxel with consolidation chemotherapy. With a median follow-up for surviving patients of 47 months, the 2-year rate of infield tumor control was 82% with a median overall survival of 25 months (41).

Rather than using mid-treatment imaging, other groups have looked at boosting any residual disease after the completion of conventional treatment with stereotactic body radiation therapy (SBRT). SBRT has proven very effective in early stage disease, especially in lesions far from major airways and the mediastinum (42). With SBRT, BEDs of at least 100 Gy total are typically delivered in 3–5 fractions with narrow margins. Four studies have looked at a SBRT boost to residual disease after conventional chemoradiation therapy to 50–60 Gy. In these small series with limited follow-up, local control at one year has been approximately 80% (43–46), comparable to that of RTOG 0617, which was a mixed group that contained patients who had a complete response. Of the 80 patients in these four studies of SBRT boosts, there have been 5 (6.3%) lethal toxicities such as pulmonary hemorrhage, particularly with boosts to central disease near the main airways and mediastinum.

A multi-institutional trial has been proposed that would use a SBRT boost to metabolically active residual disease seen on PET-CT 2–4 weeks after chemoradiation treatment completion (47). The benchmark for trial success would be that in patients with residual disease after conventional treatment, a SBRT boost would result in progression-free survival of 20–30% at 2 years, matching that seen in RTOG 0617. Now with immunotherapy recently showing a progression free benefit when used after conventional chemoradiation (48), SBRT becomes even more attractive as a boost as it could theoretically better present tumor antigens and better serve as a potentiator of the patient’s own immune system against their disease (49).

Conclusions

Locally advanced NSCLC remains a challenging disease with significant mortality and complications. Efforts to improve treatment outcomes have been only moderately successful, but the combination of hypofractionation with systemic therapy, individualized treatment adaptation, and stereotactic boosts in the era of immunotherapy offers the promise of further improvements.

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Footnote

Conflicts of Interest: The authors have no conflicts of interest to declare.

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