Combining Systemic Therapies with Radiation in Non-Small Cell Lung Cancer

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Abstract
Radiotherapy has been the mainstay of the treatment of stage III non-small cell lung cancer (NSCLC) patients. In the early nineties, combined treatment with chemotherapy was introduced. In 1995, a meta-analysis showed improved treatment outcome of the sequential use of cisplatin-based chemotherapy and radiotherapy (RCT) compared to radiotherapy alone. Subsequent randomized studies and the two meta-analyses demonstrated that concurrent radiochemotherapy (RCT) is superior (local control and overall survival) to sequential used both method. However, several questions remain unanswered concerning the optimal chemotherapy regimen and radiotherapy doses and techniques in terms of treatment outcome and toxicity profile. Targeted therapies represent a new class of drugs which interfere with specific molecular targets (typically proteins) playing critical roles in tumor growth and progression. Some combinations appear to be too toxic like the vascular epithelial growth factor antibody bevacizumab. The feasibility of adding the epidermal growth factor receptor inhibitor cetuximab has been recently reported for NSCLC patients. Strategies to safely incorporate novel antiangiogenic agents into combined-modality therapy in lung cancer are needed. The rapid development of molecular oncology will hopefully contribute to a better patient selection to particular strategies and to treatment optimization. Increasing radiotherapy doses applied according to up-to-date techniques and combinations with new biologicals might lead to further treatment improvements.

Keywords: Lung cancer; Chemotherapy; Radiotherapy

Introduction
Lung cancer is the most common cause of cancer death globally [1]. Most cases of lung cancer occur around the age of 60–70 years [2].

Treatment of non-small cell lung cancer (NSCLC) is challenging in many ways. Until the 1990s radiotherapy alone was the standard treatment for stages IIIA and IIIB non-small cell lung cancer (NSCLC). With the standard dose of 60 Gy in 30 fractions survival rates were extremely poor [3]. Indeed technical developments allowing the administration of higher radiation doses resulted in strategies to improve the treatment results include increasing doses of radiotherapy and decreasing overall treatment time [4]. For NSCLC a dose-effect relationship exists: the higher the radiation dose, the greater the probability of tumor control improved local control and survival [5]. The theoretical solution of simply increasing radiation doses to high biologically effective doses, ideally above the threshold of 100 Gy, has been suggested by several groups [6-9]. However, radiation dose escalation does not address the issue of distant or out-of-field relapses. A different option therefore is to combine radiotherapy with chemotherapy. The first report on improved survival after adding chemotherapy to the radiation was published more than 20 years ago [10]. Over the past decades, concomitant chemotherapy and radiotherapy has become the established treatment for patients with stage III non-small-cell lung cancer. In this review, we present current clinical knowledge on combining available systemic therapies with radiation.

Radiochemotherapy in Locally Advanced Non-Small Cell Lung Cancer

The strategy of exclusive radiotherapy for locally advanced inoperable NSCLC has been challenged after the publication of the meta-analysis by the Non-small Cell Lung Cancer Collaborative Group in 1995 [11]. Since then a combination of chemotherapy and radiotherapy is the recommended treatment in this group of patients. Radiotherapy preceded by (usually) two courses of chemotherapy yielded an improvement of the 2-year overall survival rate from 21% to 25%. The 5-year survival increased from 6% to 8% provided that the chemotherapy regimen included cisplatin. The effect was explained by a reduction of distant metastases. Until now this effect of a lower distant metastasis rate was observed in one study only [12]. In this study, Le Chevalier et al. compared radiotherapy alone to chemotherapy and radiotherapy. However, patients with adenocarcinoma were excluded. Since an important proportion of the NSCLC patients were not included, the results might not be representative. The 2-year survival rate was 14% in radiotherapy alone group and 21% in combined treatment group. The 3-year survival rate was 12% for the combination arm versus 4% for the radiotherapy arm (P<0.02) and local control was poor in both groups (17% and 15%, respectively). To our knowledge, these results have never been confirmed. Until recently sequential cisplatin-containing radiochemotherapy has been the standard treatment for inoperable stage IIIA and IIIB disease. Various chemotherapy schedules have been applied, but the treatment outcome did not differ significantly.

Despite this progress, both loco-regional and distant failures are frequent. Over the last 20 years, concomitant use of radiotherapy and chemotherapy has been extensively studied in various malignancies, including non-small cell lung cancer, rectal cancer, anal cancer and head and neck cancers, and has currently replaced radiotherapy alone in patients with good performance status. This strategy, through superadditive effect, not only improves local tumor control but...
also increases the overall survival [13]. The benefits of concomitant radiochemotherapy include a potential synergism between both modalities and avoiding the delay of radiotherapy. Therefore, there is a rationale for considering concomitant chemo-radiation also in patients with high-risk lung cancer. Attempts to improve the loco-regional control included increasing the radiotherapy dose using altered fractionation regimens and combining chemotherapy with radiotherapy. After phase I and phase II studies, the EORTC started a 3-arm phase III trial comparing split-course radiotherapy of 55 Gy using the same radiotherapy scheme, concurrently combined with 30 mg/m² cisplatin once a week or 6 mg/m² daily [14]. No improvement was seen after treatment with radiotherapy and weekly cisplatin. The 6 mg/m² cisplatin daily added to radiotherapy improved survival due to improved control of local disease. The difference was also significant after adjustment for known prognostic factors in a multivariate analysis. There was no effect on the distant metastasis rate, and late toxicity was not increased. These data demonstrated that cisplatin improved the radiotherapy effect by radiosensitization. The most frequently reported acute side effects were nausea and vomiting. In 1992, Trovo et al. [15] also published their randomized phase III study. Three weeks of radiotherapy, to a dose 45 Gy, were compared to the same radiotherapy dose with the addition of 6 mg/m² cisplatin daily. In this study no significant advantage of the combined treatment over radiotherapy only was found. However, this result may be due to the lower dosage of radiation used in the study (Table 1).

All phase III trials were included in a meta-analysis including 12 trials and 1921 patients by Aupérin et al. [16] indicated a 4% survival gain at 2 years and 2% at 5 years for concurrent chemoradiation versus radiotherapy alone, a comparable improvement to that observed with the sequential combination. Even though this meta-analysis was based on individual patient data it did not allow to accurately define the size of such a potential treatment benefit and the optimal schedule.
of chemotherapy. The efficacy of concurrent chemoradiotherapy versus radiotherapy also was compared in a metaanalysis including 14 randomized studies (and 2393 patients) in 2010 [17]. A Cochrane meta-analysis confirmed these conclusions: concurrent chemoradiotherapy was associated with 14% reduction in the risk of death at 2 years compared to sequential chemoradiotherapy, and a 7% reduction compared to radiotherapy alone.

If sequential and concurrent radiochemotherapy improved overall survival, so there is another question: which is better? In several trials improved 1- and 2-year overall survival rates in favour of the concurrent arm were reported [18-23]. Most of these trials were included in a new meta-analysis based on individual patient data by Auperin et al. [24] concluded that concurrent radiochemotherapy yielded superior results compared to the sequential combinations. There was a significant benefit of concomitant radiochemotherapy on overall survival (P=0.004), with an absolute benefit of 5.7% (from 18.1% to 23.8%) at 3 years and 4.5% at 5 years. There was no significant difference regarding acute pulmonary toxicity. Concomitant treatment decreased locoregional progression, but concomitant radiochemotherapy increased acute esophageal toxicity (grade 3-4) from 4% to 18%. This improved survival was accomplished because of an improved locoregional control. There were no significant differences between the regimens: single or double high-dose chemotherapy or daily low-dose cisplatin. No differences in distant metastasis rate were observed between the two approaches.

Within a few months a meta-analysis was published by O’Rourke et al. [25] reporting a 10% absolute survival benefit at two years. Six trials (1024 patients) of concurrent versus sequential chemoradiation were included. A significant benefit of concurrent treatment was shown in overall survival (hazard ratio-HR 0.74, 95% CI 0.62 to 0.89). More treatment-related deaths (4% vs 2%) in the concurrent arm without statistical significance (Relative Risk-RR 2.02, 95% CI 0.90 to 4.52). There was increased severe esophagitis with concurrent treatment (RR 4.96, 95%CI 2.17 to 11.37). The most important acute but manageable side effect was esophagitis grade 3 to 4 in 18% of the patients treated with concurrent radiochemotherapy versus 4% in the patients treated with sequential arm.

The role of timing and sequencing the treatment may also depend on the tumor type, the degree of oxygenation of tumor cells and other biochemical processes occurring during radiation. In clinical practice, a compromise option is the alternation of radiotherapy and chemotherapy, for example by insertion of radiotherapy after 2-3 cycles of chemotherapy. The clinical efficacy of this strategy has, however, not been verified in prospective clinical studies. Concurrent chemoradiation is at present the treatment of choice for patients with locally advanced NSCLC. However, due to its higher toxicity, this combination is mostly restricted to patients in a good general condition, minimal comorbidity and who are relatively young [26-29]. There is a question what proportion of patients would be suitable for concurrent chemoradiation. We found only one report on a population-based study that prospectively evaluated comorbidities in all patients diagnosed with lung cancer, stage III for NSCLC [30]. In this prospective, population-based study, more than half of the patients with stage III NSCLC were not eligible for concurrent chemoradiation.

<table>
<thead>
<tr>
<th>Authors</th>
<th>Study Design</th>
<th>Patient Characteristics</th>
<th>Radiotherapy</th>
<th>Chemotherapy</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Auperin et al. [24]</td>
<td>Meta-analysis</td>
<td>n=1295</td>
<td>Inoperable stage I, II, III and IV; ECOG 0-2</td>
<td>RT 60 and 66 Gy in two trial each, and 56 and 48.5 Gy in one trial each. In one trial the radiotherapy differed in the two arms—there was a 10 days split in the concomitant arm. In the five trials in the sequential arm, patients randomly assigned to concomitant arm received radiotherapy more frequently than those randomly assigned to the sequential arm.</td>
<td>Eleven randomised studies, sequential -cisplatin combined with one drug in four trials, or with two drugs in two trials. Vinorelbine or gemcitabine were used in two trials.</td>
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<tr>
<td>O'Rourke et al. [25]</td>
<td>Randomised control trial</td>
<td>RT vs RT and CT</td>
<td>Inoperable stage I, II, III; ECOG 0-3</td>
<td>RT 56 Gy/28 fractions- 70.2 Gy/39 fractions</td>
<td>Nineteen randomised studies, vindesine/cisplatin/mitomycin C; cisplatin/vinblastine; cisplatin/vinorelbine; carboplatin/paclitaxel</td>
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<tr>
<td>O'Rourke et al. [25]</td>
<td>Randomised control trial</td>
<td>Sequential RT and CT vs concomitant RT and CT</td>
<td>Inoperable stage I, II, III; ECOG 0-3</td>
<td>RT- 56 Gy/28 fractions- 70.2 Gy/39 fractions</td>
<td>Six randomised trials, vindesine/cisplatin/mitomycin C; cisplatin/vinorelbine; carboplatin/paclitaxel</td>
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Table 1: Chemoradiotherapy in locally advanced non-small cell lung cancer.
Targeted Therapies and Radiotherapy in Locally Advanced Non-Small Cell Lung Cancer

Targeted therapies represent a new class of drugs which interfere with specific molecular targets (typically proteins) playing critical roles in tumor growth and progression. The approved targeted therapies in lung cancer include erlotinib, gefitinib (a small-molecule tyrosine kinase inhibitor) and bevacizumab (a monoclonal anti-VEGF antibody). The accepted dogma is that antiangiogenic therapy destroys or blocks the function of tumor-associated vessels to deprive the tumor of oxygen and nutrients, thereby inhibiting tumor growth. Numerous preclinical studies indicated synergistic activity of various antiangiogenic or antivascular therapies with single-dose or fractionated radiotherapy in human and murine tumors [34,35]. However, since multiple variables contribute to the sensitivity of tumors to radiation or antiangiogenic treatment, the most effective way of their combining is virtually unknown [36,37]. Blocking survival signalling in endothelial cells after irradiation seems to increase the radiation response considerably [38]. Moreover, sensitization of endothelial cells just before exposure to radiation may be the most effective way to improve response of tumor cells to radiation [39,40]. On the other hand, induction of hypoxia via blood vessel damage may potentially induce radioprotection of the tumor. A logical and clearly proven premise for optimal multimodality therapy is therefore necessary for efficient translation of promising preclinical strategies into clinical applications. Many new biologicals have entered the therapeutic domain, several were combined with concurrent RCT regimens. Some combinations appear to be too toxic like the vascular epithelial growth factor antibody bevacizumab [41]. Also, the use of bevacizumab and erlotinib is not recommended given the lack of an efficacy signal and the substantial risk of esophageal toxicity [42] (Table 2).

<table>
<thead>
<tr>
<th>Reference (therapy)</th>
<th>Study type</th>
<th>Patients</th>
<th>RT schedule</th>
<th>Systemic therapy</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Spigel et al. [41]</td>
<td>phase II</td>
<td>n=29 (small cell) n=6 non-small cell lung cancer trial included patients with unselectable stage III nonsquamous without pleural or pericardial effusions; ECOG 0-1;</td>
<td>RT began with cycle 3, at a dose of 1.8 Gy/d to a total of 61.2 Gy</td>
<td>induction treatment included: carboplatin AUC = 5, pemetrexed 500 mg/m2, and bevacizumab 15 mg/kg; consolidative therapy with carboplatin AUC = 6, pemetrexed 500 mg/m2, and bevacizumab 15 mg/kg</td>
<td>the trial’s primary PFS end point could not be assessed due to early trial closure because of toxicity. the objective response rate was 66%</td>
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<tr>
<td>Socinski et al. [42]</td>
<td>phase I/II</td>
<td>n=45 medically inoperable or unresectable, WHO 0-1; stage IIIA or IIIB,</td>
<td>conformal radiation therapy to 74 Gy</td>
<td>induction chemotherapy (carboplatin AUC 6, paclitaxel 225 mg/m2, and bevacizumab 15 mg/kg followed by concurrent chemotherapy (carboplatin AUC 2 and paclitaxel 45 mg/m2 weekly with bevacizumab 10 mg/kg) in the phase I portion, cohort 1 received no erlotinib, whereas cohorts 2 and 3 received erlotinib at 100 and 150 mg, respectively. consolidation therapy with erlotinib (150 mg daily) and bevacizumab (15 mg/kg every 3 weeks) 3 to 6 weeks later for 6 cycles</td>
<td>the objective response rates to induction and overall treatment were 39% and 60%. median PFS 10.2 mo median OS 18.4 mo</td>
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<td>Kelly et al. [44]</td>
<td>phase III</td>
<td>n=243 inoperable stage IIIA or IIIB without pleural or pericardial effusions; ECOG 0-1;</td>
<td>initial field received 1.8 Gy/d for 5 weeks for a dose of 45 Gy; an additional radiation boost to gross disease with 2 Gy/d to 18 Gy was delivered without a break. The total radiation dose received was 61 Gy.</td>
<td>concurrent cisplatin and etoposide with thoracic radiation. cisplatin 50 mg/m2 with etoposide 50 mg/m2. 4 to 8 weeks after completion of radiation, patients without progressive disease received 3 cycles of docetaxel 75 mg/m2 3 to 6 weeks after docetaxel, patients received gefitinib 500 mg or placebo orally, once a day for 5 years or until disease progression or intolerable toxicity. Later gefitinib was amended to the 250 mg/d</td>
<td>median survival time was 23 months for gefitinib (n=118) and 35 months for placebo the toxic death rate was 2% with gefitinib compared with 0% for placebo.</td>
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<tr>
<td>Study</td>
<td>Phase</td>
<td>Therapies</td>
<td>Treatment Details</td>
<td>Toxicities</td>
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<tr>
<td>Martinez et al. [52]</td>
<td>phase II</td>
<td>erlotinib</td>
<td>unrectateble stage I-IIIA, not suitable to receive chemotherapy. ECOG 0-2</td>
<td>66 Gy in 33 fractions during 6 weeks. RT and placebo (arm a) or concomitant erlotinib 150 mg/day po maintained for 6 months (arm b). esophagitis 40% in arm A and 23% in arm B, (no grade 3-4). radiodermitis 50% in arm A (no grade 3-4 observed) and 8% in arm B, being grade 3. pneumonitis 20% in arm A (10% grade 3) and 8% in arm B (no grade 3-4 observed). main toxicities related to erlotinib were skin rash (61.5%) and diarrhea (23%). response rate in arm A was 55.5% and 83.3% in arm B. Disease progression is documented in 22.2% in arm A and 16.7% in arm B.</td>
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<td>Jensen et al. [57]</td>
<td>phase II</td>
<td>cetuximab</td>
<td>not candidates for concomitant chemoradiation (or refused), KPS at least 70, one of two trials with mandatory PET; stage IIIA or IIIB, no malignant pleural effusion. IMRT trial, 66 Gy in 33 daily fractions of 2 Gy, ENI to 50 Gy (or 40 depending on lung dose). cetuximab followed by 13 weekly consolidation cycles. median OS 19.6 mo, median PFS 8.5 mo, 63% PR, no CR,</td>
<td>cetuximab 400 mg/m(2) i.v. on day 1 followed by weekly cetuximab 250 mg/m(2) i.v. with concomitant radiation. median OS 15.1 mo, median PFS 7.2 mo, 26% PR, no CR,</td>
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<tr>
<td>Jatoi et al. [56]</td>
<td>phase II</td>
<td>Cetuximab</td>
<td>not candidates for concomitant chemoradiation, either age ≥ 65 years with ECOG 0-2 or younger but ECOG 2; stage IIIA or IIIB, no malignant pleural effusion,</td>
<td>RT 60 Gy in 30 daily fractions of 2 Gy, ENI to ipsilateral hilar and mediastinal nodes (44 Gy) cetuximab 400 mg/m(2) i.v. on day 1 followed by weekly cetuximab 250 mg/m(2) i.v. with concomitant radiation. median OS 17 mo, PFS NR, 16% PR and 7% CR at 12 months (NR at earlier time points), patterns of failure: 31% distant only, 23% local only, 7% regional only, 11% combinations of these,</td>
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<tr>
<td>Hallqvist et al. [58]</td>
<td>phase II</td>
<td>cetuximab</td>
<td>medically inoperable or unrectateble, WHO 0-1; stage IIIA or IIIB, no malignant pleural effusion with positive cytology,</td>
<td>RT 68 Gy in 34 daily fractions of 2 Gy, no ENI 2 cycles of induction cisplatin/docetaxel, cetuximab starting one week before RT. median OS NR, PFS NR, 58% PR, no CR,</td>
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<tr>
<td>Hughes et al. [59]</td>
<td>phase II</td>
<td>cetuximab</td>
<td>Inoperable, WHO 0-1; Stage IIIA or B, no pleural effusion</td>
<td>RT 64 Gy in 32 fractions of 2 Gy, in 4 cases ENI to ipsilateral hilar and mediastinal nodes (50 Gy) up to 4 cycles (median 3) of platinum-based induction CT, cetuximab starting one week before RT. median OS NR, PFS NR, 58% PR, no CR,</td>
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<tr>
<td>Blumenschein et al. [60]</td>
<td>phase II</td>
<td>cetuximab</td>
<td>Inoperable, Zubrod 0-1; Stage IIIA or B, weight loss &lt; 5%, FEV1 ≥ 1.2</td>
<td>RT 63 Gy in 35 fractions of 1.8 Gy, ENI to ipsilateral hilar and mediastinal nodes (45 Gy) cetuximab week 1-17, weekly carboplatin/paclitaxel during RT followed by 2 cycles consolidation cetuximab (7 weeks) plus 4 cycles carboplatin/ pemetrexed vs. CT without cetuximab (n = 48), afterwards 4 cycles of pemetrexed. median OS 22.7 mo, median time to progression around 14-15 mo, 29% CR, 33% PR,</td>
<td>median OS 22.7 mo, median PFS 11.9 mo, 5.7% CR, 30% PR,</td>
</tr>
<tr>
<td>Govindan et al. [61]</td>
<td>phase II</td>
<td>cetuximab</td>
<td>Inoperable, ECOG 0-1, one of two trials with mandatory PET; Stage IIIA or B, no pleural effusion, weight loss ≤ 10%</td>
<td>70 Gy in 35 fractions of 2 Gy, no ENI cetuximab (7 weeks) plus 4 cycles carboplatin/ pemetrexed vs. CT without cetuximab (n = 48), afterwards 4 cycles of pemetrexed. median OS 25.2 mo, median failure-free survival 12.3 mo, 4% CR, 68% PR,</td>
<td>median OS 25.2 mo, median failure-free survival 12.3 mo, 4% CR, 68% PR,</td>
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RT: radiotherapy; IMRT: intensity-modulated radiotherapy; CT: chemotherapy; KPS: Karnofsky performance status; ECOG: Eastern Cooperative Oncology Group performance status; WHO: World Health Organisation performance status; ENI: elective nodal irradiation; OS: overall survival; PFS: progression-free survival; PR and CR: partial and complete remission; NR: not reported; PET: positron emission tomography.

Table 2: Targeted therapies and radiotherapy in locally advanced non-small cell lung cancer.
Erlotinib and gefitinib are a small molecule inhibitor that reversibly targets the tyrosine kinase activity of the epidermal growth factor receptor (EGFR). EGFR is overexpressed and/or mutated in many cancer types, and its activation triggers pathways involved in cell growth and proliferation. Early clinical studies with gefitinib showed promising efficacy and mild toxicity in patients with advanced non-small-cell lung cancer (NSCLC). In clonogenic in vitro survival experiments, gefitinib had significant radiosensitizing effects on NSCLC cell lines [43]. Gefitinib enhances the radioreponse of NSCLC cells by suppressing cellular DNA repair capacity. But in unselected population, gefitinib did not improve survival [44]. The trial from the National Cancer Institute of Canada Clinical Trials Group showed that erlotinib monotherapy prolonged survival in patients with advanced NSCLC who had progressed after standard chemotherapy [45], and erlotinib is approved in this setting. Interestingly, EGFR expression does not seem to correlate with response to EGFR inhibitors but a recent analysis of data from this trial indicated that EGFR mutations and high copy number are predictive of response to erlotinib [46]. In addition, EGFR fluorescence in situ hybridization score was a significant predictive marker of differential survival benefit from erlotinib. Erlotinib-induced apoptosis was augmented by radiation in very high expression of HER1/EGFR cells only. In conclusion, high HER1/EGFR expression may result in a high degree of radiosensitization with erlotinib combined with radiation [47]. A strong rationale may exist for combining erlotinib with RT. Erlotinib helps disrupt cell growth pathways and enhances the sensitivity of cells to the effects of RT [48-50]. It is also possible that RT enhances the effectiveness of erlotinib by cytoreducing the tumor and creating a hypoxic environment [51]. Several studies in NSCLC were made to evaluate erlotinib in combination with RT. A prospective phase II study found that RT and concurrent erlotinib used in the treatment of patients with unresectable NSCLC shows promising results without an increase in toxicity [52]. Adverse events related to RT included esophagitis, radiation dermatitis, and pneumonitis. The addition of erlotinib to RT did not appear to increase RT-associated toxicities. Erlotinib-related adverse events included mild to moderate skin rash (61.5%) and diarrhea (23%). The RR was 55.5% in the RT-alone arm compared with 83.3% in the erlotinib arm. The Cancer and Leukemia Group B is conducting a phase II trial, CALGB 30605, of paclitaxel followed by RT and erlotinib in patients with unresectable stage III NSCLC. The study is evaluating induction chemotherapy consisting of paclitaxel and carboplatin. Patients with no disease progression outside the planned radiation field will continue to receive concurrent erlotinib and RT. Results from current studies are eagerly awaited.

Several drugs interfering with the EGFR signaling pathway have been developed e.g. cetuximab (a human-murine chimeric IgG1 monoclonal antibody that binds to the extracellular region of the EGFR). Under experimental laboratory conditions in animal models, cetuximab increases tumour radiocurability (fractionated and single dose irradiation) [53,54]. The feasibility of adding the epidermal growth factor receptor inhibitor cetuximab has been recently reported for NSCLC patients [55]. We found a few phase II clinical trials of cetuximab combined with radiotherapy for non-small cell lung cancer. Two of them have combined RT and cetuximab without any chemotherapy in patients who are not candidates for chemoradiation [56,57]. Combined radioimmunotherapy with cetuximab was safe and feasible, especially in elderly patients with multiple comorbidities. Another studies of them included patients with inoperable stage III disease and good performance status after the induction chemotherapy [58,59]. Induction chemotherapy followed by concurrent cetuximab and RT to 68 Gy was clearly feasible with promising survival. Toxicity, like pneumonitis and esophagitis was low compared to most schedules with concurrent chemotherapy. The last study has published by Radiation Therapy Oncology Group (RTOG) was a phase II study of chemoradiotherapy with carboplatin and paclitaxel plus cetuximab in patients with stage III NSCLC [60]. The combination of cetuximab with CRT is feasible and shows promising activity. The overall survival achieved with this regimen was longer than any previously reported by the Radiation Therapy Oncology Group with median survival 22.7 months, and 24-month overall survival - 49.3%. The second trial in this category with several important differences (mandatory PET, higher radiation dose of 70 Gy, only 7 weeks of cetuximab concomitant to RT, chemotherapy with carboplatin and pemetrexed) was done [61]. Median survival was 25.2 months and failure-free survival 12.3 months.

Until now no definite data can be reported. Further basic research and appropriately designed clinical studies are clearly needed to optimize scheduling of combined radiation and molecular targeted therapies. The results of the published clinical trials (one of them was a phase III study) suggest that larger randomized trials are warranted. It is very important to include the right patient population especially patients with the right genetics/mutations for these clinical trials.

Conclusions

Patients with stage III disease differ with regard to primary tumour volume and proximity/infiltration to surrounding structures, extent of lymphatic spread, cancer biology, and host factors such as age, cardiopulmonary function and other comorbidity. Treatment recommendations have to take into account these differences and stratify patients according to technical resectability, ability to tolerate high-dose radiotherapy and chemotherapy, and many more. Many patients with inoperable stage III disease are candidates for combined modality chemo- and radiotherapy (RT). In conclusion after two decades of mainly sequentially combined treatment, concurrent radiochemotherapy is nowadays the standard treatment of locally advanced NSCLC. However, there are some doubts.

Firstly, it should be realized that the trial data were collected in a period before routine staging with FDG-PET and MRI of the brain. The routine use of these tests definitely changes the population of patients enrolled in radiochemotherapy.

Secondly, the other topics for future research are RCT with more sophisticated radiotherapy techniques allowing possibly higher tumour doses and/or lower toxicities in surrounding healthy tissues. For patients with larger tumor volumes, the possibilities to increase the radiation dose were limited by normal tissue constraints (esophagus and spinal cord). Conventionally fractionated radiotherapy for stage I NSCLC has shown inferior outcomes than surgery and these results are linked to insufficient radiation doses. After the impact of RT dose for lung cancer was established, a number of trials were structured in the quest for better local control and overall survival by either dose escalation or shortening the total treatment time through conventional/ altered fractionation, even in combination with chemotherapy. The delivery of 60 Gy resulted in a 5-year survival rate of 38% for patients with primary tumours less than 2 cm in size, 22% for tumours 2-3 cm in size, 5% for tumours 3–4 cm in size, and 0% for larger tumours [62]. Based on biological and statistical modelling of tumour responses to various radiation dose levels, it has been shown that doses as high as 80 to 90 Gy ensure a progression-free survival rate of 50% [63]. The majority of studies concluded that patients receiving higher radiation doses have better treatment outcomes [64,65].
New technical advances in the application of RT enhanced the ability of targeted treatment and sparing of normal tissues, making high BED studies possible. Intensity-modulated radiotherapy (IMRT) has the potential benefit to further increase the dose that can be safely prescribed in lung cancer patients due to a better conformity index [66-68]. In Stereotactic Radiation Therapy (SABR), high doses per limited number of fractions are used, although the actual biologically equivalent dose (BED) for the eradication is not yet completely understood [69]. When a sufficient dose (BED ≥ 100 Gy) is used, it has been noted in most clinical studies that the success rate of local control is over 90%. In particular, the surprising results of the RTOG 0617 trial [70] drove attention to the importance of adverse effects, once again emphasising that future research should focus on quality of life.

Thirdly, in most of these studies authors have not taken in the analysis factors such as histological type, age, comorbid conditions. Since the incidence of NSCLC is high among elderly patients and many of them have a smoking history, the majority has severe comorbidities. Therefore, aggressive combined modality treatment might be contraindicated on priority with every patient. However, age is not an independent prognostic factor in stage III and IV NSCLC and epidemiological studies show that, with increasing age, the percentage of people treated with chemotherapy decreases [71-73]. Elderly patients with marginal renal function (creatinine clearance <70mL/min) or marginal cardiac function are eligible for administration of daily low-dose cisplatin, while administration of full-dose chemotherapy is often contraindicated. Combination of concurrent daily cisplatin with radiation appears to be a good alternative, especially in these elderly, frail patients [74,75]. Also preclinical studies on RCT support the use of daily administration for optimal radiosensitizing effects [76]. This approach, delivered in a short overall treatment time, is suitable for both the elderly and for patients with comorbidities. It also offers the opportunity to combine concomitant radiochemotherapy with new agents. Existing data concerning targeted therapies in conjunction with radiotherapy are inconsistent and do not allow for firm conclusions. The optimal timing of the administration of RT and EGFR kinase inhibitors has yet to be determined. Strategies to safely incorporate novel antiangiogenic agents into combined-modality therapy in lung cancer are needed. The studies using targeted therapies in particular address their optimal integration with radiotherapy are still in their infancy. The rapid development of molecular oncology will hopefully contribute to a better patient selection to particular strategies and to treatment optimization.

Appendix

The information was gathered from extensive PubMed searches (no limits to publication period were applied but only English-language papers are referenced). Additional references, including congress abstract presentations, are included where appropriate and in particular where there are no published studies on a discussion topic.

References


