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Decline in mortality, AIDS, and hospital admissions in perinatally HIV-1 infected children in the United Kingdom and Ireland

D M Gibb, T Duong, P A Tookey, M Sharland, G Tudor-Williams, V Novelli, K Butler, A Riordan, L Farrelly, J Masters, C S Peckham, D T Dunn, on behalf of the National Study of HIV in Pregnancy and Childhood (NSHPC) and the Collaborative HIV Paediatric Study (CHIPS)

Abstract

Objective To describe changes in demographic factors, disease progression, hospital admissions, and use of antiretroviral therapy in children with HIV.

Design Active surveillance through the national study of HIV in pregnancy and childhood (NSHPC) and additional data from a subset of children in the collaborative HIV paediatric study (CHIPS).

Setting United Kingdom and Ireland.

Participants 944 children with perinatally acquired HIV-1 under clinical care.

Main outcome measures Changes over time in progression to AIDS and death, hospital admission rates, and use of antiretroviral therapy.

Results 944 children with perinatally acquired HIV were reported in the United Kingdom and Ireland by October 2002; 628 (67%) were black African, 205 (22%) were aged ≥10 years at last follow up, 193 (20%) are known to have died. The proportion of children presenting who were born abroad increased from 20% in 1994-5 to 60% during 2000-2. Mortality was stable before 1997 at 9.5 per 100 child years at risk but fell to 2.0 in 2001-2 (trend P < 0.001). Progression to AIDS also declined (P < 0.001). From 1997 onwards the proportion of children on three or four drug antiretroviral therapy increased. Hospital admission rates declined by 80%, but with more children in follow up the absolute number of admissions fell by only 26%.

Conclusion In children with HIV infection, mortality, AIDS, and hospital admission rates have declined substantially since the introduction of three or four drug antiretroviral therapy in 1997. As infected children in the United Kingdom and Ireland are living longer, there is an increasing need to address their medical, social, and psychological needs as they enter adolescence and adult life.

Introduction

Since 1996 antiretroviral therapy with three or four drugs has been increasingly used to treat adults infected with HIV,1 3 and its substantial effect on progression of the disease has been well described.4 5 Because of difficulties in developing appropriate formulations and the lack of age specific pharmacokinetic data to guide paediatric dosing the introduction of such treatment was delayed in children. Before 1994 only one antiretroviral drug (zidovudine) was available for children. Dual antiretroviral therapy was introduced in 1995, and treatment with three or four drugs followed from mid-1997 and is now standard.6 7 Reductions in mortality and AIDS during the era of dual therapy were reported,8 9 and substantial declines attributed to therapy with three or four drugs were reported in a US paediatric cohort.10 However, data on changing rates in disease progression since 1997 and their effect on use of medical services remain sparse in children. We investigated changes over time in mortality, morbidity, hospital admission rates, and antiretroviral therapy received in perinatally HIV-1 infected children in the United Kingdom and Ireland.

Methods

Study design

The national study of HIV in pregnancy and childhood (NSHPC) is informed about children who present in the United Kingdom and Ireland with HIV-1 infection and infants born to HIV infected women through two active confidential reporting schemes run in collaboration with the British Paediatric Surveillance Unit and the Royal College of Obstetricians and Gynaecologists.11 Obstetric data include demographic information, timing of maternal diagnosis, uptake of interventions, and outcome of pregnancy; paediatric data include neonatal details and HIV status. Each year notifying paediatricians subsequently provide minimal clinical information for infected children.

In April 2000, the collaborative HIV paediatric study (CHIPS) was established between the national surveillance study at the Institute of Child Health, London, and the Medical Research Council Clinical Trials Unit, where trials in the Paediatric European Trials Network for Treatment of AIDS (PENTA) are coordinated (www.pentatrials.org). Most perinatally infected children reported to the national surveillance...
study are followed up in 17 paediatric clinics involved in PENTA. The collaborative HIV paediatric study was established to collect more detailed clinical, laboratory, and treatment information on infected children seen at the PENTA centres since 1996. During 2000, retrospective information was collected from clinical records on to standard forms; since then, information has been collected annually by questionnaire and merged with additional data from the national surveillance study and PENTA trials. The merged dataset includes demographic information, data on clinical events, hospital admissions, outpatient visits, antiretroviral therapy, and results of T cell subset and HIV RNA viral load tests. AIDS was defined according to the classification of the Centers for Disease Control (clinical category C disease).\(^1\)

Paediatric HIV surveillance started in 1986, and by October 2002, 944 perinatally infected children had been reported to the national surveillance study. All children were included in analyses of changes in progression to AIDS and death over time. In total 593 children alive in January 1996 were enrolled in the collaborative HIV paediatric study (75% of all children reported to the national surveillance study as alive on this date) and were included in analyses of antiretroviral therapy and hospital admissions.

### Statistical methods

For children whose mothers’ HIV-1 status was known before delivery, we considered the date of birth to be the date of the child’s first presentation to medical services with HIV-1. For remaining children, we estimated the date of presentation using the earliest date of: first positive results of HIV-1 antibody test, T cell subsets, or viral load tests; HIV related clinical events; clinic visit; or hospital admission. In all analyses, each child began to contribute to follow up from his or her date of presentation.

We censored records at the date children were last known to be alive in the death analysis and at the date of last clinical assessment in the AIDS/death analysis. To reduce presentation bias, we excluded children not identified at birth from analyses of AIDS or death if they first presented within one month of AIDS diagnosis or death, respectively. The effect of calendar period was further examined by using Cox proportional hazards models, adjusted for how the children were identified (at birth or later), ethnicity, sex, and place of birth (United Kingdom and Ireland or abroad). We also carried out separate adjusted analyses by place of birth. Time was measured from birth by using late entry at the age of first presentation.\(^1\) In any calendar period, children were considered at risk only from the first day of last clinical assessment in the AIDS/death analysis.

### Table 1 Characteristics of perinatally HIV-1 infected children in the United Kingdom and Ireland up to October 2002 according to place of birth. Figures are numbers (percentages) of children

<table>
<thead>
<tr>
<th>Region of follow up</th>
<th>Born in UK or Ireland (n=586)</th>
<th>Born abroad (n=353)</th>
<th>Unknown (n=0)</th>
<th>Total (n=944)</th>
</tr>
</thead>
<tbody>
<tr>
<td>London</td>
<td>416 (71)</td>
<td>248 (70)</td>
<td>4</td>
<td>668 (71)</td>
</tr>
<tr>
<td>Rest of England</td>
<td>92 (16)</td>
<td>73 (20)</td>
<td>0</td>
<td>165 (17)</td>
</tr>
<tr>
<td>Scotland</td>
<td>36 (6)</td>
<td>8 (2)</td>
<td>0</td>
<td>44 (5)</td>
</tr>
<tr>
<td>Ireland</td>
<td>35 (6)</td>
<td>18 (5)</td>
<td>1</td>
<td>54 (6)</td>
</tr>
<tr>
<td>Wales and N Ireland</td>
<td>7 (1)</td>
<td>6 (2)</td>
<td>0</td>
<td>13 (2)</td>
</tr>
</tbody>
</table>

### How child was identified:

- Prospective from birth: 121 (21)
- After birth, asymptomatic: 157 (27)
- After birth, symptomatic: 287 (49)
- Unknown: 21 (4)

### Age at first presentation (years):

- At birth: 121 (21) 11 (3) 0 132 (14)
- <1: 254 (43) 27 (8) 0 281 (30)
- 1-4: 203 (35) 19 (5) 0 222 (23)
- 5-9: 100 (17) 123 (21) 2 225 (23)
- ≥10: 6 (1) 41 (12) 1 48 (5)

### Table 1 Characteristics of perinatally HIV-1 infected children in the United Kingdom and Ireland up to October 2002 according to place of birth. Figures are numbers (percentages) of children

- Median (range): 0.46 (0.0-13.4) 4.6 (0.2-15.3) 5.0 (2.7-12.8) 1.6 (0.0-15.3)
- Median (range): 0.46 (0.0-13.4) 4.6 (0.2-15.3) 5.0 (2.7-12.8) 1.6 (0.0-15.3)

### Age at last follow up (years):

- ≤1: 104 (18) 11 (3) 0 115 (12)
- 1-4: 203 (35) 99 (28) 2 304 (32)
- 5-9: 158 (24) 121 (34) 1 300 (34)
- ≥10: 67 (11) 97 (27) 2 166 (18)
- ≥15: 14 (2) 26 (7) 0 39 (4)

### Stage of disease at last follow up:

<table>
<thead>
<tr>
<th>Stage</th>
<th>Born in UK or Ireland (n=586)</th>
<th>Born abroad (n=353)</th>
<th>Unknown (n=0)</th>
<th>Total (n=944)</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>275 (47)</td>
<td>226 (64)</td>
<td>5</td>
<td>506 (54)</td>
</tr>
<tr>
<td>B</td>
<td>159 (27)</td>
<td>86 (24)</td>
<td>0</td>
<td>245 (26)</td>
</tr>
<tr>
<td>C</td>
<td>152 (26)</td>
<td>41 (12)</td>
<td>0</td>
<td>193 (20)</td>
</tr>
</tbody>
</table>

*According to classification of Centers for Disease Control.\(^1\)
of the calendar period or the date of presentation, if later. We fitted calendar period as a time varying covariate, with adjustment for an interaction with age.

We used Kaplan-Meier curves to compare survival after initial diagnosis of AIDS by first indicator disease, grouped in hierarchical order: Pneumocystis carinii pneumonia, other opportunistic infection, HIV encephalopathy, failure to thrive, severe recurrent bacterial infections. We excluded six children in whom cancer was the first indicator disease. Hospital admission rates were compared with Poisson regression.

Results

Half of the 944 children reported were girls and 628 (67%) were black African (table 1). The proportion of children born abroad increased from 20% in 1994-5 to 60% in 2000-2. In total 132 (14%) children were identified at birth, 281 (30%) during the first year, and 202 (21%) at ≥5 years. Whereas the median age at presentation among children born in the United Kingdom and Ireland has remained constant at around 6 months, among children born abroad it increased from 2.5 years before 1991 to 5.2 years in 1994-5 and remained constant thereafter. At last follow up, 205 (22%) children were aged ≥10 years and 39 (4%) ≥15 years (table 1). A total of 193 children are known to have died (including 54 since 1997); and 41 left the country or were otherwise lost to follow up before 1996. Among 593 children in the collaborative HIV paediatric study, 534 (90%) were in follow up in 2001-2, 30 (5%) in 2000, and 29 (5%) were last seen in 1999 or earlier.

Mortality and progression to AIDS

While the number of perinatally infected children in follow up has increased steadily over time, the annual number of deaths declined markedly after 1996 (table 2). Among 734 children identified at birth, or who survived at least a month after presentation, the crude mortality was stable before 1997 at 9.3 per 100 child years but declined thereafter to 3.3, 2.7, 1.2, 1.3, and 2.0 in 1997, 1998, 1999, 2000, and 2001-2, respectively (log rank test for trend, P < 0.001). Figure 1 shows the change in risk relative to 1996, adjusted for place of birth, ethnicity, sex, and whether the child was identified at birth or later. However, improvement in survival since 1997 was more marked in children aged >1 year (adjusted hazard ratio 0.19, 95% confidence interval 0.12 to 0.30) than in those aged <1 year (log rank test for trend, P < 0.001). Figure 1 shows adjusted hazard ratios were similar when we carried out separate analyses for children aged >1 year (adjusted hazard ratio 0.19, 95% confidence interval 0.12 to 0.30) than in those aged <1 year (0.72, 0.35 to 1.48; P = 0.003, heterogeneity for age) (table 3).

Information on antiretroviral therapy was available for 36 children who died; 14 (including six infants) had not received three or four drug therapy, six died within two months of starting it, and 16 died a median of 10 (range 2-53) months after starting it.

A total of 438 children developed an AIDS indicator disease, but 13 of these died before AIDS was diagnosed. Two hundred and seventeen children (50%) developed AIDS within one month of presentation (table 2). Of the children with AIDS, 119 (28%) had more than one indicator disease for AIDS at diagnosis. The proportion of children with opportunistic infections was higher in 1997-2002 than previously (57% versus 29% for P carinii pneumonia; 57% versus 22% for other opportunistic infections), but the distribution of other indicator diseases was similar in the two periods. Prognosis from initial AIDS diagnosis varied: mortality was higher in children with P carinii pneumonia or HIV encephalopathy compared with mortality in children with other opportunistic infections, failure to thrive, or severe recurrent bacterial infection (fig 2).

Among 587 children who had not progressed to AIDS within one month of presentation after birth, crude progression rates declined from 15.9 per 100 child years before 1997 to 3.0 per 100 child years at

![Fig 1 Risk of death or AIDS/death by calendar year relative to 1996, adjusted for age, sex, ethnicity, place of birth, whether the child was identified prospectively from birth or later (y axis on log scale)](image-url)

![Table 2 Deaths and children with AIDS by calendar time](table-url)

<table>
<thead>
<tr>
<th>Year</th>
<th>No in follow up</th>
<th>No who died</th>
<th>No developing first AIDS indicator disease*</th>
</tr>
</thead>
<tbody>
<tr>
<td>1989</td>
<td>69</td>
<td>17</td>
<td>30</td>
</tr>
<tr>
<td>1990</td>
<td>80</td>
<td>4</td>
<td>12</td>
</tr>
<tr>
<td>1991</td>
<td>108</td>
<td>9</td>
<td>17</td>
</tr>
<tr>
<td>1992</td>
<td>158</td>
<td>15</td>
<td>27</td>
</tr>
<tr>
<td>1993</td>
<td>207</td>
<td>24</td>
<td>38</td>
</tr>
<tr>
<td>1994</td>
<td>252</td>
<td>18</td>
<td>46</td>
</tr>
<tr>
<td>1995</td>
<td>283</td>
<td>23</td>
<td>35</td>
</tr>
<tr>
<td>1996</td>
<td>326</td>
<td>29</td>
<td>37</td>
</tr>
<tr>
<td>1997</td>
<td>356</td>
<td>12</td>
<td>42</td>
</tr>
<tr>
<td>1998</td>
<td>427</td>
<td>14</td>
<td>40</td>
</tr>
<tr>
<td>1999</td>
<td>465</td>
<td>6</td>
<td>24</td>
</tr>
<tr>
<td>2000</td>
<td>524</td>
<td>6</td>
<td>38</td>
</tr>
<tr>
<td>2001-2</td>
<td>606</td>
<td>16</td>
<td>53</td>
</tr>
<tr>
<td>Total</td>
<td>944</td>
<td>193</td>
<td>438</td>
</tr>
</tbody>
</table>

*Includes 13 children who died of HIV-related causes without diagnosis of AIDS. 217 (50%) children developed AIDS or died within one month after presentation with HIV infection; a further eight children born abroad first experienced an AIDS indicator disease at least one month before arrival in UK or Ireland.
risk in 2001-2 (test for trend P < 0.001). Figure 1 shows the adjusted risks relative to 1996. As with mortality, this decline was more marked in children aged >1 year (hazard ratio 0.37, 95% confidence interval 0.25 to 0.53, for before 1997 compared with 1997 onwards, table 3), while progression rates were similar in both periods for children aged <1 (1.03, 0.60 to 1.77, P = 0.002, heterogeneity for age).

Antiretroviral therapy
Of 593 children followed since 1996 in the collaborative HIV paediatric study, 137 (23%) had not started antiretroviral therapy at last report (121) or by the time of death (16). The percentage of child time spent on three or four drug antiretroviral therapy increased from 1% in 1996 to 56% in 1999 and 69% in 2001-2 (fig 3). There was no evidence that this differed in children aged <1. There was a shift over time from initial drug regimens containing a protease inhibitor to those containing a non-nucleoside reverse transcriptase inhibitor (table 4). The proportion of children who had previously been on therapy but had currently stopped remained constant after 1996 at around 5% of child time (fig 3).

Among 371 children on antiretroviral therapy when they were last seen in 2000 or later, 48 (13%) were taking four drugs, 301 (81%) three, and 22 (6%) two. Overall 91 different drug combinations were used.

Hospital admissions
The number of children seen in collaborative HIV paediatric study centres increased from 299 during 1996 to 493 in 2001-2, while the number of hospital admissions fell by 26% from 350 to 258. Admission rates declined by 80% from 4.4 per 100 child years of follow up in 1996 to 0.9 in 2001-2 (test for trend P < 0.001). As expected, the rates were lower in children born to mothers whose HIV status was known in pregnancy compared with those in whom it was not (relative risk 0.49, 0.42 to 0.57, P < 0.001). Among those presenting after birth, rates of admission were higher in the first six months after presentation than later (4.21, 3.79 to 4.69, P < 0.001).

Discussion
Mortality
From 1997 onwards we have seen reductions of around 80% in mortality and 50% in progression to AIDS among children perinatally infected with HIV-1 in the United Kingdom and Ireland. In the collaborative HIV paediatric study, hospital admission rates fell substantially. The introduction and increased

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**Table 3** Adjusted hazard ratios (95% confidence intervals) for factors associated with progression to death and AIDS/death

<table>
<thead>
<tr>
<th>Calendar year</th>
<th>Death Ratio</th>
<th>P value</th>
<th>AIDS/death Ratio</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Before 1997</td>
<td>1</td>
<td></td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>1997-2002</td>
<td>0.26 (0.17 to 0.36)</td>
<td>P&lt;0.001</td>
<td>0.49 (0.36 to 0.67)</td>
<td>P&lt;0.001</td>
</tr>
<tr>
<td>1997-2002, age &lt;1 year</td>
<td>0.72 (0.55 to 1.68)</td>
<td>P&lt;0.001</td>
<td>1.03 (0.60 to 1.77)</td>
<td>P=0.002*</td>
</tr>
<tr>
<td>1997-2002, age ≥1 year</td>
<td>0.19 (0.12 to 0.30)</td>
<td>P&lt;0.001</td>
<td>0.37 (0.25 to 0.53)</td>
<td>P&lt;0.001*</td>
</tr>
</tbody>
</table>

Sex:
- Male: 1
- Female: 0.84 (0.61 to 1.15) | P=0.3 | 1.06 (0.81 to 1.40) | P=0.7 |

Ethnicity:
- White: 1
- Black: 0.67 (0.54 to 1.27) | P=0.77 (0.55 to 1.08) |
- Other or not known: 1.42 (0.85 to 2.37) | P=0.1 | 1.09 (0.69 to 1.72) | P=0.1 |

How child was identified:
- Prospectively from birth: 1
- After birth: 0.49 (0.31 to 0.78) | P=0.001| 0.44 (0.30 to 0.64) | P=0.001 |

Place of birth:
- UK or Ireland: 1
- Abroad: 0.96 (0.62 to 1.48) | P=0.9 | 1.06 (0.73 to 1.53) | P=0.8 |

*Test of heterogeneity for age.

---

**Table 4** Change in % composition of initial three or four drug antiretroviral therapy in treatment of children with HIV before 1999 compared with 1999 onwards

<table>
<thead>
<tr>
<th>Antiretroviral drug combinations including:</th>
<th>Before 1999 (%)</th>
<th>1999 onwards (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Protease inhibitor</td>
<td>67</td>
<td>30</td>
</tr>
<tr>
<td>Non-nucleoside reverse transcriptase inhibitor</td>
<td>22</td>
<td>54</td>
</tr>
<tr>
<td>Both drug types</td>
<td>5</td>
<td>1</td>
</tr>
<tr>
<td>Triple nucleoside reverse transcriptase inhibitors</td>
<td>6</td>
<td>15</td>
</tr>
</tbody>
</table>

**Fig 2** Survival after initial AIDS diagnosis, by presenting indicator disease. Six cancer cases are excluded. Only one indicator disease is shown for children presenting with more than one according to hierarchy: P canini pneumonia, other opportunistic infection, HIV encephalopathy, failure to thrive, serious bacterial infections

**Fig 3** Proportion of time children with HIV spent having never received any drug treatment and while receiving none (previously treated), one, two, or three/four drugs, by calendar year
uptake of three or four drug antiretroviral therapy accompanied these changes and is likely to be the major contributing factor. Similar findings, concurrent with the widespread uptake of antiretroviral therapy, have been reported from cohorts of children in the United States and Italy. Follow up in these studies was only to 1999, but we observed that survival stabilised during 2000-2 after dramatic reductions during 1997-9. A similar pattern has been observed in adult cohorts.

There are clearly limitations in using cohort studies to ascribe changes in patterns of disease to particular interventions. As in the Italian study, we allowed for late entry at first presentation to reduce bias due to children with a good prognosis surviving longer and being more likely to be enrolled. In addition we repeated analyses separately for children born in the United Kingdom and Ireland or born abroad and observed similar results; this is particularly important in our cohort with the increasing proportion of children born abroad in recent years. As we included all children ever reported in the United Kingdom and Ireland, biases related to referral patterns to specialist centres are not an issue.

AIDS and hospital admissions
Our study is among the first to report reductions in hospital admissions over time in children with HIV, paralleling reductions in mortality and morbidity. The pattern of AIDS diagnoses before and since 1997 were similar, except for a trend towards a higher proportion of opportunistic infections in the later period. Improvements in the uptake of antenatal HIV testing and interventions to prevent transmission from mother to child have substantially increased the proportion of women diagnosed before delivery (from 32% in 1997 to 77% in 2001 in the United Kingdom) and led to a corresponding decrease in the proportion of infected infants. As expected, mortality, progression to AIDS, and hospital admission rates were lower in infected children born to mothers whose diagnosis was known before the child was born. Caution must be exercised in interpreting these results to suggest that antenatal testing has benefited beyond prevention of transmission by improving the outcome of infected children; presentation bias could play a part as asymptomatic infected children born to undiagnosed women will not be identified until they present with symptoms. Despite the marked reduction in hospital admission rates, total admissions have declined by only one quarter since 1996, underlying the need for continued inpatient as well as outpatient services for HIV infected children.

Infants compared with older children
Whereas in children aged over 1 year under clinical care mortality decreased by 81% and progression to AIDS or death decreased by 63%, we saw only modest improvements among infants. This may reflect a decrease in the proportion of infants in the cohort followed from birth, due to the reduction in mother to child transmission rates, although we attempted to control for this. Other explanations include the possibility that three or four drug antiretroviral therapy is not started early enough in infants. Pharmacokinetic issues may also hamper the effectiveness of antiretroviral therapy in this age group. In recent years, most infants presented with symptoms as they had been born to mothers undiagnosed at delivery. As reported here and previously, some infants die from P carinii pneumonia or cytomegalovirus disease, or both, before antiretroviral therapy can be started. The outlook for infants infected despite maternal diagnosis in pregnancy cannot easily be deduced from this study as numbers are small. However, we can speculate that transmission of maternal viruses that are drug resistant or more virulent may be also important. Ethnicity, sex, and place of birth had no significant effect on outcome.

Antiretroviral therapy
We observed a change over time from first use of antiretroviral therapy regimens based on protease inhibitors to those containing a non-nucleoside reverse transcriptase inhibitor. Although an increasing proportion of children receive four drug antiretroviral therapy, nearly all start with triple therapy. Further analyses of changes in prescribing practices for antiretroviral therapy and associated changes in HIV RNA and CD4 cell count responses are planned.

Conclusions
Rates of death, progression to AIDS, and hospital admission in children with HIV in the United Kingdom and Ireland, biases related to referral patterns to specialist centres are not an issue.

AIDS and hospital admissions
Our study is among the first to report reductions in hospital admissions over time in children with HIV, paralleling reductions in mortality and morbidity. The pattern of AIDS diagnoses before and since 1997 were similar, except for a trend towards a higher proportion of opportunistic infections in the later period. Improvements in the uptake of antenatal HIV testing and interventions to prevent transmission from mother to child have substantially increased the proportion of women diagnosed before delivery (from 32% in 1997 to 77% in 2001 in the United Kingdom) and led to a corresponding decrease in the proportion of infected infants. As expected, mortality, progression to AIDS, and hospital admission rates were lower in infected children born to mothers whose diagnosis was known before the child was born. Caution must be exercised in interpreting these results to suggest that antenatal testing has benefited beyond prevention of transmission by improving the outcome of infected children; presentation bias could play a part as asymptomatic infected children born to undiagnosed women will not be identified until they present with symptoms. Despite the marked reduction in hospital admission rates, total admissions have declined by only one quarter since 1996, underlying the need for continued inpatient as well as outpatient services for HIV infected children.

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Whereas in children aged over 1 year under clinical care mortality decreased by 81% and progression to AIDS or death decreased by 63%, we saw only modest improvements among infants. This may reflect a decrease in the proportion of infants in the cohort followed from birth, due to the reduction in mother to child transmission rates, although we attempted to control for this. Other explanations include the possibility that three or four drug antiretroviral therapy is not started early enough in infants. Pharmacokinetic issues may also hamper the effectiveness of antiretroviral therapy in this age group. In recent years,
Kingdom and Ireland have significantly fallen. As antenatal detection rates improve and fewer children born to infected women are themselves infected, children presenting to paediatric services with HIV are likely to be older and have to be born abroad. This, combined with improved life expectancy, means that the demand for specialist paediatric HIV services will continue to increase. \(^{21}\) Transition links with adult services are required to deal with the medical, social, and psychological needs of children entering adolescence and adult life.

National surveillance of paediatric HIV is undertaken by the National Study of HIV in Pregnancy and Childhood (NSHPC) at the Institute of Child Health, London, in collaboration with the Health Protection Agency Communicable Disease Surveillance Centre and the Scottish Centre for Infection and Environmental Health. NSHPC relies on active reporting from paediatricians through the British Paediatric Surveillance Unit of the Royal College of Paediatrics and Child Health, and from obstetric respondents reporting through an active reporting scheme run under the auspices of the Royal College of Obstetricians and Gynaecologists. We thank all those reporting to the NSHPC and particularly the staff, families, and children from the following 17 CHIPS centres: KB, E Hayes, M O’Mara, R Griffin (Our Lady’s Hospital for Sick Children, Dublin), AR (Heartlands Hospital, Birmingham), A Foot, H Kershaw (Bristol Royal Hospital for Sick Children, Bristol), W Tarnow-Mordi, J Petrie (Ninewells Hospital and Medical School, Dundee), K Sloper, V Shah (Ealing Hospital, London), J Mok (Royal Edinburgh Hospital for Sick Children), DMG, VN, K Klein, LF, M Clapson, B ÓMolana-Keen (Great Ormond St Hospital for Children NHS Trust, London), C Ball, D Navigam, D Graham, A Waters (King’s College Hospital, London), DMG, E Cooper, T Fisher, R Barrir, S Wong (Newham General Hospital, London), V Van Someren, K Moshal, S McKenna (Royal Free Hospital, London), M Sharland, S Donaghy, W Faulknall (St George’s Hospital, London), GT-E, H Lyall, S Walters, J White, S Head (St Mary’s Hospital, London), G Du Mont, R Cross (St Thomas’ Hospital, London), J Stremlau (University Hospital, Lewisham, London), A Riddell (John Radcliffe Hospital, Oxford), S Choo, R Lakshman, J Hobbs, F Shackleay (Children’s Hospital, Sheffield), and H Lyall, P Seery (Chelsea and Westminster Hospital, London).

Contributors: The CHIPS steering committee are all authors on this paper. Additional contributions from the MRC Clinical trials Unit are A Babiker, D Johnson, V Leclezio, B Glickstein, G O’Connor, A S Walker, G Wait, and Y Sovannmi. DMG, DTD, TD and MS conceived the idea for this paper. DMG and MS developed the protocol for the CHIPS cohort and secured additional funding in collaboration with CSP and PAT. PAT is responsible for the NSHPC at the Institute of Child Health, and JM manages data collection and entry. At the MRC Clinical Trials Unit, LF and K Doebelhaupt supervised data collection and data entry for CHIPS, assisted by S Masters, B Glickstein, G Wait, V Leclezio, G O’Connor, D Johnson and Y Sovannmi provided Oracle database support. TD conducted the statistical analyses under the supervision of DTD and input from A S Walker and A G Babiker. The paper was prepared by DMG and TD with PAT, CSP, A S Walker, and DTD. MS, GT-E, VN, KB, and AR ensured the quality of prospective data from their centres and commented on drafts of the paper. DMG is the guarantor for this study.

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Competing interests: None declared.

Ethical approval: The national study of HIV in pregnancy and childhood received renewed ethics approval, including follow up in the collaborative HIV paediatric study, in 2001 from Great Ormond Street Hospital for Children NHS Trust and the Institute of Child Health research ethics committee.

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the process is not fit for purpose. In particular there is confusion about whether revalidation is intended to detect poor performance, and if so, whether the process will suffice. Formative appraisal and summative revalidation are seen as uneasy bedfellows.

For most doctors the process will entail participation in an appraisal system, which must be aligned with the headings in Good Medical Practice and quality assured to the satisfaction of the General Medical Council. The Council states that such participation will be “a powerful indicator of a doctor’s current fitness to practise,” but makes no claim that the process will be sensitive (identify poor performance), specific (identify educational needs), valid (reflect actual clinical practice), or reliable (behave consistently across cohorts of doctors). Where there is any doubt doctors will be invited to submit more information and may be subjected to the performance procedures.

Conclusion

It is difficult to escape the conclusion that the purpose of revalidation is as a form of professional regulatory enforcer to ensure the NHS implements appraisal in a designated manner. This may be enough to encourage doctors to develop and to seek help early in case of difficulty. Alternatively, doctors relying for their revalidation on five appraisals might be tempted to set easily achievable objectives in their personal development plans, rather than risk failing to meet a challenge. The problems with this dual purpose have long been recognised. In Pringle’s view, the most likely outcome is the worst of all worlds, where the developmental and formative nature of appraisal is lost, and where revalidation fails to identify poor performers.

Whether patients and the government will be satisfied remains to be seen, particularly if (and predictably when) a recently revalidated doctor is found to have been a poor performer. Ultimately it may be more sensible to separate revalidation from appraisal. Doctors themselves could choose the independent route to revalidation, by submitting other evidence of minimal fitness to practise—appropriate tools are being developed. Or we could move wholesale to such a model (as in Canada), where a screening tool provides fitness to practise, rather than risk failing to meet a challenge. The problems with this dual purpose have long been recognised. In Pringle’s view, the most likely outcome is the worst of all worlds, where the developmental and formative nature of appraisal is lost, and where revalidation fails to identify poor performers.

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Patients and others might be more convinced by this. If doctors are found to be poor performers, the GMC’s “fitness to practise” criteria should be applied. These doctors are then investigated in more depth. In Pringle’s view, the most likely outcome is the worst of all worlds, where the developmental and formative nature of appraisal is lost, and where revalidation fails to identify poor performers.

Finally, the impact on doctors and patients should not be underestimated. In just over a decade, the NHS has moved from being an organisation based on high trust relationships to one where explicit written down standards, which are monitored, have become the norm for individuals and institutions. Revalidation is part of this increased bureaucratic control being applied to professional self regulation. It may increase apparent accountability, but may not foster a culture which increases patients’ trust and doctors’ professionalism.

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10 Nesbitt D. Purpose of revalidation process must be agreed on. BMJ 2001;322:558.
11 Wakeford R. GMC’s proposals for revalidation would not be accurate, economical, or fair. BMJ 2000;321:1220.

Corrections and clarifications

Decline in mortality, AIDS, and hospital admissions in perinatally HIV-1 infected children in the United Kingdom and Ireland

Mis-reading of the alignment in table 5 of this paper (table 2 in the abridged version) by D M Gibb and colleagues led us to print the hazard ratios for death and AIDS “death incorrectly for “How the child was identified” (BMJ 2003;327:1019-23). The hazard ratios for “prospectively from birth” and “after birth” should be inverted: the values for children identified after birth were therefore 1, and for children identified prospectively from birth 0.40 (95% confidence interval 0.31 to 0.78) and 0.44 (0.30 to 0.64).

ABC of smoking cessation: Nicotine replacement therapy

The table showing the formulations and availability of nicotine replacement products in this article by Andrew Molyneux contained an error (21 February, pp 134-6). Nicorette nasal spray is licensed as a Pharmacy (P) medicine and is available over the counter at pharmacies; it is not a prescription-only product, as stated in the table.

Effects of low dose ramipril on cardiovascular and renal outcomes in patients with type 2 diabetes and raised urinary albumin: randomised, double blind, placebo controlled trial (the DIABHYCAR study)

A wrong reference number persisted to publication of the full version of this article by Michel Marre and colleagues (28 February, pp 495-9). In the fifth paragraph of the Discussion, in the sentence starting “Conversely, only 30% of the diabetic patients . . .” the reference for the HOPE and MICRO-HOPE studies should be reference 9 (not 11, as stated). The references in the abridged version are correct.

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