Pulmonary function after treatment for acute lymphoblastic leukaemia in childhood

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Summary The aim of this study was to examine pulmonary function after acute lymphoblastic leukaemia in childhood and identify risk factors for reduced pulmonary function. We studied a population-based cohort of 94 survivors of acute lymphoblastic leukaemia in childhood who were in first remission after treatment without spinal irradiation or bone marrow transplantation. Pulmonary function test results were compared with reference values for our laboratory, based on 348 healthy subjects who had never smoked from a local population study. A median of 8 years after cessation of therapy (range 1–18 years) the participants had a slight, subclinical, restrictive ventilatory insufficiency and reduced transfer factor and transfer coefficient. The changes in lung function were related to younger age at treatment and to more dose-intensive treatment protocols that specified more use of cranial irradiation and higher cumulative doses of anthracyclines, cytosine arabinoside and intravenous cyclophosphamide than previous protocols. We conclude that, 8 years after treatment without bone marrow transplantation or spinal irradiation, survivors of childhood acute lymphoblastic leukaemia in first remission were without pulmonary symptoms but had signs of slight restrictive pulmonary disease including reduced transfer factor. The increased dose intensity of many recent protocols for childhood acute lymphoblastic leukaemia may lead to increased late pulmonary toxicity.

Keywords: acute lymphoblastic leukaemia; childhood; combination chemotherapy; pulmonary function; late effects of therapy

During the last decades the survival rate after childhood acute lymphoblastic leukaemia (ALL) has been considerably improved (Pui, 1995). Consequently, the frequency and severity of late effects in survivors of childhood ALL have gained importance.

A few reports have been published on the lung function after childhood leukaemia (Miller et al. 1986; Shaw et al. 1989; Jenney et al. 1995; Turner-Gomes et al. 1996). Two groups of 15 and 31 childhood leukaemia survivors had a slight restrictive pulmonary disease 3–7 years after the end of therapy (Miller et al. 1986; Shaw et al. 1989). In contrast, 19 ALL survivors in first remission had normal lung volumes and transfer factor 4 years after the end of therapy (Turner-Gomes et al. 1996). None of these studies identified any risk factors for reduced pulmonary function.

In the largest study to date, 70 childhood leukaemia survivors, examined 4 years after the end of therapy, had lower lung volumes and transfer factor than 146 matched control subjects (Jenney et al. 1995). Chest infections, cyclophosphamide and craniospinal irradiation were risk factors for reduced lung volumes, whereas chest infections, doxorubicin, craniospinal irradiation and bone marrow transplantation were risk factors for reduced transfer factor. Chest infections were defined as the number of lower respiratory tract infections requiring hospitalization during or subsequent to the treatment for leukaemia, irrespective of whether or not an underlying cause was identified (Jenney et al. 1995).

Most of the previous studies had several limitations, such as few participants, many patients lost to follow-up, short follow-up, lack of control groups, or many different diagnoses, stages of disease and treatments included. The two largest studies (Shaw et al. 1989; Jenney et al. 1995) both included patients treated with spinal irradiation and patients treated with total body irradiation and bone marrow transplantation, which are treatments known to cause reduced pulmonary function (Jakacki et al. 1995; Nysom et al. 1996). Thus, the lung function of the majority of childhood ALL survivors – patients in first remission treated without bone marrow transplantation or spinal irradiation – is not well described.

We studied the pulmonary function of 94 survivors of childhood ALL who were in first remission several years after diagnosis and had never been treated with bone marrow transplantation or spinal irradiation.

PATIENTS AND METHODS

Patients

From the population-based Danish Cancer Registry (de Nully Brown et al. 1989) 304 cases of ALL were identified. These patients were at most 14 years old and diagnosed between 1970 and 1990 (inclusive) while residing in east Denmark. On 1 January 1993, 11 patients were still on therapy and 127 had died. Only four patients who had emigrated were lost to follow-up.

Of 162 eligible patients, 128 (79%) gave their informed consent to participate in the present study. Thirty-two patients declined to participate for personal reasons. One was pregnant, and one had a relapse before being studied. Two participants could not perform the pulmonary function tests because of young age. To study the pulmonary function after standard ALL therapy, we excluded 24 patients treated for a relapse, one treated for a second malignancy and seven other patients treated with bone marrow transplantation.
mediastinal irradiation or Carmustine. This leaves 94 participants off therapy in first remission of ALL for the present analysis.

The median age of the participants was 3.9 years at diagnosis (range 0.5–14.8 years), 6.9 years at completion of therapy (3.5–19.7 years) and 16.2 years at time of the study (5.3–34.2 years). The median length of follow-up was 10.6 years from diagnosis (3.4–23.4 years) and 7.6 years from completion of therapy (1.2–18.3 years).

The participants had received chemotherapy and cranial irradiation as described in Table 1. The dose of cranial irradiation was 15–18 Gy (n = 23) or 24 Gy (n = 16). No participants received other types of irradiation.

The height of the participants at time of the study ranged from 119 to 179 cm in female subjects and from 110 to 185 cm in male subjects. The average height did not differ significantly from national reference data (mean standardized residual −0.1, 95% confidence interval −0.3 to 0.1, range −2.1 to 3.0) (Andersen et al. 1982).

Participants who had received cranial irradiation were significantly shorter than the other participants (estimated difference between means 0.8 standardized residual, 95% confidence interval 0.4–1.2), but their sitting height to standing height ratio did not differ significantly from local reference data (Holm. 1996).

At the time of the study 18 participants smoked, 4 had previously smoked, whereas 72 had never smoked. The four who had previously smoked were all considered as smoking subjects as they had consumed the same median amount of cigarettes as the 18 current smoking subjects and as they had stopped smoking only 2–17 months before being studied. This gave a group of 22 smoking subjects in all (seven men, P = 0.05 for more women who smoked), who had consumed a median of 1.5 pack-years of cigarettes (1 pack-year is 20 cigarettes a day for 1 year; range 0.1–10.5) over a median period of 3 years (1–13 years).

### Pulmonary function testing

All pulmonary function tests were performed in the same laboratory in accordance with the European recommendations (Quanjer and Tammelng, 1983). The forced expiratory volume in 1 second (FEV₁) and the forced vital capacity (FVC) were measured with a pneumotachograph (Jaeger, Germany). In patients who terminated the forced expiration in less than 1 s, the forced expiratory volume in 0.5 s was recorded instead. The ratio of FEV₁/FVC was calculated as a percentage. Flow–volume curves were evaluated by one of us (BH). Total lung capacity (TLC) was measured by the helium dilution technique (Jaeger, Germany) and the transfer factor for carbon monoxide with the single-breath technique according to the recommendations given by the American Thoracic Society (Anonymous, 1987), except that TLC and transfer factor were only determined once if cooperation was good. The equipment detected carbon monoxide with an infrared spectrophotometer (Jaeger, Germany) and helium with a thermal conductivity method. The transfer coefficient was calculated as transfer factor divided by alveolar volume.

Pulmonary function test results were compared with reference values for our laboratory, which were generated by adjusting published reference values (Quanjer and Tammeling, 1983; Table 1: Treatment protocols)

| Protocol | n | Induction | Consolidation | CNS prophylaxis | VCR/ PRED | Anthra- cyclines | Cyclo- phosphamide | Unusual cases |
|----------|---|-----------|---------------|----------------|-----------|----------------|-------------------|---------------|
| 70–76    | 14| VCR/PRED  |               |                | +          |                |                   |               |
|          |    | –         |               |                |           |                |                   |               |
| 77–81    | 19| VCR/PRED  |               |                | +          |                |                   |               |
|          |    | –         |               |                |           |                |                   |               |
| 81 SR    | 8 | VCR/PRED  |               |                | +          |                |                   |               |
|          |    | –         |               |                |           |                |                   |               |
| 81 IR    | 11| VCR/PRED/DOX |           |                | +          |                |                   |               |
|          |    | –         |               |                |           |                |                   |               |
| 81 HR    | 10| VCR/PRED/DOX/L-ASP |       |                | +          |                |                   |               |
| 86 SR    | 6 | VCR/PRED  |               |                | +          |                |                   |               |
|          |    | –         |               |                |           |                |                   |               |
| 86 IR    | 17| VCR/PRED/DNR |             |                | +          |                |                   |               |
|          |    | –         |               |                |           |                |                   |               |
| 86 HR    | 7 | PRED→DEX/CYC/Ara-C→VM-26/IDM→VCR/DN/L-ASP | |                | +          |                |                   |               |
| Other    | 1 | VCR/PRED/DOX |           |                | +          |                |                   |               |
|          |    | –         |               |                |           |                |                   |               |

All participants received intrathecal methotrexate and maintenance therapy with oral 6-mercaptopurine and methotrexate. Participants diagnosed after May 1981 were treated according to protocols of the Nordic Society of Paediatric Haematology and Oncology (NPHO) (Schroeder et al., 1995) Doses of anthracyclines and cyclophosphamide are given as cumulative mg m⁻² of body surface area; 6MP: 6-mercaptopurine; 6TG: 6-thioguanine; Ara-C: cytosine arabinoside; CNS: central nervous system; CYC: intravenous cyclophosphamide; DEX: dexamethasone; DNR: daunorubicin; DOX: doxorubicin; HDM: high-dose methotrexate (1 g m⁻²); HR: high risk; IDM: intermediate-dose methotrexate (0.5 g m⁻²); IFOS: ifosfamide; IR: intermediate risk; L-ASP: L-asparaginase; MTX: methotrexate; PRED, prednisone; RT: radiotherapy of the CNS; SR: standard risk; VCR, vincristine; VM-26, teniposide; VP-16, etoposide; *VCR and PRED reinductions; *three patients also received RT, *one patient received 23 mg m⁻², one 120 mg m⁻² and one 281 mg m⁻² of DNR. **two patients also received RT, **two patients received 75 mg m⁻² of DNR for suspected, but never confirmed, relapse; *MTX 4 g m⁻² and MTX/methylPRED/Ara-C intrathecal.
Rosenthal et al. 1993; Quanjer et al. 1995; Stam et al. 1996) to fit 348 healthy 13- to 24-year-old Caucasian subjects who had never smoked from a local population study (Nysom et al. 1997). The height of the control subjects ranged from 143 to 182 cm in women and from 147 to 200 cm in men. In addition to evaluating each individual pulmonary function test variable, an integrated evaluation of the pulmonary function was also performed, based on the combination of the different variables. The evaluation was carried out without knowledge of the leukaemia treatment protocol of the patient. Pulmonary function was classified as (a) restrictive pattern (reduced FVC or TLC; with or without reduced transfer factor or coefficient; normal or elevated FEV/FVC); (b) restrictive flow--volume curve pattern but with volumes and capacities within reference limits; (c) reduced transfer factor or transfer coefficient with normal lung volumes and flow--volume curves; (d) obstructive pattern (low FEV/FVC); or (e) normal.

Data analysis
To make data comparable, pulmonary function test results and heights were analysed as standardized residuals: (observed value – predicted value) divided by the residual standard deviation (Quanjer and Tammelng, 1983). Standardized residuals are equivalent to standard deviation (Z) scores. The distribution of pulmonary function test results and heights did not differ significantly from a normal distribution (Shapiro–WilK test), so these results are given as mean values. 95% confidence intervals of mean and ranges. All other data are reported as median values with ranges. Student’s t-test was used for determining whether height and pulmonary function test results differed significantly from the reference values (i.e. a standardized residual of 0), and for comparing the results of groups of patients. Pulmonary function test results were considered abnormal if they were more than 1.645 residual standard deviation from the predicted mean value (Quanjer and Tammelng, 1983). If raised as well as reduced values of a variable are considered abnormal (TLC, FEV/FVC, transfer coefficient) this corresponds to two-sided 90% prediction limits for reference data. If only reduced values are considered abnormal (FVC, FEV , transfer factor) this corresponds to one-sided 95% limits. Parametric (Pearson) and non-parametric (Spearman) correlation analysis and simple and multiple linear regression models were used according to the distribution of data to evaluate possible predictive variables of pulmonary function. In the multiple regression analysis, variables that did not significantly improve the models were removed, one at a time, in a step-down procedure. The continuity adjusted chi-square test and Mann–Whitney’s unpaired test were used for comparing baseline characteristics between groups of patients (Altman, 1991). Probabilities below 0.05 were considered statistically significant and data were analysed using the SAS computer software package (SAS Institute, Cary, NC, USA).

Ethics
All participants and the parents of the children younger than 18 gave their written informed consent. The study was in accordance with the Helsinki II declaration and was approved by the local medical ethics committee of Copenhagen, Denmark (approval no. KF V92-097).

RESULTS
Participants
At the time of the study one participant received captopril for arterial hypertension, and one used beclomethasone inhalations for asthma. Ten participants had pulmonary symptoms: one complained of dyspnoea, two complained of cough at night, five complained of cough on exertion, one complained of cough at night and on exertion, whereas one complained of dyspnoea and cough at night. Eighty-four participants had no pulmonary symptoms. Fourteen participants (including five smoking subjects) considered their physical work capacity better than that of other people of their own age. 63 (14 smoking subjects) considered it equal to that of other people of their own age. 14 (three smoking subjects) considered it a bit inferior, and three (non-smoking subjects) much inferior. Five participants had haemoglobin concentrations 0.2–1.1 g dl⁻¹ below the lower limits of normal of our laboratory. All others had values evenly distributed within the normal range of our laboratory (5–7 years: 11.9–14.8, 7–13 years: 12.1–15.6, female subjects >13 years: 11.3–16.1, male subjects >13 years: 12.9–17.7 g dl⁻¹).

| Variable | Number evaluable | Mean (95% CI) | Range | Number with reduced/ raised values* |
|----------|------------------|--------------|-------|------------------------------------|
| Forced vital capacity | 94 | −0.4 (−0.6 to −0.2) | −2.9–2.3 | 14/3 |
| Forced expiratory volume in 1 s | 83 | −0.2 (−0.5 to −0.01) | −2.6–3.1 | 7/4 |
| Ratio of FEV₁ to FVC | 83 | 0.1 (−0.1 to 0.3) | −2.4–2.0 | 1/5 |
| Total lung capacity | 89 | −0.2 (−0.5 to −0.03) | −2.1–2.3 | 10/5 |
| Transfer factor | 67 | −0.4 (−0.7 to −0.2) | −3.4–1.7 | 9/1 |
| Non-smokers | 22 | −0.7 (−1.0 to −0.4) | −2.0–0.8 | 3/0 |
| Smokers | 66 | −0.3 (−0.5 to −0.1) | −2.2–2.0 | 4/2 |
| Transfer coefficient | 22 | −0.9 (−1.4 to −0.4) | −2.9–1.5 | 6/0 |

Results are given as standardized residuals; CI confidence interval; FEV₁ forced expiratory volume in 1 s; FVC, forced vital capacity; *standardized residuals <=−1.645 / >=1.645.

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Lung volumes

Pulmonary function test results (Table 2) were poorly related to self-estimated physical work capacity (plots not shown), and only four of the ten participants with pulmonary symptoms had abnormal pulmonary function tests. Similarly, standardized residuals for lung function and height were not related (plots not shown). For five participants measurements of TLC and transfer factor were impossible because of poor co-operation at the single-breath procedure.

The mean FVC, FEV, and TLC standardized residuals were all slightly, but significantly reduced, and 7–14 participants (one smoking subject) had values below the fifth percentile (Table 2). In 11 participants, the forced expiration was terminated in less than 1 s, but in ten of these the forced expiratory volume in 0.5 s was larger than the predicted FEV\textsubscript{1}.

FEV\textsubscript{1}/FVC ratio and flow–volume curves

The mean FEV\textsubscript{1}/FVC standardized residual was normal (Table 2). Two flow–volume curves were classified as obstructive, and one was also restrictive. None of these curves were from smoking subjects or patients with short expirations. Eighteen other flow–volume curves appeared restrictive, including the curves of seven patients (one smoking subject) with a low TLC.

Transfer factor and transfer coefficient

The mean transfer factor and transfer coefficient standardized residuals were significantly reduced both in non-smoking subjects and in smoking subjects (Table 2).

Table 3  Regression models for total lung capacity

| Variable                                           | R\textsuperscript{2} | Regression coefficient (95% CI) | P   |
|----------------------------------------------------|-----------------------|---------------------------------|-----|
| Simple regression models                           |                       |                                 |     |
| Doxorubicin (mg m\textsuperscript{-2})              | 0.03                  | −0.0022 (−0.0051 to 0.0007)     | 0.13|
| Daunorubicin (mg m\textsuperscript{-2})             | 0.01                  | −0.0018 (−0.0035 to 0.0019)     | 0.3 |
| Cyclophosphamide intravenously (g m\textsuperscript{-2}) | 0.04                  | −0.11 (−0.23 to 0.001)          | 0.07|
| Cytosine arabinoside (g m\textsuperscript{-2})      | 0.05                  | −0.26 (−0.50 to −0.02)          | 0.04|
| 6-Thioguanine (g m\textsuperscript{-2})             | 0.03                  | −0.39 (−0.89 to 0.11)           | 0.12|
| Anthracyclines (mg m\textsuperscript{-2})          | 0.03                  | −0.0017 (−0.0038 to 0.0003)     | 0.10|
| High-dose methotrexate (number of cycles)          | 0.00                  | −0.0065 (−0.0082 to 0.00740)    | 0.9 |
| Cranial irradiation                                | 0.05                  | −0.45 (−0.88 to −0.02)          | 0.04|
| Intermediate- or high-risk protocol\textsuperscript{a} | 0.03                  | −0.36 (−0.79 to 0.06)           | 0.09|
| Smoking                                            | 0.02                  | +0.31 (−0.18 to 0.80)           | 0.2 |
| Female sex                                         | 0.00                  | +0.024 (−0.40 to 0.45)          | 0.9 |
| Follow-up after diagnosis (years)                  | 0.04                  | +0.036 (−0.004 to 0.076)        | 0.08|
| Age at diagnosis (years)                           | 0.05                  | +0.055 (0.001 to 0.109)         | 0.045|
| Age at follow-up (years)                           | 0.08                  | +0.044 (0.012 to 0.076)         | 0.01|

| Multiple regression models\textsuperscript{a}      |                       |                                 |     |
| Doxorubicin                                        | 0.09                  | −0.0063 (−0.0107 to −0.0020)    | 0.005|
| Doxorubicin × age at diagnosis\textsuperscript{a}  |                       | +0.00065 (0.00013 to 0.00116)  | 0.01|
| Cyclophosphamide intravenously                     | 0.10                  | −0.14 (−0.25 to −0.02)          | 0.02|
| Age at diagnosis                                   | 0.11                  | −0.068 (0.014 to 0.122)         | 0.01|
| Cytosine arabinoside                               | 0.11                  | −0.30 (−0.54 to −0.07)          | 0.01|
| Age at diagnosis                                   | 0.11                  | −0.066 (0.012 to 0.119)         | 0.02|
| Cranial irradiation                                | 0.11                  | −0.53 (−0.95 to −0.11)          | 0.01|
| Age at diagnosis                                   | 0.11                  | +0.066 (0.013 to 0.119)         | 0.02|
| Intermediate- or high-risk protocol\textsuperscript{a} | 0.09                  | −0.89 (−1.50 to −0.28)          | 0.005|
| Intermediate- or high-risk protocol × age at diagnosis\textsuperscript{a} | 0.09                  | +0.085 (0.012 to 0.157)         | 0.02|

CI, confidence interval; \( R^2 \) of a model indicates how large a proportion of the total variation the model explains; \( * \) cumulative dose of doxorubicin + daunorubicin; \( \dagger \) the intermediate and high-risk protocols from 81 and 86 (Table 1); \( \ddagger \) daunorubicin, anthracyclines, and 6-thioguanine were non-significant in the multiple regression models, and consequently were removed during the backward selection procedure; \( \ast \) interaction term (see text).

The transfer factor values given were not corrected for haemoglobin concentration in the present study because the reference values for pulmonary function for our laboratory were based on transfer factor measurements without haemoglobin correction and because the haemoglobin concentration of nearly all participants was within normal limits. Correcting the transfer factor to the average age-specific haemoglobin concentration of our laboratory according to the equation of Cotes (Quanjer and Tammeling, 1983) had no influence on conclusions concerning transfer factor (data not shown).

Integrated evaluation of pulmonary function

When all available measures of pulmonary function were considered together, 25 participants (one smoking subject) had a restrictive pattern, six of them with normal lung volumes and capacities. Ten (six smoking subjects) had a reduced transfer factor or transfer coefficient with normal lung volumes and flow–volume curves, and two (one smoking subject) had an obstructive pattern. Sixty-one per cent of the participants were estimated to have a normal pulmonary function.

Risk factors for reduced pulmonary function

The age at diagnosis and the length of follow-up after diagnosis were not significantly different for patients with a reduced TLC and patients without a reduced TLC, but nine out of ten patients with a reduced TLC had been treated according to the intermediate or high-risk protocols of the 1980s (Table 1). These protocols used more cytosine arabinoside, 6-thioguanine, anthracyclines and intravenous...
cyclophosphamide than the other protocols. The mean TLC was 
-0.4 (-0.8 to -0.1) in 40 patients treated according to intermediate or 
high-risk protocols, compared with -0.1 (-0.3 to 0.2) in the 49 other 
patients (estimated difference between means 0.4 standardized 
residual, 95% confidence interval -0.1 to 0.8).

Using simple and multiple linear regression analysis we 
analysed the relationship between TLC standardized residual and 
cumulative doses of the five most characteristic drugs of the inter-
mediate and high-risk protocols (doxorubicin, daunorubicin, intra-
venous cyclophosphamide, cytotoxic arabinoside, 6-thioguanine), 
the dose of anthracyclines (doxorubicin plus daunorubicin doses), 
the number of high-dose methotrexate cycles, cranial irradiation, 
treatment according to intermediate or high-risk protocols, 
smoking, sex, the length of follow-up after diagnosis and the ages 
at diagnosis and follow-up (Table 3). In the simple regression 
analysis there was a significant relationship between a lower TLC 
and a larger cumulative dose of cytotoxic arabinoside, cranial ir-
radiation, younger age at diagnosis and younger age at follow-up.

The cumulative doses of the five drugs considered and the use 
of cranial irradiation were all strongly correlated with each other 
(P < 0.0001) except for doxorubicin with daunorubicin when 
P = 0.008). Therefore, testing several of these variables in one 
multiple regression model would be meaningless. As age at diag-
nosis was not significantly correlated with any of these variables, 
we subsequently tested multiple regression models considering 
age at diagnosis, one of the five drugs mentioned or cranial irradi-
ation and a term representing interaction between the treatment 
variable and age at diagnosis. A statistically significant interaction 
term indicates that the pulmonary toxicity of a certain treatment 
depends on the age at which this treatment is given. Variables that 
did not improve the models significantly were removed, one at a 
time, and the resulting models are shown in Table 3. Age at follow-
up was significantly correlated with TLC and also with the treat-
ment related variables considered (non-parametric correlation 
coefficient -0.37 to -0.59, P ≤ 0.0002). Neither age at follow-up, 
length of follow-up, nor smoking improved any of the multiple 
regression models significantly.

When we applied the multiple regression models in Table 3 to 
the 52 participants who were within the height and age range of 
the local control group, the estimated regression coefficients were all 
within the confidence intervals given in Table 3. However, the esti-
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were also risk factors for reduced lung function in the study by Jenney et al (1995). As we excluded patients treated with craniospinal irradiation or bone marrow transplantation and as we did not test chest infections as an explanatory variable, we could not evaluate the influence of these factors.

Cranial irradiation was apparently a risk factor for reduced TLC in our multiple regression analysis. This may be because persons treated with cranial irradiation had abnormal body proportions, which resulted in an inadequately high predicted lung function. However, the body proportions of the participants were normal, and reduced lung function was not related to height. It therefore seems more likely that cranial irradiation was significant in our regression models simply because it reflects the more dose-intensive treatment protocols.

Age at follow-up was the strongest predictor of TLC in the simple regression models (Table 3). This could be explained by a gradual recovery of pulmonary function over decades after the end of chemotherapy. However, there was also a significant correlation between age at follow-up and the doses of the drugs in the multiple regression models. It therefore seems more likely that age at follow-up was significant in a simple regression model because it reflects the change in treatment protocols.

Most participants with reduced transfer coefficient smoked. This, and the fact that the best regression model of transfer factor included only TLC and tobacco smoking, indicates that the reduced transfer factor after chemotherapy seems to be caused mainly by reduced lung volumes. The present data therefore do not suggest thickening of the pulmonary diffusion membrane or reduced pulmonary capillary blood volume.

Mechotrexate treatment was related to lung toxicity in some early studies (Clarysse et al, 1969; Nesbit et al, 1976). In contrast, the more recent studies found no lung toxicity of mechotrexate (Miller et al, 1986; Shaw et al, 1989; Jenney et al, 1995). The present data support the latter finding because lung function was not associated with the number of high-dose mechotrexate cycles and because the TLC was normal in the 49 patients treated according to other than intermediate or high-risk protocols, although these patients received a higher cumulative dose of oral mechotrexate than patients treated according to intermediate or high-risk protocols (data not shown). Cytosine arabinoside has been reported to cause acute pulmonary oedema (Shearer et al, 1994), but usually after higher doses than those received by our patients. Cytosine arabinoside has not previously been linked to abnormal pulmonary function tests.

Anthracyclines could affect the lung function by inducing congestive heart failure. The cumulative doses of anthracyclines received by our participants, however, were lower than what has yet been reported to cause late clinical cardiotoxicity (Lipshultz et al, 1991; Steinherz et al, 1991), and only a few patients reported any cardiopulmonary symptoms, which makes this explanation unlikely.

In a large population study the transfer factor was reduced shortly after the onset of smoking, whereas lung volumes were unchanged during the first decade of smoking (Knudson et al, 1989). Our data were in accordance with this and did not suggest increased pulmonary toxicity of tobacco smoking in survivors of childhood ALL compared with the background population. However, because of the small sample size and the relatively mild tobacco exposure of participants, our study does not have the statistical power to exclude a clinically relevant increased toxicity of tobacco smoking in those treated with chemotherapy in childhood. Tobacco smoking is undoubtedly toxic to the lung, childhood cancer survivors are at increased risk of developing second malignant neoplasms, and many of them have had subclinical damage to several organs causing a reduced functional reserve. Consequently, long-term survivors of childhood cancer should be eagerly warned against tobacco smoking, irrespective of whether the tobacco toxicity to their lungs is actually greater than in the background population or not.

As in a previous study (Jenney et al, 1995), pulmonary function correlated poorly with self-estimated physical work capacity. Thus, self-estimated physical work capacity can apparently not be used for screening patients for reduced pulmonary function. The prevalence of asthmatic symptoms was lower in our participants than reported in a local population survey study (Backer, 1995).

Selection bias should not influence our results because patients and controls were selected from population-based cohorts and a high percentage of eligible subjects participated. It is a limitation of our study that the age and height range of participants was wider than that of control subjects. However, our local predictive equations for lung function were generated by adjusting the best available predictive equations from the literature to fit the large group of local control subjects optimally. The reference equations from the literature that we adjusted are based on subjects covering the full height and age range of our participants. Furthermore, the estimated regression coefficients of our multiple regression models were only minimally affected by limiting the analysis to patients within the height and age range of the control subjects.

The present study was cross-sectional in design and treatment was non-randomly assigned. Consequently, year of diagnosis, length of follow-up, age at follow-up, risk group and type of therapy were all closely related and could therefore not be tested against each other. A cohort effect must be considered as a source of error in a cross-sectional study, but the expected cohort effect would tend to cause the opposite results of what was found. If the drugs given had similar pulmonary toxicity, patients treated during the last part of the study period would be expected to have suffered less complications during therapy than patients treated during the first part because of improved supportive care and increasing clinical experience. In contrast with this, patients treated during the last part of the study period actually experienced more pulmonary toxicity.

Studies of late effects are historic in nature: when the long-term toxicity of a treatment protocol is assessable the protocol itself has often been revised several times already, thus necessitating a study of the late effects of the altered protocol. The recent tendency to increase the dose intensity of treatment protocols for a number of childhood cancers also calls for new studies of late effects. Using a repeated follow-up study of the present cohort, it would be possible to determine whether pulmonary function recovers, stays unchanged or deteriorates late after childhood ALL.

In conclusion, several years after therapy for childhood ALL, participants had a subclinical, slight restrictive pulmonary disease including reduced transfer factor. The reduced pulmonary function was related to young age at treatment and to more dose-intensive protocols. The abnormalities appear mild, but the limited number of patients studied to date, the cross-sectional design of the few available studies and the increased dose intensity of many recent ALL protocols make long-term clinical follow-up of survivors of childhood ALL necessary.

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