Response to influenza immunisation during treatment for cancer

J C Chisholm, T Devine, A Charlett, C R Pinkerton, M Zambon

Abstract
Aims—To assess the annual risk of influenza infection in children with cancer and the immunogenicity of a trivalent split virus influenza vaccine in these children.

Methods—Eighty four children with cancer were tested for susceptibility to the circulating strains of influenza virus in autumn 1995 and 1996. Non-immunised children were reassessed following the spring for serological evidence of natural infection. Forty two patients received two doses of influenza vaccine. These children were receiving continuing chemotherapy for acute lymphoblastic leukaemia or were within six months of completing chemotherapy.

Results—Among the 84 children tested for influenza virus susceptibility only 8% of patients were fully protected (antibody titres ≥ 40) against all three of the prevalent influenza virus strains; 33% were susceptible to all three viruses. Evidence of acquired natural infection was seen in 30% of unimmunised patients. Among immunised susceptible patients, 66% made some protective response to the vaccine and 55% showed protective antibody titres to all three viral strains following vaccination. Older age was associated with increased response to the H1N1 and H3N2 vaccine components, but total white cell count or neutrophil count at immunisation, type of cancer, or length of time on treatment for acute lymphoblastic leukaemia did not affect response.

Conclusions—Most children with cancer studied were at risk of influenza infection. A significant response to immunisation was seen, supporting annual influenza vaccination for children being treated for cancer.

Keywords: influenza; immunisation; susceptibility; cancer

Influenza is a common cause of respiratory tract infection and hospitalisation during the influenza season in adult patients with leukaemia,1 2 and following bone marrow transplantation,3 4 where a high incidence of complications, especially pneumonia, and significant mortality have been reported. Annual influenza infection rates in children vary from 18% to 48%,5 but the infection is more common in children with cancer than in healthy control children.6 Although in children with cancer the illness usually runs a mild course it may result in hospitalisation and the interruption of cancer therapy.6 7 Severe and fatal complications have also been reported in these children.6 8

British9 and American10 guidelines recommend annual influenza vaccination of high risk groups, including immunosuppressed children and adults, but uptake in high risk patients remains low.9 11 Contributing to the poor uptake is uncertainty over the efficacy of the vaccine in certain patient groups.9 Indeed, previous data suggest an impaired response to influenza immunisation in children on treatment for cancer compared with healthy children.12

This study is the first to report on the response to a trivalent split virion influenza vaccine in children during cancer therapy. The inclusion of a control, unvaccinated group in whom paired sera were obtained allowed assessment of the annual risk of disease in our population.

In the first year an unselected group of all consenting patients attending the paediatric oncology outpatient department or resident on the paediatric inpatient ward between 28 September 1995 and 20 October 1995 was tested for susceptibility to the prevalent strains of influenza virus (A/Taiwan/1/86 [H1N1], A/Johannesburg/34/94 [H3N2], and B/Beijing/184/93 [B]). Susceptibility was defined as antibody titre < 40 (see below). Follow up serum was taken in May 1996 to document seroconversion in unimmunised patients. No attempt was made to document respiratory infections in these patients.

Among patients showing susceptibility to one or more of the prevalent virus strains, the following subgroups were offered immunisation in November 1995: (1) patients with acute lymphoblastic leukaemia (ALL) between weeks 9 and 18 of the UKALL XI treatment schedule; (2) patients with ALL between weeks 24 and 100 of the UKALL XI treatment schedule except those receiving the third intensification block (that is, on maintenance chemotherapy, between first and second intensification blocks); (3) all patients who had finished chemotherapy within the past six months (including patients with leukaemia, solid tumours, and bone marrow transplant recipients). The following
children were ineligible for vaccination: age less than 6 months; neutropenic (less than 1.0 × 10⁹/l) or lymphopenic (less than 1.0 × 10⁹/l) at time of vaccination; previously vaccinated against influenza; or egg allergy. Serum was taken to assess response four to six weeks after the second vaccination.

In the second study year vaccination was offered to the same subgroups of patients as above. The vaccine was also given to one patient with relapsed ALL and one child receiving chemotherapy for a brain tumour. Previously vaccinated patients were excluded. Prevacination and 4–6 week post-vaccination sera were obtained, although, in this cohort, the vaccine was given without knowing the immune status.

All immunised children received two doses of inactivated influenza vaccine (split virion; Aventis Pasteur MSD), subcutaneously, four weeks apart (0.5 ml for children over 4 years and 0.25 ml for children ≤ 4 years). In 1995 the influenza vaccine strains were A/Taiwan/1/86, A/Johannesburg/34/94, and B/Beijing/184/93 according to the recommendations of the World Health Organisation. In 1996, A/Wuhan/359/95 replaced the previous H3N2 component.

Sera were analysed in the Enteric, Respiratory, and Neurological Virus Laboratory, Central Public Health Laboratory, Colindale. Antibodies to the prevalent A and B strains were tested by haemagglutination inhibition (HI). Responses were assessed in terms of geometric mean titre (GMT) and protective response. Antibody levels were expressed on a doubling scale (10, 20, 40, 80, etc) with a level of 40 or more considered as protective.

**STATISTICS**

McNemar’s test of paired proportions was used to assess increase in proportion of children exhibiting a protective response. For each strain the child was placed into one of three categories: immune prior to vaccination; protective response made to vaccine; and susceptible after vaccination. Associations with the response made to the vaccine were assessed using the Kruskal–Wallis test and Fisher’s exact test for continuous and categorical variables respectively.

A within subject linear model was used to assess increase in antibody titre. The dependent variable in this analysis was the HI titre, which was log transformed to remove skewness in this variable. For sera where the antibody level was below the lower limit of detection (HI = 10) a value of 5 was used in this analysis. A blocking factor was fitted for subject. The pre/post-vaccination factor and the interaction between this factor and the variables that may influence response were fitted as within subject effects. The assumption of normally distributed residuals and equality of residual variance were assessed using the Shapiro–Francia W’ test and the Cook–Weisburg test respectively.

**RESULTS**

Sera were taken from 67 children in 1995. Of these children, 25 fulfilled the criteria for vaccination and received influenza vaccine. Paired sera were obtained in 27/42 children who were not eligible for immunisation. The remaining 15 children had no follow up serum taken. Twelve of these children died from progressive disease, one was followed up elsewhere, and in two a follow up sample was never taken. A further 17 children were immunised in 1996. Table 1 shows diagnoses of children in the study.

**SUSCEPTIBILITY ANALYSIS**

Among 84 children tested, 60 (71%) were susceptible to H1N1, 47 (56%) to H3N2, and 58 (69%) to B. Although the greatest pre-existing protection was to H3N2, half of all patients tested were susceptible to this virus. Twenty eight patients (33%) were susceptible to all three viruses and only seven patients (8%) showed protective titres against all three viruses prior to immunisation (data not shown).

Among the 27 unimmunised patients in whom paired sera were available, eight of 26 (30%) patients susceptible to at least one virus showed evidence of exposure to influenza A over the winter months (four to H1N1, four to H3N2), developing antibody where none was detected previously. Only one of these patients had a blood product transfusion in the three months before the second serum sample, indicating that seroconversion was the result of natural infection.

Ten of 19 (53%) unimmunised patients with pre-existing full or partial protection lost protective antibody levels to one or more virus types over the winter months. Five patients had blood or platelet transfusions in the two weeks before the initial blood test and may have had false positive tests on the initial screening. However, the other five patients had no history of recent blood product transfusion.

**RESPONSE TO IMMUNISATION**

Forty two patients were immunised with influenza vaccine. No adverse events were reported. Prior to vaccination, only four patients (5%) were fully protected against all three viruses (all had been immunised in 1996 without knowing status), whereas post-vaccination 23 patients (55%) were fully protected. Partial protective responses to immunisation were seen in a
Table 2  Numbers of children making response to each virus subunit

<table>
<thead>
<tr>
<th>Virus</th>
<th>Pre-vaccination</th>
<th>Post-vaccination</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>HI &lt; 40</td>
<td>HI ≥ 40</td>
</tr>
<tr>
<td>H1N1</td>
<td>15</td>
<td>14</td>
</tr>
<tr>
<td>H3N2</td>
<td>7</td>
<td>16</td>
</tr>
<tr>
<td>B</td>
<td>10</td>
<td>18</td>
</tr>
<tr>
<td>Total</td>
<td>32</td>
<td>42</td>
</tr>
</tbody>
</table>

Table 3  Response to immunisation by geometric mean titres

<table>
<thead>
<tr>
<th>Virus</th>
<th>Pre-vaccination Geometric mean</th>
<th>95% CI</th>
<th>Post-vaccination Geometric mean</th>
<th>95% CI</th>
<th>p value</th>
</tr>
</thead>
<tbody>
<tr>
<td>H1N1</td>
<td>12.6</td>
<td>8.6 to 19.2</td>
<td>12.0</td>
<td>6.0 to 21.6</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>H3N2</td>
<td>13.2</td>
<td>13.2 to 40.8</td>
<td>12.0</td>
<td>10.0 to 40.8</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>B</td>
<td>10.4</td>
<td>12.4 to 21.6</td>
<td>8.0</td>
<td>6.0 to 16.0</td>
<td>&lt;0.0001</td>
</tr>
</tbody>
</table>

Discussion

This study confirms previous reports that children with cancer are highly susceptible to influenza virus. Our estimated natural infection rate of 30% in unimmunised children is similar to that of Borella and Webster, who reported that 46% of unimmunised children with leukaemia suffered with a flu-like illness in one influenza season. The loss of protective antibody levels in some unimmunised patients noted in this study, possibly as the result of immunosuppressive therapy, might contribute to the increased risk of clinical infection in children with cancer noted in previous studies.7 13

Previous studies of influenza vaccination in children with cancer have used a variety of different vaccines and have involved both children on treatment for cancer and children who have completed their treatment.14–21 All these studies have shown the safety of the vaccine in this patient group. However, the immunogenicity of influenza vaccine in children on chemotherapy varies in different reports according to the type of vaccine used and the viral strains involved. It is clear that two vaccine doses produce a better response than a single dose,14 15 and prior exposure to the circulating viral strains seems to increase the likelihood of response to the vaccine.14 16 Protective antibody levels to individual viral strains following immunisation are reported in 29–75% of children on chemotherapy,16–18 compared to around 70–90% of healthy children.21 The results of this study are in keeping with an impaired response to vaccination in children with cancer, but nevertheless show a very useful response to immunisation. Similar impaired but useful response to influenza vaccine has been seen in some groups of adult patients receiving chemotherapy.15 22–25

By contrast with patients on treatment for cancer, children who have completed their treatment seem to show response rates more comparable with those expected in healthy controls.16–18 For H3N2, the response was better in patients off treatment, although the difference was not statistically significant in the small number of patients off treatment in this study. The situation is different in bone marrow transplant recipients, a particularly high risk group with prolonged immunological impairment, who show no response at all until at least six months post-transplant.27

In our subgroup analysis we found no significant effect of tumour type (solid tumour versus ALL) on response to vaccination, but again a suggestion that response to H3N2 was better in solid tumour patients. As all the solid tumour patients were off treatment, the effect of tumour type could not be isolated from on versus off treatment. Children with solid tumours on treatment showed a significantly better response to A/New Jersey/8/76 than children being treated for ALL, although no difference was seen for A/Victoria/3/75.16 Differences in response between patients with solid and haematological malignancies could...
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relate to the intrinsic disease associated immunosuppression of some haematological malignancies as well as the type and intensity of chemotherapy treatment. With the increased intensity of many chemotherapy regimes for solid tumours in recent years, it would be relevant to study response to a trivalent vaccine in a larger number of solid tumour patients currently on treatment.

Our study showed an interesting relation between age and response to immunisation for H1N1 and H3N2. In the two to three years prior to this study, H3N2 was the predominant circulating influenza strain, with limited H1N1. Influenza B had been largely absent.16,20 The effect of age on response to H3N2 and H1N1 may therefore largely reflect previous exposure or "priming", with older children more likely to have been exposed. Gross found no difference in response to split virus H1N1 between 3–5 and 6–18 year olds where H1N1 had been absent from recent natural circulation,25 but Wright et a. showed a non-significant trend of better response with increasing age for H3N2 and B where these had been the previous circulating strains.24 These results support a predominant effect of prior exposure rather than age alone on response.

One remaining area of uncertainty is whether an antibody level of 40, normally considered protective in healthy individuals, is actually protective in the immunocompromised host. In one study, 24% of immunised children with cancer developed proven influenza infection despite influenza virus titres of >1/32,2 but another study reported an incidence of flu like illness in only 10% of immunised children with leukaemia, compared to 46% of unimmunised children, suggesting that immunisation was associated with a reduced incidence of clinical infection.13 The latter study showed a rise in antibody titres following immunisation, but post-immunisation titres were not discussed in relation to clinical infection in individual patients. The discrepancy between the studies could be explained if there is high rate of loss of protective antibody levels in patients receiving chemotherapy, a finding shown in the current study. If the latter explanation is correct, it may be that annual immunisation prior to the influenza season in children with cancer is of greater benefit than suggested by the change in the proportions of patients with protective antibody titres following immunisation, because it may reduce loss of protective immunity in some patients as well as facilitating new protective immunity in others. In one study of leukaemic children on and off treatment, antibody levels returned to baseline by 12 months following vaccination,17 but in other children, mainly off treatment, levels were maintained at least until six months following vaccination,17 suggesting that protection would be maintained during the highest risk time for influenza even if it is subsequently lost. Taken with previous data, our study suggests that the risk of influenza infection in children with cancer may be reduced substantially by immunisation.

Even though influenza usually runs a mild course in children with cancer, the occasional morbidity and mortality and high risk of interruption of potentially curative chemotherapy make it worth immunisation. It seems reasonable that the household contacts of such children and the hospital staff caring for them, as well as the children themselves, should receive annual immunisation against influenza.19,20 Although to our knowledge there is no direct evidence to support this approach. When proven influenza occurs despite immunisation, judicious use of zanamivir may reduce hospital admission rates and complications in high risk patients, but data in these groups are lacking.35 In conclusion, this study supports existing recommendations that all children with cancer receiving chemotherapy should receive annual immunisation with influenza vaccine. We suggest that their household contacts and appropriate hospital staff should also be immunised. Two doses of vaccine should be given at the first immunisation but data are needed on whether one or two doses of vaccine are required for second and subsequent vaccinations in patients on chemotherapy. Further data are also needed on the efficacy of the vaccine in children with solid tumours on treatment.

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Immunisation debates

Whether we like it or not “the media” are the most powerful influence on public opinion. There have been times when unbalanced reporting has done harm but recently (December 2000–January 2001) there has been evidence of a well-considered approach, at least to the subject of immunisation. The apparent success of group C meningococcal vaccination has been well publicised and MMR vaccination defended. Commenting on a recent fall in MMR acceptance rates, the Times concluded (January 5, 2001) that parents who refuse the vaccine for their children “shirk, for selfish reasons and on the basis of wholly inconclusive scientific research, a manifest social responsibility”. Strong stuff, and probably unfair to many concerned parents, but at least fighting on the side of the angels.

Whenever questions are raised about immunisation there are fears about vaccine refusal, the great spectre being the example of pertussis in the early 1980s when disease rates soared after fears of vaccine-induced encephalopathy resulted in low acceptance rates. But questions must be asked. It may be irresponsible to endanger vaccination programmes on the basis of inadequate data but it is never irresponsible to ask sensible questions.

There is no doubt that parents do have their own thoughts about immunisation. In a national survey in the USA (Bruce G Gellin and colleagues. Pediatrics 2000;106:1097–102) 23% agreed that “children get more immunisations than are good for them”, and 25% agreed, or expressed concern, that “too many immunisations could weaken the child’s immune system”. Could such concerns have any rational basis? Immunisation almost certainly has non-specific effects on immune function. Both measles and BCG immunisations probably boost immunity in non-specific ways and so may other immunisations. In a randomised study in immune function. Both measles and BCG immunisations probably boost immunity in non-specific ways and so may other immunisations.

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