Treatment-Related Predictive and Prognostic Factors in Trimodality Approach in Stage IIIA/N2 Non-Small Cell Lung Cancer

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While there are no established pretreatment predictive and prognostic factors in patients with stage IIIA/pN2 non-small cell lung cancer (NSCLC) indicating a benefit to surgery as a part of trimodality approach, little is known about treatment-related predictive and prognostic factors in this setting. A literature search was conducted to identify possible treatment-related predictive and prognostic factors for patients for whom trimodality approach was reported on. Overall survival was the primary endpoint of this study. Of 30 identified studies, there were two phase II studies, 5 “prospective” studies, and 23 retrospective studies. No study was found which specifically looked at treatment-related predictive factors of improved outcomes in trimodality treatment. Of potential treatment-related prognostic factors, the least frequently analyzed factors among 30 available studies were overall pathologic stage after preoperative treatment and UICC downstaging. Evaluation of treatment response before surgery and by pathologic tumor stage after induction therapy were analyzed in slightly more than 40% of studies and found not to influence survival. More frequently studied factors—resection status, degree of tumor regression, and pathologic nodal stage after induction therapy as well as the most frequently studied factor, the treatment (in almost 75% studies)—showed no discernible impact on survival, due to conflicting results. Currently, it is impossible to identify any treatment-related predictive or prognostic factors for selecting surgery in the treatment of patients with stage IIIA/pN2 NSCLC.

Keywords: predictive factors, prognostic factors, stage IIIA/pN2, non-small cell lung cancer, trimodality therapy

INTRODUCTION

Stage III non-small cell lung cancer (NSCLC) is a heterogeneous disease presentation based on the range of patient and tumor characteristics included under its heading. Concurrent radiochemotherapy (RT-CHT) is the standard treatment approach for locally advanced stage III NSCLC (1–3). In patients with stage IIIA/pN2 disease, thoracic oncologists have explored various means of optimizing treatment approach. Treatment regimens consisting of surgery alone or in combination with adjuvant CHT- and/or RT have been historically among the most common approaches used,
with recent decades showing a trend for induction regimens followed by surgery. The latter have included induction CHT followed by surgery (4–7), and, in more recent years, induction RT-CHT followed by surgery (8–34). When tested in prospective randomized phase III studies, preoperative RT (PORT)-CHT (with or without preceding CHT) followed by surgery brought no improvement in overall survival (OS) or local control compared to definitive concurrent RT-CHT, and was associated also with an increase in treatment-related mortality (8, 10, 35, 36).

Notwithstanding this high level evidence, there exists reluctance among many clinicians to rule out the use of surgery in stage III NSCLC. For example, in an unplanned subgroup analysis of the Intergroup 0139 trial (10), it appeared that patients who underwent lobectomy as a component of their trimodality regimen had improved OS compared to the patients who received definitive RT-CHT. Given the controversial role of surgery in the management of this disease, looking at patient, tumor, and treatment variables from the range of studies in stage III NSCLC might then provide guidance as to which might favor including surgery as part of patient care.

Recently, a survey of the literature tested the role of surgical treatment in the care of stage III NSCLC patients (37). Important finding was that there were no pretreatment factors that identified any patient subgroup whose outcomes were improved by the addition of surgery. In addition, the majority of studies did not consistently evaluate prognostic factors that may have had an impact on treatment outcomes, but when done, none of the three most frequently analyzed factors (age, gender, histology) was found to clearly influence survival. From this, it was concluded that there were no specific pretreatment factors which favorably selected patients for surgery as a part of trimodality program (over definitive concurrent RT-CHT).

While little is known about treatment-related factors, the aim of the present study is to identify if potential treatment-related (i.e., after completion of induction therapy) predictive and prognostic factors exist that can help optimize the selection of patients for a trimodality approach for their stage IIIA/pN2 disease.

**MATERIALS AND METHODS**

References were identified through a literature search using PubMed, Google Scholar, EMBASE the terms “non-small cell lung cancer,” “surgery,” “radiotherapy,” “chemotherapy,” and “Stage IIIA” for the interval from 1990 until October 2017 as well as through searches of the references of the identified articles as well as of the authors’ own files. The year 1990 was selected as the starting point for the survey to mark for the contemporary era of RT delivery. Findings were restricted to English language publications in which potential treatment-related predictive and prognostic factors in the trimodality setting of concurrent RT-CHT (with or without preceding CHT) followed by surgery primarily focusing on Stage IIIA/pN2 (Stage III) were investigated. We principally sought out the highest level of evidence (from prospective randomized phase III trials). Upon finding none, the search was expanded to include prospective research (formal phase II or studies described only as “prospective”), and following that, all available retrospective studies using a trimodality approach meeting the criteria detailed above. Due to substantial heterogeneity in the sources, studies invariably included those with a mixture of patients (i.e., with both early and late stages NSCLC); they have also included patients treated with a variety of adjuvant therapies [e.g., PORT, or postoperative CHT (POCHT)]. Studies involving patients with superior sulcus/Pancoast/thoracic inlet tumors were excluded from the search. No study was excluded based on the form of induction CHT (drugs, regimens, administration, number of cycles), the form of RT (total doses delivered, once- or twice-daily fractionation, split- or continuous course), the type of surgery (pneumonectomy, bilobectomy, lobectomy) or adjuvant therapies (PORT or POCHT) employed. Any potential predictive or prognostic factor that was considered in a given study was included in the analysis.

Using Clark methodology (38, 39) predictive factor was defined as the one which defines the effect of treatment on the tumor, while a prognostic factor was defined as one which defines the independent impact of a given variable on the patient outcome. If one attempts to meaningfully use such predictors to guide the selection patients for trimodality approach, evaluation of clinically relevant endpoints (e.g., survival) should be used while the independent influence of these factors on a relevant endpoint (e.g., survival) distinguished from their ability to predict a differential clinical benefit from the specific treatment (e.g., PORT-CHT followed by surgery) (38, 39).

The purpose of this analysis is to determine if the survival benefit of patients treated surgically relative to patients treated with exclusive concurrent RT-CHT differs by any treatment-related characteristic, and thus establish the significance of a potential predictor. To establish the significance of such potential prognostic factors, it would be necessary to demonstrate its independent influence on treatment outcome. The current analysis was, therefore, restricted to studies that have provided such data and excluded studies that either did not discuss these or those in which they were described in univariate analyses only. Also, when the study using both univariate and multivariate analysis (MVA) disclosed factors that were not shown to influence survival on univariate analysis (and, therefore, subsequently, not entered into a multivariate model), such study was excluded from consideration. Finally, we excluded from consideration all studies that used multivariate analyses for subgroups of patients that were a part of the whole cohort of patients entering the published study.

In both prospective and retrospective series, multivariate analyses are used to investigate (treatment-related) prognostic factors, while the detection of treatment-related predictive factors requires more stringent criteria (38, 39). For the latter, one must determine if the survival benefit of patients on the investigational arm relative to patients on the control arm differ by treatment characteristic status. As an example, when survival by treatment arm for patients with documented mediastinal downstaging is compared to that for patients without this finding, one may find that the relative treatment benefit is similar. Then, looking at the survival results for all combinations of treatments and in whom documented mediastinal downstaging or not is determined, and if tests for interaction between mediastinal downstaging and treatment is not statistically significant, one may conclude that
although mediastinal downstaging is a strong prognostic factor, it does not predict a differential survival benefit between the experimental treatment and the control treatment dependent on mediastinal downstaging.

Overall survival was the primary endpoint in this analysis and was provided in all identified studies. Loco-regional and distant metastasis-free survival as well as treatment-related toxicity, including serious adverse events, were not considered endpoints for this study, since they were absent in some of the identified studies. When presenting the study finding we used a qualitative approach (yes vs no) (Table 1). Tests of significance for a given study were not included in our summary table. Individual patient data was not sought for this analysis.

RESULTS

Using Clark methodology (38, 39), we did not identify any study investigating treatment-related predictive factors for the potential superiority of surgical outcomes in patients with Stage IIIA (mostly pN2) NSCLC. A total of 30 studies which provided information on prognostic factors in trimodality studies using MVA (Cox proportional hazards model) were found for the interval of interest. Of these, there were two prospective phase II studies (16, 40), five prospective studies (11, 12, 18, 22, 41), and 23 retrospective studies (13–15, 20, 21, 23–32, 34, 42–48) (Table 1). Of these 30 studies, 17 were single arm studies, while the remaining 13 were two-arm studies. Of the latter, five studies compared concurrent RT-CHT vs a trimodality approach, one compared surgery alone with a trimodality approach, while the remaining seven studies compared induction CHT followed by surgery with induction RT-CHT followed by surgery. Of the total of 30 studies, 12 studies either allowed (in cases of high-risk features) or mandated (irrespective of risk features) the use of various forms of adjuvant or postoperative treatment. This included PORT alone, POCHT alone as well as postoperative RT-CHT. The total number of patients in these 30 studies was 3,397, of which 3,071 (90%) patients underwent surgical resection. Total of 14 studies included Stages IIIA and IIIB patients, three studies included patients with earlier (IB, II, and IIB) and/or higher (IV) stages. The 13 remaining studies defined staging through status confirming N2 may have had worse prognosis than those with pN2 found at surgery and, possibly, could have different predictive and prognostic factors in the trimodality setting of Stage IIIA NSCLC.

DiSCUSSiON

The present study looked at treatment-related factors that could aid in optimizing patient selection in the setting of trimodality treatment approach for Stage IIIA (primarily pN2) NSCLC. Similar to a recent study on pretreatment predictive and prognostic factors (37), we first noted inconsistent use of clinical definitions in this setting by clinicians. While we employed the definition of Clark (38, 39) to identify potential predictive factors, we found that that term was inaccurately used in most studies which then led to their incorrect inclusion in MVAs. In other words, MVAs reveal prognostic factors and not predictive ones.

Second, we found wide heterogeneity among studies with respect to diagnostic, staging, and treatment procedures, as well as different systems used to detect tumor response in surgical specimens. This led to significant inconsistencies in how potential treatment-related factors could be investigated. In particular, the majority of studies have used mediastinoscopy to prove pN2 while in some studies pathological mediastinal disease was found during surgery. It introduces potential bias in interpretation of the study results as patients with preoperatively confirmed N2 may have had worse prognosis than those with pN2 found at surgery and, possibly, could have different predictive and prognostic factors in the trimodality setting of Stage IIIA NSCLC.

Mindful of these shortcomings, we were unable to identify any prospective randomized study which investigated predictive factors in the setting of trimodality therapy, or the group of retrospective studies which compared bi- and trimodality approach. Third, we found that potential treatment-related prognostic factors were not consistently evaluated. Factors, such as post-induction therapy staging (ypStaging) or UICC downstaging, were looked at in less than 20% of all eligible studies; while factors, such as the degree of tumor regression, status of surgical margins, evaluation of response before surgery, or post-induction tumor status (ypT), were evaluated slightly more than 40% of all available studies. This limited our ability to detect the potential influence of any of these factors on treatment outcome. The factor more often investigated, post-induction nodal status (ypN), presented the same difficulty. No firm conclusions could be drawn possibly because of inconsistent definitions used for this across studies: some investigators used different comparators, either focusing on mediastinal nodal clearance (i.e., ypN0-1 vs ypN2), or only on ypN0 vs ypN+ or simply testing for the independent influence of a variable ypN (yes vs no).

The variable use of adjuvant treatments further contributed to the difficulty in assessing post-induction factors: while the majority of studies did not involve their use, for those that did,
| Reference          | Study                        | Response before surgery* | Treatment* (all possible variables) | Resection status | Degree of tumor regression* | yp Stage | ypT* | ypN | UICC downstaging |
|--------------------|------------------------------|--------------------------|------------------------------------|-----------------|----------------------------|----------|------|-----|------------------|
| Adelstein et al. (16) | Prospective phase II         | No—clinical              | Not evaluated                       | Not evaluated   | Yes—pCR or pPR vs pNR     | Not evaluated | Not evaluated | No—pN0 | Not evaluated |
| Park et al. (40)    | Not evaluated                | No—R0                    | Not evaluated                       | Yes (R0 vs R+)  | No (resectability)        | Yes—pN0-1 | Not evaluated |       |                 |
| Choi et al. (11)    | Prospective                  | No                       | Yes (CR—R0)                         | Not evaluated   | Yes (yp0 vs ypN1-2)       | Not evaluated | Not evaluated |       |                 |
| Thomas et al. (12)  | Not evaluated                | Yes (R0 vs R+)           | Yes (>90 vs <90%)                   | Not evaluated   | Yes (ypN0 vs ypN1-2)      | Not evaluated | Not evaluated |       |                 |
| Pezzetta et al. (22) | No—induction CHT vs induction RT-CHT | No—ypT downstaging       | No—pT0-pT4                         |                |                            |           | Not evaluated |       |                 |
| Kim et al. (18)     | No—number of CHT cycles      | No—(resectability)       | Yes                                 | No—pT0-pT4     |                            |           | Not evaluated |       |                 |
| Isobe et al. (41)   | Not (clinical response)      | No (resectability)       | Yes                                 | No (pN0 vs pN2-3) |                            |           | Not evaluated |       |                 |
| Cyjon et al. (20)   | Retrospective                | No                       | Yes—CR (R0)                         | Not evaluated   |                            |           | Not evaluated |       |                 |
| Sawabata et al. (21)| Not evaluated                | No—R0 vs R1              | No—pT0-pT4                         | Yes—pN0-pN2 (single) vs pN2 (multiple)-N3 | Not evaluated | Not evaluated |       |                 |
| Takeda et al. (13) | No                           | Yes—surgery; no—CHT      | No—Regression score >IIb (<10% viable tumor tissue) | No—pT0-pT4     |                            |           | Not evaluated |       |                 |
| Caglar et al. (14)  | Not evaluated                | Yes—pN0-1 vs pN2 (medialstinal downstaging) | Not evaluated | No—pT0-pT4     |                            |           | Not evaluated |       |                 |
| Steger et al. (15)  | No—Extent of surgery (pneumonectomy) | No—ypT downstaging       | No—ypT (pneumonectomy)              |                |                            |           | Not evaluated |       |                 |
| Li (23)             | No (CR-PR vs SD-PD)          | No—RT; no—POCHT          | Yes—pCR or pPR vs pNR              |                |                            |           | Not evaluated |       |                 |
| Li J (24)           | Not evaluated                |                           |                                     |                |                            |           | Not evaluated |       |                 |
| Weder et al (25)    | No                           | No—type of surgery       | No—pStage                           |                |                            |           | Not evaluated |       |                 |
| Meacci et al. (26)  | No—extent of surgery         | Yes (R0 vs R1)           | Not evaluated                       |                |                            |           | Not evaluated |       |                 |
| Kim et al (27)      | No—RT, No—CHT; no—extent of surgery | No—ypN0-1 vs pN2 (medialstinal downstaging) | Not evaluated |                |                            |           | Not evaluated |       |                 |
| Shintani et al. (29)| No                           | Not evaluated             |                                     |                |                            |           | Not evaluated |       |                 |

(Continued)
| Reference          | Study                          | Response before surgery | Treatment* (all possible variables) | Resection status | Degree of tumor regression* | yp Stage | ypT* | ypN | UICC downstaging |
|--------------------|--------------------------------|-------------------------|-------------------------------------|------------------|-----------------------------|----------|------|-----|------------------|
| Steger et al. (30) |                                | Not evaluated           | Yes (trimodality superior to surgery only) | No—ypT0N0M0 | No | No | Yes—ypUICC |
| Gomez-Caro et al. (31) |                               |                         | No—pStage (I-II/III-IV) | No—yp1-2 vs ypT3-4 | No | Not evaluated | Not evaluated |
| Toyooka et al. (32) |                                |                         | Yes—(induction CRT superior to induction CT); no—extent of surgery | No—pCR (+ vs -) | No | ypT | Yes—yp0-1 vs ypN2 (mediastinal downstaging) |
| Askoxylakis et al. (42) |                               |                         | No—induction CHT; no—non-CDDP | No | Not evaluated | No | Not evaluated |
| Lee et al. (43)     |                                |                         | Yes—type of CHT; yes—type of surgery (worse for pneumonectomy); yes—PORT; POCHT | No—resection margin (negative vs close/positive) | No | ypT0 vs ypT1-4 | Yes—ypN0 vs ypN+ |
| Renaud et al. (45)  |                                |                         | No (pN downstaging after induction treatment) | No (type of surgery) | No | Yes |
| Lim et al (34)      |                                |                         | No—clinical response; no—tumor size on post-induction treatment; yes—% change of target lesions on post-induction treatment | No—type of surgery; no—adjuvant treatment | No | No—total tumor size on surgical specimen; yes—viable tumor size on surgical specimen | No—pN0, pN1, pN2 |
| Shien et al. (46)   |                                |                         | No (radiological response) | No (type of surgery) | No (pCR) | No (pT3 vs pT0-2) | No (pN0-1 vs pN2-3) |
| Darling et al. (47) |                                |                         | Not evaluated | Yes (surgery) | Not evaluated | Not evaluated | Not evaluated |
| Kim et al. (44)     |                                |                         | Not evaluated | Yes—type of surgery (worse for pneumonectomy) | Yes—close or positive margins | Yes—advanced pT | Yes—persistent N2 |
| Yang et al. (48)    |                                |                         | Yes (surgery) | No (type of surgery) | No | Not evaluated | Not evaluated |
| Summary             |                                |                         | 1/13/18 | 10/19/10 | 8/4/18 | 5/5/21 | 0/5/25 | 2/11/18 | 11/8/11 | 2/0/28 |

*More than one finding per study (the same for both RT, chemo, surgery).

n.e., not evaluated; UICC, International Union Against Cancer; yp, pathological evaluation done after induction treatment (on operative specimen); pCR, pathological complete response; pPR, pathological partial response; pNR, pathological no response; R0, negative surgical margins; R1, microscopically positive resected tumor margins; R+, positive resected tumor margins; RT, radiotherapy; CHT, chemotherapy; PORT, postoperative RT; POCHT, postoperative CHT.
some allowed it in high-risk patients while others mandated their use, irrespective of surgical/pathological findings. It is possible that by allowing post-surgical therapy to be employed, the natural course of the disease treated with trimodality therapy was improved by, perhaps, improving both locoregional tumor control (with PORT) and/or microscopic tumor burden (with POCHT), and thus potentially achieving superior results when compared to those studies using no adjuvant therapy. Indeed, while resection status has long been considered as one of the major factors in predicting poor surgical outcomes, such that R0 is considered superior to any R+ status, we observed no such consistent association between studies and could not establish the importance of this factor.

Radiographic (clinical) response evaluations after induction therapy had a limited role in our analysis. In that regard, it is generally accepted that these are unreliable measures of the benefits of therapy as they are easily confounded. Contemporary studies that use novel approaches (e.g., PET-CT based response evaluations) and which could help in identifying such factors remain to be published. Future studies are required that incorporate this form of imaging which is then correlated with surgical/pathological findings.

The form of the surgical approach (all approaches considered) was also shown not to influence survival. The extent of surgery (pneumonectomy vs lesser operations) may simply be a surrogate for the bulk of the disease, so that the larger tumor burden by its nature carries the worse prognosis independent of surgical approach. A consistent definition of tumor burden prior to surgical intervention would clarify the impact of surgery in future studies where pneumonectomy is allowed since the location of the primary and nodal disease. This seems to be very important aspects, especially after Intergroup 0139 study results became available (10) showing worse survival for patients treated with trimodality approach using pneumonectomy vs lobectomy. Likewise, consistent stratification factors and trial eligibility criteria across studies and between research consortia would help better interpret conflicting results such as those in the histological domain which in EORTC study (8) favored squamous cell carcinoma while in Nordic study (36) favored adenocarcinoma.

The findings of the current study highlight our inability to use published study-derived treatment-related factors as either predictors or prognosticators to evaluate their impact in this clinical setting and to aid in the decision-making process regarding the appropriateness of surgery in stage III NSCLC. They follow our previous effort which focused on the pretreatment patient and tumor predictive and/or prognostic factors in this setting (37). Similar to it (37), although with slightly different number of included studies, the current one also could not identify such factors. This brings significant uncertainty in the process of decision-making before and after induction treatment was administered. It also reemphasizes the need for a more universal approach to defining and exploring treatment-related predictive and prognostic factor analysis in future studies, which needs to be coordinated across groups and countries. Possible avenues for clinical research in this setting would include more PRCTs with statistical analysis including predictive factor analysis being mandatory. In addition, a number of studies identified treatment-related factors that need to be further tested for its predictive and/or prognostic influence in this setting. This could be done as a preliminary step in the research scenario and, if repeatedly positive, across of various groups and institutions as to identify likely candidates for entering more patient and/or treatment-shaped trial milieu.

**AUTHOR CONTRIBUTIONS**

BJ: study design; literature search, data collection, manuscript draft, and final approval of the manuscript. FC, PD, AG-C, NC, GV, and II: literature search, data collection, manuscript draft, and final approval of the manuscript.

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**Conflict of Interest Statement:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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