Feasibility of neuropsychological assessment in leukaemia patients shortly after diagnosis: directions for future prospective research

N C Jansen, A Kingma, P Tellegen, R I van Dommelen, A Bouma, A Veerman, W A Kamps

Aims: To study neuropsychological functioning of newly diagnosed children with acute lymphoblastic leukaemia (ALL) within two weeks after diagnosis in order to determine the feasibility of a sibling controlled prospective study design.

Methods: Fifty consecutive patients (median age at testing 6.6 years, range 4–12) were included in a prospective, longitudinal, nationwide study. Treatment would include intrathecal and systemic chemotherapy according to the DCLSG ALL-9 protocol. Children were evaluated with an extensive neuropsychological battery including measures of intelligence, memory, attention, language, visuoconstructive function, and fine-motor abilities within two weeks after start of the chemotherapy. The control group consisted of 29 healthy siblings (median age at testing 8.2 years, range 4–12), who were tested <4 weeks after the patients’ assessment.

Results: Mean scores on Wechsler Intelligence Scales did not differ significantly between patients and siblings; mean IQ scores for both the patients and the controls were high average. To examine specific neuropsychological functions, norm scores based on the exact age were acquired by fitting procedures, but no significant differences were found.

Conclusions: Neuropsychological assessment of patients during early hospitalisation is feasible. The results indicate no adverse effect of illness and psychological factors on IQ and neuropsychological functioning of patients with recently diagnosed ALL. The prospective design of this study of cognitive late effects of chemotherapy will allow discrimination between adverse sequelae of disease and treatment.

Acute lymphoblastic leukaemia (ALL) is the most common form of childhood cancer. Approximately 80% of newly diagnosed children with ALL are curable with modern treatment. Following this improved survival rate, an increasing number of studies has focused on the quality life of the survivors. Patients who have been treated with cranial irradiation (CI) and additional chemotherapy have shown intellectual deterioration and specific neuropsychological deficits. To date, prospective longitudinal studies on neuropsychological sequelae in children treated for ALL with chemotherapy only are rare or have yielded inconsistent results. These inconsistencies may be understood from less suitable control groups, different ages at time of testing (age effect), and selection of neuropsychological measures.

Moreover, pretreatment neuropsychological assessments are rarely conducted. It is commonly thought that testing shortly after diagnosis is not feasible because children diagnosed with ALL are seriously ill and have to cope with medical procedures and intensive treatment immediately after diagnosis. Leukaemia or leukaemia treatment can furthermore cause emotional, non-organic distress in patients and families, which may influence test behaviour of the children.

In 1999, we initiated a prospective longitudinal and nationwide study in the Netherlands, which includes siblings as controls, applies a comprehensive test battery, and has a broad age spectrum. Here we report the results of the neuropsychological assessment in patients shortly after diagnosis, and their healthy siblings. The results will eventually be used to investigate both early and late neuropsychological effects of chemotherapy according to the Dutch Childhood Leukaemia Study Group (DCLSG) ALL-9 protocol. In this report, we review the results of neuropsychological assessment shortly after diagnosis of both patients and their healthy siblings.

METHODS

Patients and sibling controls

From January 1999 to June 2001, 79 consecutive patients from six participating paediatric oncology centres in the Netherlands were eligible for this study. Criteria of eligibility were: newly diagnosed patients with high or standard risk ALL; age between 4 years and 12 years 3 months; and Dutch as primary language. Informed consent was obtained according to each hospital’s rules. Patients with initial CNS leukaemia and patients with pre-existent disorders that could interfere with normal cognitive development were excluded.

Sixteen (20%) parents refused participation because of the expected burden, and 19 (24%) cases were missed due to logistical problems.

Between March 1998 and January 1999, six consecutive patients has been enrolled in a pilot study in the hospital which coordinated the study. These patients did not significantly differ from the children in the main study; hence, a combined group of 50 patients entered the study.

The control group consisted of 25 healthy siblings who met the same inclusion criteria as the patients. If the patient had more than one sibling, the child (1) closest in age to the patient and (2) the same sex was chosen. Table 1 shows the characteristics of patients and siblings.

Abbreviations: ALL, acute lymphoblastic leukaemia; CI, cranial irradiation; DCLSG, Dutch Childhood Leukaemia Study Group; SES, socioeconomic status
visual-motor integration, attention, cognitive flexibility, cognitive functions as verbal-auditory and visual memory, hours, including measures of intelligence and specific psychological assessment of these children took about three weeks.

The neuropsychological testing was performed by one qualified child neuropsychologist who travelled to the hospitals where the children were tested. All participants were nationwide tested by one qualified child neuropsychologist who travelled to the hospitals where the children were treated.

Patients and healthy sibling controls were evaluated with an age appropriate comprehensive standardised neuropsychological test battery (table 2). Children aged 4–6 years were administered a developmental screening test and measures of intelligence, visual-motor integration, and, if >=5 years, fine-motor functioning. Participants aged 6–12 years were assessed with a more extensive test battery. The neuropsychological assessment of these children took about three hours, including measures of intelligence and specific cognitive functions as verbal-auditory and visual memory, visual-motor integration, attention, cognitive flexibility, verbal fluency, and fine-motor functioning. If necessary, the assessment was split into two sessions.

**Statistical analysis**

Performances of patients were compared to those of sibling controls using non-directional two tailed Student’s *t* tests for paired groups.

For the Wechsler Pre-School and Primary Scales of Intelligence (WPPSI-R), Experimental Dutch-Flemish version, and for the Wechsler Intelligence Scale for Children-revised (WISC-R, Dutch version), mean norm scores are 100 (SD = 15). For the remaining tests, norms have been acquired by fitting procedures based on the raw scores and the exact ages resulting in norm scores (mean = 50; SD = 10). The fitting procedures were based on the published norm data (means and standard deviations for different age groups) in the respective test manuals or other publications. This procedure enables comparisons of standardised scores between subjects of any specific age.21 22 Significance levels were established at *p* < 0.05. Statistical analyses were performed using version 10 of the Statistical Package for the Social Sciences (SPSS Inc., Chicago, IL, USA).

**RESULTS**

Included patients (*n* = 50) did not significantly differ from missing patients (*n* = 35) in terms of sex, age at diagnosis, and initial characteristics of disease and prognostic risk group. We had no indication of differences in socioeconomic status between included and missing patients. The latter mainly emanated from two hospitals; patients were missed due to illness of the psychologists who should have referred eligible patients. Patients and siblings aged 4–6 years at diagnosis were assessed as essentially normal on the Denver Developmental Scales. Patients aged 4–6 years scored significantly higher than siblings on WPPSI-R FS-IQ and WPPSI-R VIQ (table 3). Comparing patients aged 6–13 and siblings, no significant differences were found for any WISC-R factor. IQs were high average for patients on the WPPSI-R and both patients and siblings on the WISC-R.

Table 4 shows results for the remaining cognitive measures. No significant differences between the groups were found for any test. Overall, patients and siblings had average scores.

**Table 1** Characteristics of patients and siblings at the first neuropsychological evaluation shortly after diagnosis of ALL

<table>
<thead>
<tr>
<th>Group</th>
<th>Male n (%)</th>
<th>Female n (%)</th>
<th>Age at testing Median (range)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Patients</td>
<td>30 (60)</td>
<td>20 (40)</td>
<td>6.6 (4–12)</td>
</tr>
<tr>
<td>Healthy controls (siblings)</td>
<td>11 (38)</td>
<td>18 (62)</td>
<td>8.2 (4–12)</td>
</tr>
</tbody>
</table>

**Table 2** Neuropsychological battery

<table>
<thead>
<tr>
<th>Neuropsychological domain</th>
<th>Measures</th>
<th>Age</th>
<th>No. patients</th>
<th>No. siblings</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mental, motor, and social development</td>
<td>Denver Developmental Scales12</td>
<td>4–6</td>
<td>17</td>
<td>6</td>
</tr>
<tr>
<td>Intelligence</td>
<td>Wechsler Pre-School and Primary Scales of Intelligence (WPPSI-R), 10 subtests14</td>
<td>4–6</td>
<td>12</td>
<td>6</td>
</tr>
<tr>
<td></td>
<td>FS-IQ</td>
<td>26</td>
<td>23</td>
<td></td>
</tr>
<tr>
<td></td>
<td>V-IQ</td>
<td>30</td>
<td>23</td>
<td></td>
</tr>
<tr>
<td></td>
<td>P-IQ</td>
<td>26</td>
<td>23</td>
<td></td>
</tr>
<tr>
<td>Intelligence</td>
<td>Wechsler Intelligence Scale for Children (WISC-R), 10 subtests15</td>
<td>6–13</td>
<td>28</td>
<td>23</td>
</tr>
<tr>
<td></td>
<td>FS-IQ</td>
<td>26</td>
<td>23</td>
<td></td>
</tr>
<tr>
<td></td>
<td>V-IQ</td>
<td>30</td>
<td>23</td>
<td></td>
</tr>
<tr>
<td></td>
<td>P-IQ</td>
<td>26</td>
<td>23</td>
<td></td>
</tr>
<tr>
<td>Verbal-auditory learning and memory</td>
<td>Dutch version of Rey’s Auditory-Verbal Learning Test (RAVLT)16</td>
<td>6–13</td>
<td>26</td>
<td>23</td>
</tr>
<tr>
<td>Visual memory</td>
<td>Rey-Osterreith Complex Figure Test delayed (CFT) recall17</td>
<td>6–13</td>
<td>20</td>
<td>21</td>
</tr>
<tr>
<td></td>
<td>Animal naming fluency test</td>
<td>6–13</td>
<td>30</td>
<td>23</td>
</tr>
<tr>
<td>Sustained attention/speed</td>
<td>Bourdon-Vos; self-paced, continuous performance cancellation task13</td>
<td>6–13</td>
<td>47</td>
<td>28</td>
</tr>
<tr>
<td>Cognitive flexibility</td>
<td>Wisconsin Card Sorting Test (WCST)18</td>
<td>6–13</td>
<td>26</td>
<td>23</td>
</tr>
<tr>
<td>Perceptual-motor skills</td>
<td>Beery Developmental Test of Visual-Motor Integration (VMI)19</td>
<td>4–13</td>
<td>47</td>
<td>28</td>
</tr>
<tr>
<td></td>
<td>Rey-Osterreith Complex Figure Test (CFT) copy17</td>
<td>6–13</td>
<td>26</td>
<td>23</td>
</tr>
<tr>
<td>Fine-motor function</td>
<td>Purdue Pegboard</td>
<td>5–13</td>
<td>29</td>
<td>27</td>
</tr>
</tbody>
</table>
DISCUSSION

We have shown that recently diagnosed children with the life threatening disease ALL can be reliably assessed with an extensive standardised neuropsychological test battery shortly after diagnosis. An important observation in this study was that the majority, even the very young children, enjoyed the assessment, which was rather a distraction among numerous medical procedures than an emotional burden. Moreover, this study is strengthened by the inclusion of healthy siblings as controls, who were also pleased to be involved in the study and enjoyed the special attention. This control group enables appropriate comparison with the healthy population. Decrements in test results within the patient group can be detected, even if the results are still overrated, we could expect above average IQs in the siblings as well. There were no demographic differences explaining the IQ differences between patients and siblings aged 4–6 years. The scores of the children tested with the WISC-R are as well. The scores of the children tested with the WISC-R are

The present study can be criticised for the high number of missing patients, which could possibly account for bias in these test results. However, this is unlikely because included patients did not significantly differ from missed patients concerning demographic and initial disease characteristics. Missed patients should mainly have been referred by two ill psychologists. Fortunately, patients in these hospitals represent a random patient population, hence we have no indication that characteristics of the missed children differed from those who could be included. With the current numbers we would detect IQ differences of 0.7 SD (10.5 IQ points) to obtain an adequate power of 80%. To illustrate the meaning of 10.5 SD, a difference between 105 and 95 would be significant, but both IQs are considered average and children with both IQs would be in a regular school class. We could not control for SES. Given the overall average results, bias does not seem likely. In general, patients showed greater standard deviation on both the intelligence tests and neuropsychological tests. However, differences between patients and siblings did not result from one or few individuals with extreme scores.

The norm scores of the Experimental Dutch-Flemish version of the WPPSI-R were recently evaluated as disputable, which could explain the above-average IQs in the young patients (table 3). However, if the patients’ IQs are overrated, we could expect above average IQs in the siblings as well. There were no demographic differences explaining the IQ differences between patients and siblings aged 4–6 years. The scores of the children tested with the WISC-R are high average as a result of the Flynn effect, accounting for an average IQ difference of 5.1 SD, a difference between 105 and 95 would be significant, but both IQs are considered average and children with both IQs would be in a regular school class. We could not control for SES. Given the overall average results, bias does not seem likely. In general, patients showed greater standard deviation on both the intelligence tests and neuropsychological tests. However, differences between patients and siblings did not result from one or few individuals with extreme scores.

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<table>
<thead>
<tr>
<th>Test measures</th>
<th>Patients Mean (SD)</th>
<th>Siblings Mean (SD)</th>
<th>t value</th>
<th>p value</th>
<th>95% CI</th>
</tr>
</thead>
<tbody>
<tr>
<td>RAVLT Immediate recall</td>
<td>50.6 (8.0)</td>
<td>52.2 (8.7)</td>
<td>-0.67</td>
<td>0.506</td>
<td>-6.4 to 3.2</td>
</tr>
<tr>
<td>Delayed recall</td>
<td>52.4 (11.0)</td>
<td>49.2 (9.8)</td>
<td>1.06</td>
<td>0.295</td>
<td>-2.9 to 9.2</td>
</tr>
<tr>
<td>Fluency Test: animal naming</td>
<td>63.7 (10.5)</td>
<td>60.4 (9.7)</td>
<td>1.17</td>
<td>0.248</td>
<td>-2.4 to 9.0</td>
</tr>
<tr>
<td>Bourdon-Vos Speed</td>
<td>53.8 (14.3)</td>
<td>48.8 (8.7)</td>
<td>1.36</td>
<td>0.183</td>
<td>-2.5 to 12.4</td>
</tr>
<tr>
<td>Accuracy</td>
<td>51.0 (10.4)</td>
<td>49.7 (7.1)</td>
<td>0.45</td>
<td>0.658</td>
<td>-4.4 to 6.8</td>
</tr>
<tr>
<td>WCST Errors</td>
<td>49.4 (11.1)</td>
<td>50.7 (8.8)</td>
<td>-0.44</td>
<td>0.664</td>
<td>-7.1 to 4.6</td>
</tr>
<tr>
<td>Perseverations</td>
<td>49.3 (10.7)</td>
<td>50.8 (8.6)</td>
<td>-0.54</td>
<td>0.591</td>
<td>-7.1 to 4.1</td>
</tr>
<tr>
<td>Trials administered</td>
<td>50.0 (10.7)</td>
<td>50.7 (8.9)</td>
<td>-0.28</td>
<td>0.784</td>
<td>-6.5 to 4.9</td>
</tr>
<tr>
<td>Beery VMI</td>
<td>47.6 (9.7)</td>
<td>51.4 (11.6)</td>
<td>-1.52</td>
<td>0.133</td>
<td>-8.7 to 1.2</td>
</tr>
<tr>
<td>Rey-Osterreith CFT Copy</td>
<td>54.3 (5.9)</td>
<td>54.7 (5.5)</td>
<td>-0.26</td>
<td>0.795</td>
<td>-3.7 to 2.9</td>
</tr>
<tr>
<td>Delayed recall</td>
<td>44.5 (8.4)</td>
<td>46.3 (6.1)</td>
<td>-0.91</td>
<td>0.370</td>
<td>-6.5 to 2.5</td>
</tr>
<tr>
<td>Purdue Pegboard Dominant hand</td>
<td>45.5 (10.5)</td>
<td>45.7 (10.1)</td>
<td>-0.11</td>
<td>0.916</td>
<td>-5.8 to 5.2</td>
</tr>
<tr>
<td>Non-dominant hand</td>
<td>48.3 (8.9)</td>
<td>46.4 (8.6)</td>
<td>0.81</td>
<td>0.424</td>
<td>-2.8 to 6.4</td>
</tr>
<tr>
<td>Both hands</td>
<td>49.0 (11.1)</td>
<td>48.3 (7.3)</td>
<td>0.25</td>
<td>0.804</td>
<td>-4.5 to 5.7</td>
</tr>
<tr>
<td>Assembly</td>
<td>52.5 (11.7)</td>
<td>51.9 (9.1)</td>
<td>0.20</td>
<td>0.840</td>
<td>-5.1 to 6.3</td>
</tr>
</tbody>
</table>
IQ rise of about 6 points, since test norms were collected in the early 1980s.22 If evaluated with more recent test norms, these children would probably have average results.

Generally, it is often suggested that emotional, non-organic distress influences the test results. However, such an effect is very unlikely given the normal outcome. Even measures of attention and memory, known to be sensitive for emotional distress,23 did not differ between patients and siblings.

Conclusion

The present data strongly suggest that patients do not suffer from neuropsychological deficits related to acute disease or early treatment. In the future, patients’ baseline scores can be used to discriminate between possible adverse sequelae of disease and/or treatment and eventually, to optimise treatment protocols compromising between high cure rate and good quality of life. Ideally, neuropsychological assessment early after hospitalisation also selects patients who need early intervention for mental or academic deficits, but this was not the aim of this study.

Neuropsychological assessment of children with ALL shortly after diagnosis with sibling controls is feasible and essential to discriminate between adverse sequelae of treatment. Prospective, longitudinal study designs should become the standard for evaluating possible treatment effects.

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