Early death during chemotherapy in patients with small-cell lung cancer: derivation of a prognostic index for toxic death and progression

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Summary Based on an increased frequency of early death (death within the first treatment cycle) in our two latest randomized trials of combination chemotherapy in small-cell lung cancer (SCLC), we wanted to identify patients at risk of early non-toxic death (ENTD) and early toxic death (ETD). Data were stored in a database and logistic regression analyses were performed to identify predictive factors for early death. During the first cycle, 118 out of 937 patients (12.6%) died. In 38 patients (4%), the cause of death was sepsis. Significant risk factors were age, performance status (PS), lactate dehydrogenase (LDH) and treatment with epipodophyllotoxins and platinum in the first cycle (EP). Risk factors for ENTD were age, PS and LDH. Extensive stage had a hazard ratio of 1.9 ($P = 0.07$). Risk factors for ETD were EP, PS and LDH, whereas age and stage were not. For EP, the hazard ratio was as high as 6.7 ($P = 0.0001$). We introduced a simple prognostic algorithm including performance status, LDH and age. Using a prognostic algorithm to exclude poor-risk patients from trials, we could minimize early death, improve long-term survival and increase the survival differences between different regimens. We suggest that other groups evaluate our algorithm and exclude poor prognosis patients from trials of dose intensification.

Keywords: small-cell lung cancer; chemotherapy; early death; prognostic factors; sepsis

Early death (ED), defined as death within the first treatment cycle, was more frequently observed in our two latest randomized trials of combination chemotherapy in unselected patients with small-cell lung cancer (SCLC) compared with our previous trials from the Copenhagen Lung Cancer Group. A total of 12.6% of all patients died during the first treatment cycle, 6.7% in limited disease patients and 18.9% in extensive disease. In our preceding trials, early death was encountered in only 4.2% of all the patients, 3.6% in limited disease and 4.8% in extensive disease.

The increased frequency of early death could be caused by either inferior or more toxic treatment or by an increased number of patients with poor prognosis. However, a recent analysis of our treatment outcome during two decades did not encounter stage migration or change in the distribution of prognostic factors over time (Lassen et al, 1998). According to the International Association for the Study on Lung Cancer (IASLC), toxic death, defined as death during septic episodes, should not exceed 5% (Kristjansen et al, 1990). The risk of early death caused by toxicity, i.e. infections, bleeding episodes etc., may be reduced by identification of high-risk patients. They should primarily be found among those with poor prognostic factors, such as extensive stage of disease, poor performance status (PS), high age and high serum lactate dehydrogenase (LDH). Prompted by the observed increased frequency of early death, the present analysis was carried out focusing on treatment intensity and risk factors for early death, especially early toxic death, in patients with SCLC.

PATIENTS AND METHODS

Treatment intensity

The rates of grade IV leucopenia (< 1000 white blood cells mm$^{-3}$) in the recent trials with a higher rate of early death were compared with the trials conducted from 1976 to 1981. The details of these trials have previously been published (Østerlind et al, 1983, 1986, 1991; Hirsch et al, 1987). Early death was defined as death within the first treatment cycle (4 weeks). Early toxic death (ETD) was defined as early death during episodes with neutropenia and fever, and early non-toxic death (ENTD) was defined as early death in all other instances including progression and concurrent diseases. Patients with neutropenia, but without fever or other signs of infection, were allocated to the ENTD group. Data were obtained from a database of patient records and clinical report forms of trials conducted in our group. A detailed comparison of patient characteristics during this time have recently been performed and published (Lassen et al, 1997).

Prognostic factors

A detailed analysis of patients included from 1981 to 1991 in our two latest randomized trials was performed. All trials were conducted jointly at Rigshospitalet and Bispebjerg Hospital, Copenhagen, Herlev University Hospital and Hillerød Sygehus, Denmark, and Renströmska Hospital, Gothenburg, Sweden. The
studies included a total of 941 consecutive patients, and 937 patients were evaluable for this analysis (Pedersen et al, 1987; Lassen et al, 1996).

There were no limitations for inclusion into the studies with respect to PS, the presence of CNS or bone marrow metastases, liver function tests etc., resulting in a considerably higher fraction of patients with poor prognostic factors. Table 1 describes the study design of the two trials.

All data were stored in a database, and the association between treatment, pretreatment factors and early death was analysed with special reference to toxic deaths. The factors included stage, liver, bone marrow and brain metastases, gender, age, PS and biochemical and haematological variables. Pretreatment characteristics of ETD and ENTD patients were compared with the characteristics of those who survived the first 4 weeks (controls) to identify prognostic factors.

**Statistical analysis**

Data were analysed using an interactive program (SPSS, statistical package, 1995). Patient characteristics were compared using $\chi^2$ test for discrete data, and the Kruskal–Wallis test for ordinal scale variables and continuous non-parametric data. Logistic regression analysis was used to analyse factors of prognostic significance for events such as early death, ENTD and ETD. In the analyses, LDH was categorized according to the multiplicity of upper normal levels (450 units l$^{-1}$) and age was dichotomized to <65 or $\geq$65 years. Survival analysis was calculated by the Kaplan–Meier method, and groups were compared for statistically significant difference using the log-rank method.

**RESULTS**

**Treatment comparison**

The frequencies of ED and ETD were less than 3.4% and 1.4%, respectively, in our trials performed in 1976–81 (Österlind et al, 1983, 1986, 1991). In a later randomized trial, these figures changed in a treatment arm with early inclusion of etoposide, based on in vivo cell cycle analysis. In this arm, ED and ETD were encountered in 19% and 10%, respectively, and grade IV leucopenia was observed in 51% of the patients (Hirsch et al, 1987).

Table 2 shows the distribution of grade IV leucopenia, median white blood cell (WBC) nadir, ENTD and ETD in the different treatment regimens of the trials. The combination of epipodophyllotoxins and platinum (EP) in the induction regimens resulted in an increased number of toxic deaths ($P = 0.0002$), whereas treatment had no significant impact on ENTD. The sequential arm in the first trial was more myelotoxic than the other treatment arms (median WBC nadir 760 mm$^{-3}$, $P < 0.001$). In the latest trial, ETD was more frequently observed in the EP regimens even though low median WBC nadir and grade IV leucopenia were more pronounced in the non-EP regimens. The duration of the leucopenic episodes have not been recorded. In general, significantly more ETD occurred among patients receiving EP ($P = 0.04$).

When early death patients were excluded from the two trials, the outcome was significantly changed. In the 1981–85 trial, the survival inferiority of the EP regimens disappeared and no survival difference was present ($P = 0.15$). In the 1985–91 trial, the significant superiority ($P = 0.04$) of the EP regimens increased ($P = 0.002$), as reflected by a difference in median survival of 2 months compared with the previous 1 month. Figure 1 compares the survival curves of all patients, early deaths and the patients surviving the first 4 weeks of treatment.
Prognostic factors

A total of 941 patients were included in the two trials from 1981 to 1991, and 932 patients were evaluable for the analysis of prognostic factors. One hundred and eighteen patients died during the first 4 weeks after start of treatment (12.6%). The most frequent cause of death was disease progression, but whether a patient died from progression or concurrent diseases could not always be defined because some patients died at home and because autopsy was not performed routinely. A majority of ED occurred during the second week after initiation of chemotherapy when blood cell counts were at the lowest level, and ETD was recorded in 38 patients (4%). In some instances, ED may have been caused by a combination of toxicity, progression or concurrent diseases. Table 3 shows the pretreatment characteristics of ENTD patients, ETD patients and the control group. Extensive disease and elevated liver enzymes were significantly more frequent among patients who died within the first 4 weeks, whereas the sex ratio was equal between the groups. Also, poor PS was more frequently observed in early deaths. However, 27% of ETD had initial PS 1 and 27% had initial PS 2. The median age and the median values of lactate dehydrogenase (LDH) and alkaline phosphatase (AP) were significantly higher in patients dying early (Kruskal–Wallis). These factors were equally distributed between the ENTD and ETD group, except extensive stage which was less frequently observed with ETD ($P = 0.05$, $\chi^2$ test).

A variable indicating induction regimens containing epipodophyllotoxins and platinum was introduced (EP). This variable was added to the statistically significant variables from the univariate analysis in subsequent logistic regression analyses of factors with prognostic impact on early death, ETD and ENTD (Table 4). Significant risk factors of early death were high age, poor performance status, elevated LDH and EP treatment. High age, poor performance status and elevated LDH were statistically significant risk factors for ENTD. Extensive stage of disease resulted in a hazard ratio of 1.9, but the confidence interval did not reach significance ($P = 0.07$). EP had no influence on ENTD, and age was not a significant factor in a separate analysis of limited Early death in small-cell lung cancer

517 British Journal of Cancer (1999) 79(3/4), 515–519© Cancer Research Campaign 1999

Table 3 Pretreatment characteristics for early deaths and controls

| Factor                          | ENTD $n = 80$ | ETD $n = 38$ | Controls $n = 819$ | Significance |
|---------------------------------|--------------|--------------|-------------------|-------------|
| Age (median, years)             | 65           | 64           | 61                | $P<0.0005^c$|
| Men/women (%)                   | 63/37        | 66/34        | 65/36             | NS$^b$      |
| Stage L/E (%)                   | 24/76        | 42/58        | 58/42             | $P<0.0005^b$|
| PS (%)                          |              |              |                   |             |
| WHO 0                           | 3            | 3            | 23                | $P<0.0005^c$|
| WHO 1                           | 19           | 27           | 44                |             |
| WHO 2                           | 24           | 27           | 19                |             |
| WHO 3                           | 33           | 24           | 10                |             |
| WHO 4                           | 21           | 19           | 4                 |             |
| AP (median, U $1^{-1}$)         | 444          | 293          | 217               | $P=0.01^c$  |
| LDH (median, U $1^{-1}$)        | 1030         | 860          | 436               | $P=0.0005^c$|
| Liver metastases (%)            | 38           | 40           | 18                | $P<0.005^b$|
| Bone marrow metastases (%)      | 30           | 35           | 14                |             |
| Brain metastases (%)            | 9            | 5            | 3                 |             |

L/E, limited/extent stage; PS, performance status; ENTD, early non-toxic deaths; ETD, early toxic deaths; AP, alkaline phosphatase; LDH, lactate dehydrogenase. $^a$Stage: ENTD vs ETD, $P = 0.05$; $^b$chi-squared test; $^c$Kruskal–Wallis test.

Table 4 Logistic regression analysis of factors of prognostic significance for early death (ED), early non-toxic death (ENTD) and early toxic death (ETD)

| Characteristic | Hazard ratio | 95% CI | Significance |
|----------------|--------------|--------|--------------|
| Early death    |              |        |              |
| Age            | 2.21         | 1.32–3.71 | 0.003        |
| PS             | 2.06         | 1.66–2.56 | <0.0005      |
| Stage          | 1.15         | 0.65–2.03 | 0.6          |
| LDH            | 1.62         | 1.35–1.95 | <0.0005      |
| EP             | 2.32         | 1.42–3.79 | 0.0008       |
| Early non-toxic death |          |        |              |
| Age            | 2.58         | 1.38–4.83 | 0.003        |
| PS             | 1.88         | 1.46–2.42 | <0.00005     |
| Stage          | 1.92         | 0.94–3.92 | 0.07         |
| LDH            | 1.45         | 1.18–1.78 | 0.0004       |
| EP             | 1.15         | 0.66–2.00 | 0.6          |
| Early toxic death |            |        |              |
| Age            | 1.36         | 0.65–2.82 | >0.4         |
| PS             | 1.86         | 1.34–2.57 | 0.0002       |
| Stage          | 0.55         | 0.23–1.32 | 0.2          |
| LDH            | 1.59         | 1.20–2.10 | 0.001        |
| EP             | 6.66         | 2.66–16.70 | 0.0001     |

PS, performance status (WHO); LDH, lactate dehydrogenase; EP, epipodophyllotoxins and platinum during first cycle; CI, confidence interval.

Table 5 Algorithm for predicting risk of early death and poor prognosis

| Characteristic                  | Score |
|--------------------------------|-------|
| Performance status 3 or 4      | 2     |
| LDH > twice upper normal level | 1     |
| Age ≥65 years                  | 1     |

Sum score ≥2 allocates patients to risk group. LDH, lactate dehydrogenase; upper normal level 450 U $1^{-1}$.

Figure 2 Kaplan–Meier survival curve of all 937 patients with SCLC (–), median survival 309 days (95% CI 289–329), compared with the favourable prognosis group, $n = 737$ (---) median survival 351 days (95% CI 331–371), and the poor prognosis group, defined as patients with PS 3 or 4 or LDH > 900 units $1^{-1}$ and age ≥65 years, $n = 200$ (- - -) median survival 133 days (95% CI 86–180).
the control regimen indicates the complexity of this problem. The schedule of administration of epipodophyllotoxins has previously been shown to be important for outcome. Based on in vivo cell kinetic observations, it has been shown that etoposide has maximal activity against cells in the S and G2 phases (Kalwinsky et al., 1983). In most regimens, epipodophyllotoxins, therefore, are administered for 3–5 days early in each cycle. This was also the case in our regimens. This could possibly result in nadir periods of longer duration compared with regimens in which the drugs are administered in 1 day. Unfortunately, our haematological surveillance did not include daily blood cell counts. This was only the case when grade IV toxicity had resulted in hospitalization of the patients. Therefore, a comparison of nadir durations among the different regimens could not be performed.

Risk of early death due to infection seems to be more independent of stage, and fatal toxicity occurred in both stages. Increased LDH was an independent prognostic factor, presumably related to tumour burden rather than to hepatic function. PS was found to be the most powerful risk factor of both toxic and non-toxic early death. Stratification for PS and LDH is extremely important in both disease stages because these patients have an increased risk of early toxic death when treated with extraordinary myelotoxic combination chemotherapy. It is noteworthy that more than 50% of ETD occurred among patients with PS 1 or 2. A majority of these patients had increased LDH and were more than 65 years old. Therefore, it is difficult to propose a simple algorithm predicting risk of ETD. Patients with PS 1 in general have a favourable prognosis and should be included in clinical trials. Our data suggest that good performance patients should receive increased haematological surveillance if they are more than 65 years old and have moderately elevated LDH, they should be excluded from trials with potent myelosuppressive agents.

Most phase II and III trials of high-dose chemotherapy and dose intensification have been disappointing with regard to survival (Stahel et al., 1984; Johnson et al., 1987; Jackson et al., 1988; Klasa et al., 1991; Katakami et al., 1996; Fetscher et al., 1997; Murray et al., 1997). Most of these trials only included patients with good performance status and thereby excluded approximately 15% of the patients. If patients with other poor prognostic factors had been excluded, the results might have been in favour of treatment intensification. Epipodophyllotoxins can safely be administered to elderly patients (Bork et al., 1991, 1997). The risk of toxic death is correlated to dose intensity and scheduling with other agents. According to this and other similar analyses, the goal for treatment of patients with poor prognosis primarily is palliative. It should not be too intensive (Girling et al., 1996a), albeit monotherapy with etoposide may be suboptimal (Girling et al., 1996b; Souhami et al., 1997) compared with intravenous combination chemotherapy. Our analysis did not address how to treat such patients; epipodophyllotoxins and platinum are widely used, and dosage and scheduling will always depend on a narrow balance between efficacy and toxicity.

Our algorithm included predictive factors for both ENTD and ETD. The frequency of early death among the excluded group of patients was as high as 41% when receiving EP. We propose that other groups use and evaluate our algorithm to exclude patients.
with poor prognosis from regimens with potent myelosuppressive agents and trials with dose intensification or dose escalation. Exclusion of such patients may reveal that dose intensification will result in a significantly improved outcome. This should encourage further randomized studies in SCLC in patients with a favourable prognosis. These account for approximately 80% of the patients who better tolerate intensive treatment, and an increased toxicity is acceptable provided more patients become long-term survivors.

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