Improved outcome in children with advanced stage B-cell non-Hodgkin’s lymphoma (B-NHL): results of the United Kingdom Children Cancer Study Group (UKCCSG) 9002 protocol

A Atra¹, JD Imeson², R Hobson², M Gerrard³, IM Hann⁴, OB Eden⁵, RL Carter¹, CR Pinkerton¹ on behalf of the UKCCSG

¹Department of Paediatric Oncology, The Royal Marsden Hospital NHS Trust, Downs Road, Sutton, Surrey SM2 5PT, UK; ²UKCCSG, University of Leicester, Department of Epidemiology and Public Health, 22–28 Princess Road West, Leicester LE1 6TP, UK; ³Sheffield Children’s Hospital NHS Trust, Western Bank, Sheffield S10 2TH, UK; ⁴The Hospital For Sick Children, Great Ormond Street, London WC1 SJH, UK; ⁵Academic Unit of Paediatric Oncology, Christie Hospital NHS Trust, Manchester M20 9BX, UK

Summary
From July 1990 to March 1996, 112 children with stage III or IV B-cell non-Hodgkin’s lymphoma (B-NHL) with up to 70% FAB L3-type blasts (n = 42) in the bone marrow without central nervous system (CNS) disease were treated on the United Kingdom Children Cancer Study Group (UKCCSG) 9002 protocol (identical to the French LMB 84). The median age was 8.3 years. There were 81 boys and 31 girls. According to the extent of the primary disease, patients were sub-staged into three groups: IIIA with unresectable abdominal tumour (n = 39); IIIB with abdominal multiorgan involvement (n = 57) and IIIX with extra-abdominal primary lymphoma often presenting as pleural effusion (n = 16). Univariate and multivariate analyses were carried out to evaluate the prognostic significance of lactate dehydrogenase (LDH) level at diagnosis, the sub-stage and the time to achieve complete remission (CR). With a median follow up of 48 months (range 12–92), the overall and event free survival (EFS) is 87% (95% confidence interval (CI) 79.2–92.1%) and 83.7% (95% CI 76.3–89.2%) respectively. Six patients (5.4%) never achieved CR, of whom one is alive following high-dose therapy. Eight patients (7.1%) relapsed after achieving CR, three are alive after second-line therapy. There were three early toxic deaths (2.7%), mainly from infection, and one late death from a second cancer. There was no significant difference in EFS according to LDH level at diagnosis, the sub-stage or the time to CR. This study confirms the overall good prognosis and low rate of toxic deaths in patients with advanced B-NHL treated with this intensive regimen. No significant difference in EFS according to the sub-stage, the time to achieve CR or LDH level at diagnosis making it difficult to identify a group that should not receive intensive therapy. © 2000 Cancer Research Campaign

Keywords: childhood; B-cell non-Hodgkin’s lymphoma; advanced stage III and IV; UKCCSG 9002 protocol

Over the last two decades, various studies have shown consistent improvement in the overall prognosis of children with B-cell non-Hodgkin’s lymphoma (B-NHL) (Philip et al, 1982; Al-Attar et al, 1986; Patte et al, 1990). This is attributable to both a reduction in early toxic deaths (Lynch et al, 1977; Allegretta et al, 1985; Miron et al, 1997) and a decrease in relapse rate following an increase in the intensity of systemic and central nervous system (CNS)-directed therapy (Hann et al, 1990; Patte et al, 1991; Cairo et al, 1996). This improvement in the results of treatment has involved patients with all stages of disease (Pinkerton et al, 1991; Patte et al, 1997; Atra et al, 1998). Radiotherapy no longer appears to have a role in the treatment of primary CNS disease (Patte et al, 1986; Bowman et al, 1996), and radical surgery with its potential risks, has become unnecessary (Al-Attar et al, 1989; Miron et al, 1997). Relapse still occurs and is the principle cause of treatment failure. Patients who relapse after first-line chemotherapy can occasionally be salvaged with more intensive chemotherapy with or without megatherapy and stem cell rescue (Philip et al, 1993; Sweetenham et al, 1994).

The United Kingdom Children Cancer Study Group (UKCCSG) 9002 protocol was based on the French LMB-84 and it involved 3 months of treatment with sequential doses of chemotherapy including intrathecal (i.t.) therapy. No radiotherapy was given. We report the results of this regimen in the treatment of children with stage III and IV B-NHL (L3 blasts up to 70% in the bone marrow). The prognostic significance of serum lactate dehydrogenase (LDH) level at diagnosis, tumour substage and time to achieve complete remission (CR) were prospectively analysed in an attempt to identify different risk groups and to facilitate delivery of risk-directed therapy in the future.

PATIENTS AND METHODS
Children under 18 years with newly diagnosed Murphy stage III or IV B-NHL and up to 70% FAB L3-type blasts in the bone marrow (but with no CNS disease), with B-cell histology (Burkitt’s or Burkitt’s-like or large cell) and/or phenotype (surface membrane immunoglobulin-positive) were eligible for this protocol. This is the group of patients identified by the LMB group to be of...
The schema and details of the UKCCSG 9002 protocol are shown in Figure 1 and Table 1 respectively. The first course of treatment consisted of low-dose cyclophosphamide, vincristine and prednisolone (COP) in an attempt to avoid severe drug toxicity, often associated with tumour lysis syndrome at diagnosis. This course could be repeated 1 week later if the clinical condition of the patient remained poor before proceeding with the more intensive treatment of cyclophosphamide, vincristine, prednisolone, doxorubicin and high-dose methotrexate (COPADM). All patients received regular intrathecal (i.t.) hydrocortisone with either methotrexate or cytarabine. Patients were assessed clinically and radiologically to confirm response and exclude progressive disease after COP. Full assessment was done to confirm CR prior to the second block of cytarabine and high-dose methotrexate (CYM2) (Figure 1). Those who achieved CR continued on protocol. In the absence of CR, patients were treated with more intensive chemotherapy as in the UKCCSG 9003 protocol (Atra

| Regimen                  | Dose             | Administration          |
|--------------------------|------------------|-------------------------|
| COP                      | Cyclophosphamide | 300 mg m^2 i.v. bolus day 1 |
|                          | Vincristine      | 1 mg m^2 i.v. bolus day 1 |
|                          | Prednisolone     | 60 mg m^2 Oral days 1–7 |
|                          | i.t. MTX + HC    | day 1                   |
| COPADM 1                 | Vincristine      | 2 mg m^2 i.v. bolus day 1 |
|                          | Doxorubicin      | 60 mg m^2 i.v. over 6 h day 2 |
|                          | Cyclophosphamide | 250 mg m^2 dose i.v. bolus 12 h days 2–4 (with mesna) |
|                          | High-dose MTX    | 3 m^2 i.v. over 3 h day 1 |
|                          | Folinic acid rescue | 15 mg m^2 i.v. from day 2 |
|                          | Prednisolone     | 60 mg m^2 oral days 1–5 |
|                          | i.t. MTX+HC      | day 2, 6                |
| COPADM 2 as COPADM 1 except COPADM 2 as COPADM 1 except | Vincristine | 2 mg m^2 i.v. bolus day 1, 6 |
|                          | Cyclophosphamide and mesna | 500 mg m^2 dose i.v. bolus 12 hr days 2–4 |
| CYM 1+2                  | Cytarabine       | 100 mg m^2 day . continuous infusion days 2–6 |
|                          | High-dose MTX    | 3 m^2 . over 3 h day 1   |
|                          | Folinic acid rescue | 15 mg m^2 i.v. from day 2 |
|                          | i.t. MTX + HC    | day 2                   |
|                          | i.t. Cytarabine + HC | day 7                |
| COPADM 3 AS COPADM 1 (expect that cyclophosphamide is given for 2 days only) | COPADM 3 AS COPADM 1 (expect that cyclophosphamide is given for 2 days only) |

**Table 1 UKCCSG 9002 protocol**

*standard risk*. Those with up to 70% L3-type blasts in the bone marrow but no blasts in peripheral blood (who would be defined by the Murphy scheme as B-ALL) are included. Informed consent was obtained from all patients and their parents as appropriate. Staging investigations included bilateral bone marrow aspirations, cerebrospinal fluid (CSF) cell count and cytospins, a chest radiograph and ultrasonography, with or without computerized tomography (CT) scan of the primary tumour. All pathology specimens, biopsies, pleural or ascitic fluid, bone marrows and CSF were reviewed centrally. Surgical attempts were limited to obtain adequate biopsy material for histopathological diagnosis.

One hundred and thirteen consecutively diagnosed and previously untreated patients were entered between July 1990 and March 1996 (one patient was identified after entering the study to be HIV-positive and was subsequently excluded from the analysis). Eighty-one patients were boys and 31 girls. Ages at diagnosis ranged from 2.2 to 17.5 years (median 8.3 years). For the purpose of this study, patients were substaged according to the French Society of Paediatric Oncology (SFOP) classification, into two groups according to the extent of primary disease (Philip et al., 1987). Thirty-nine patients were stage IIIA as defined by the presence of limited but unresectable disease usually localized to the abdomen with or without ascites. Patients with abdominal primary B-NHL, particularly those with pleural involvement often presenting with effusion, are believed to have a worse outcome (Sandlund et al., 1990); in this study those are classified as IIIX (n = 16). Among the 112 patients, 42 had involvement of the bone marrow at diagnosis with up to 70% L3-blasts. No patient included in this study had CNS disease at diagnosis.

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et al, 1998), followed by high-dose chemo/radiotherapy and stem cell rescue. Patients were formally assessed at the end of treatment to confirm remission with 1–2 monthly review for the first year after the end of treatment. The follow-up was by clinical examination and radiological imaging was only performed if there was clinical concern.

All patients received appropriate supportive care immediately after diagnosis. Antimicrobials, blood and platelet transfusions and total parental nutrition (TPN) were used as indicated. During induction therapy hyper-hydration was given using dextrose–saline 3 l m–2 per day and allopurinol 10 mg kg–1 day–1. Dialysis (peritoneal or haemodialysis) was performed according to the clinical situation. No patient received prophylactic granulocyte colony-stimulating factor (G-CSF).

Statistical methods

The main end point used to evaluate the protocol treatment was event-free survival (EFS). This was defined as the length of time from diagnosis to the earliest detection of relapse/progression or death from any cause. Surviving relapse-free patients were censored at the date of the last clinical follow-up and survival curves were calculated by standard methods (Kaplan and Meier, 1958; Rothman, 1978). Univariate analysis was carried out using the log-rank test (Peto et al, 1977) to evaluate the differences in prognosis between patients in relation to the sub-stage – IIIA vs IIIB vs IIIX; LDH level at diagnosis twice or more vs less than twice the institutional upper limit normal (ULN) and the achievement of CR within less than 28 versus 28 days or more. Multivariate analysis was carried out including all these factors together to test for independent effects (SAS, 1991).

RESULTS

Six patients (5.4%) never achieved CR. All died 1–6 months from diagnosis (median 3.6 months) except one who continued to have residual lymphoma (biopsy proven) at the primary site. He went into CR after high-dose chemotherapy/radiotherapy and autologous bone marrow rescue and is alive 2 years from diagnosis.

At the time of analysis, eight patients (7.1%) have relapsed after achieving CR. Sites of relapse were combined primary site and bone marrow (n = 2), bone marrow (n = 2), primary site (n = 3) and one patient relapsed in the bone marrow, primary site and CNS. Five of these eight patients died 6–14 months from diagnosis (median 9.2 months) despite receiving subsequent more intensive chemotherapy (n = 4) or high-dose chemo/radiotherapy with autologous stem cell rescue (n = 1). Three are alive in CR after receiving further intensive chemotherapy using the UKCCSG 9003 protocol or CHOP, followed by autologous stem rescue (n = 2) or matched unrelated bone marrow (n = 1), 2–6 years from diagnosis (median 3.6 years).

Toxicity

Three patients (2.7%) died of early toxicity. Causes of death were disseminated aspergillosis at 35 days (n = 1), sepsis with post-mortem evidence of anthracycline-induced cardiomyopathy at 30 days (n = 1) and peritonitis with renal cortical necrosis and...
congestive cardiac failure 52 days from diagnosis ($n = 1$). There was one late death in a patient who developed chronic myeloid leukaemia 21 months from diagnosis. He was treated with three intensive blocks of AML-type chemotherapy with good initial response, but no further treatment was given because of severe cardiac toxicity. He subsequently progressed to develop an acute myeloblastic crisis and died 15 months later. Acute mucosal and myelotoxicity was significant, but manageable as previously reported (Patte et al, 1991).

**Survival**

With a median follow-up of 48 months (range 12–92), the overall survival and EFS are 87% (95% confidence interval (CI) 79.2–92.1%) and 83.7% (95% CI 76.3–89.2%) respectively (Figure 2). Results of univariate and multivariate analyses of EFS are shown in Table 2.

LDH levels at diagnosis were available in 79 patients with a range of 154–7575 iu l$^{-1}$ (median 1118). The median time to achieve CR was 70 days (range 7–181). The overall survival and EFS are high and differences in EFS according to the three sub-stages identified, LDH level or the time to achieve CR are small at 4 years from diagnosis (Figures 3, 4 and 5). The multivariate analysis does not appear to change the conclusions from the univariate analysis.

**DISCUSSION**

The consistent improvement in the prognosis of children with B-NHL has been mainly due to the use of higher dose systemic chemotherapy, frequent i.t. therapy and the reduction in early toxic death by avoiding unnecessary surgery coupled with aggressive treatment of biochemical and infectious complications soon after diagnosis or during induction (Lynch et al, 1977; Hann et al, 1990; Patte et al, 1991; Cairo et al, 1996; Miron et al, 1997). In the last decade the emphasis has been on trying to find prognostic factors that help to stratify patients and to deliver treatment according to risk group. The SFOP classification subdivided patients with stage

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**Table 2** Results of univariate and multivariate analyses of EFS

|                      | Log rank $P$-value | Relative risk | $P$-value multivariate |
|----------------------|--------------------|--------------|------------------------|
|                      | Univariate         |              |                        |
| LDH ($>2$ vs $<2$ ULN) | 0.75               | 0.85         | 0.76                   |
| CR                   | 0.47               | 1.33         | 0.71                   |
| (>28 days vs <28 days) | 0.82               | 0.56         | 0.64 (0.38)            |
| Substage (IIIA vs IIIB) |                   | 0.71         | (0.67)                 |
| (IIIX vs IIIB)       |                    |              |                        |

ULN – upper limit of normal

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**Figure 3** Event free survival in relation to the substage

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III disease into two main groups, namely IIIA and IIIB, according to the extent of primary disease (Philip et al, 1987). The German BFM group used a similar approach by dividing patients with stage III B-NHL into two risk groups based on LDH levels less than or more than 500 u l\(^{-1}\) (Reiter et al, 1995). Children with LDH less than 500 (R2 Regimen) and those with LDH more than 500 u

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**Figure 4** Event free survival in relation to serum LDH level at diagnosis

**Figure 5** Event free survival in relation to time to achieve CR
We conclude that with a short LMB type regimen of moderately intensive multi-agent chemotherapy, patients with a stage III and IV (blasts less than 70%) B-NHL have an excellent prognosis and as therapy has become more intensive and effective, the significance of various prognostic factors disappear. The priority in this group is to try and reduce both early and late morbidity. This is the aim of the current UK, French and US collaboration (FAB LMB 96) where reduction in both dose intensity and duration are evaluated in a randomized study. It is apparent that, with such effective chemotherapy, no clear prognostic subgroups can be identified at presentation. Something could emerge in the eventual analysis of FAB LMB 96 where treatment has been reduced for some patients, although few in number, the salvage rate for these patients remains relatively poor.

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