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De novo malignancy after paediatric renal replacement therapy

H M Coutinho, J W Groothoff, M Offringa, M P Gruppen, H S A Heymans

Abstract

Aims—To determine frequency, type, determinants, and outcome of malignancies in children with end stage renal failure. Methods—All Dutch patients, aged less than 15 years, who started chronic renal replacement therapy between 1972 and 1992 and who were at least 18 years old on 1 January 1997, were retrospectively studied.

Results—Mean follow up from first renal replacement therapy was 15.5 years. Twenty two malignancies were found in 21 of 249 patients. Skin cancer accounted for 59% and non-Hodgkin lymphoma for 23% of malignancies. At 25 years after first renal replacement therapy, the probability of developing a malignancy was 17% (95% CI: 9 to 24%). Compared to the general population the incidence rate for overall cancer was tenfold higher. For non-melanoma skin cancer and non-Hodgkin lymphoma, standardised risks were 222 and 46 respectively. The use of more than 20 mg/kg cyclophosphamide showed an association with increased risk of malignancy. Six patients died as a result of their malignancy, accounting for 9.5% of overall mortality. Whereas four out of five patients with non-Hodgkin lymphoma died, the most frequent malignancy, skin cancer, did not contribute to mortality.

Conclusion—The long term risk of certain malignancies is significantly increased in children who have undergone renal replacement therapy. As an important contributor to overall mortality, awareness of this risk of malignancy in these patients is necessary, especially after treatment with cyclophosphamide.

Key words: renal replacement therapy; malignancy; follow up

The incidence of de novo malignancies in adult recipients of renal transplants is between six and seven times higher than in the general population.1 2 This is thought to be related to the use of immunosuppressive drugs for prevention of graft rejection. Mechanisms through which malignancies can arise include impaired immune surveillance, increased susceptibility to oncogenic viruses, and direct mutagenic effects of the drugs themselves.3–5 An increased risk of malignancy has also been described in adult dialysis patients: Inamoto et al found this group to have a 1.4-fold increased risk.4

In adult kidney recipients, post-transplant malignancy is known to be an important cause of morbidity and mortality, with cumulative incidences in long term follow up studies varying from 2.6% to 19.4%.1 2 3 7 Non-melanoma skin cancer is the most frequent malignancy in this group.1 2 3 7 8 11

In young adults with renal insufficiency since childhood, malignancy seems to be much less of a problem, with cumulative incidences varying from 0.8% to 3.9%.12–14 Lymphoma is the most frequent malignancy.12–16 However, current studies are either based on incomplete data registry or follow up time is relatively short. The long term risk of malignancy in young adults with renal insufficiency since childhood has not been evaluated sufficiently.

We present data from a Dutch national cohort study on Late Effects of Renal Insufficiency in Children (LERIC). The aim of this study, in which completeness of the cohort has been thoroughly assessed, is to determine frequency, type, determinants, and outcome of malignancies in this group of children.

Methods

STUDY POPULATION

The medical history of all Dutch patients, under 15 years of age, who started chronic renal replacement therapy (RRT) between 1972 and 1992 and who were at least 18 years old on 1 January 1997 was reviewed. Patients who needed RRT for more than four consecutive months were considered to be patients on chronic RRT. The cohort formation was based on data provided by the National Dutch Registry of patients on RRT (RENINE, Rotterdam, Netherlands). RENINE was founded in 1985. The completeness of data approaches 100% as central registration is obligatory for reimbursement of RRT. RENINE is the Dutch source of the European Dialysis and Transplantation Association (EDTA). Retrospective registration of patients with RRT starting before 1985 was checked by comparing RENINE data with the database of all Dutch centres for paediatric dialysis and kidney transplantation as well as the database of all centres for adult dialysis and transplantation. A list of all patients who fitted our age criteria was submitted to us by RENINE; it contained registry numbers, treatment modality, and the name of the last physician and hospital of treatment. All nephrologists in the Netherlands received a list of those cohort patients, who were under their treatment or had been so during the time of death. We contacted all physicians, asking them whether the list sent by RENINE was consistent with their registry base. Data were collected.

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Accepted 13 August 2001
Table 1  Patient characteristics of the LERIC cohort

<table>
<thead>
<tr>
<th></th>
<th>LERIC cohort</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of patients (male/female)</td>
<td>249 (136/113)</td>
</tr>
<tr>
<td>Total follow up time from start of RRT</td>
<td>3870 patient years</td>
</tr>
<tr>
<td>Mean follow up time from start of RRT, y (range)</td>
<td>15.5 (0.2–30.0)</td>
</tr>
<tr>
<td>Mean age at start of RRT, y (range)</td>
<td>10.6 (1.9–14.9)</td>
</tr>
<tr>
<td>Mean age at first transplantation, y (range)</td>
<td>11.5 (0.9–27.0)</td>
</tr>
<tr>
<td>Number of deaths</td>
<td>63</td>
</tr>
<tr>
<td>Number of patients never transplanted</td>
<td>18</td>
</tr>
<tr>
<td>Mean follow up time of patients never transplanted, y (range)</td>
<td>1.8 (0.3–9.8)</td>
</tr>
</tbody>
</table>

between November 1998 and July 2000. The study was approved by the Medical Ethical Committee of all participating centres.

DATA COLLECTION

To determine in which centres each patient had been treated, data collection was started in the centres of last treatment according to RE-NINE. From data found here, a list of all centres of treatment was made for each patient. For these centres, visits were arranged. If patients had been under treatment at two centres simultaneously, both centres were visited, unless all required data were found in one centre. If patients were found to have been treated in additional centres, these were added to the list and were visited as well.

A total of 37 hospitals in the Netherlands were visited by members of the LERIC team to collect and standardise medical histories of all patients included in the study. To achieve this, all available medical charts were reviewed. If charts had been lost or destroyed, data for a particular period could not be collected. These data were listed as missing, although important events such as death and occurrence of malignancy could be obtained from more recent medical information. Recorded data included date of birth, sex, kidney disease, date of first RRT, RRT history, use and type of immunosuppressive agents, total duration or cumulative dose of immunosuppressive agents irrespective of start of RRT, development of malignancy, date of diagnosis, type and outcome of malignancy, and occurrence and cause of death. Data were recorded from the first moment that renal disease led to hospitalisation until either patient death or last registered hospital visit up to July 2000. If uncertainties about malignancies remained, additional information was gathered by contacting the patient’s doctor. Malignancies prior to RRT and benign tumours were not included.

DATA ANALYSIS

Standardised risks were determined by calculating the expected number of malignancies based on the 1996 report of Incidence of Cancer in the Netherlands of the Netherlands Cancer Registry. A total of 3870 patient years was recorded, with data of 81 patient years missing. Mean follow up was 15.5 years per patient. For 82 patients (33%) follow up was more than 20 years. At the end of the study no patients were lost to follow up. Eighteen patients never underwent transplantation; mean follow up time from first RRT was 1.8 years and 17 patients died within a mean time of 1.4 years after start of RRT (range 0.3–4.0). Sixty seven patients were transplanted once, 118 twice, 38 three times, and eight four times. Treatment modality at the from the National Registry of Hospital Discharge Diagnosis (Landelijke Medische Registratie). After extensive checks for inconsistencies and duplicate records, the data are entered into the national database. All malignant and in situ malignancies are registered, except for basal cell carcinomas of the skin and carcinoma in situ of the cervix. As these malignancies are often removed without affirmation of the diagnosis by pathologist laboratories, registration of their incidence would not be reliable. Skin cancer is recorded per type per patient, unless new lesions arise more than three months after diagnosis. A malignancy first discovered at autopsy is also included in the cancer registry. For all comparisons with the LERIC cohort we used an age and gender matched group of the general population. Calculation of standardised risks was based on the incidence rates in the Netherlands between 1992 and 1996, whereas our study cohort consists of all Dutch patients, aged under 15 years, who started chronic RRT between 1972 and 1992 and were at least 18 years old on 1 January 1997. No national registry for cancer incidence existed in the Netherlands until 1989. We therefore assumed that the incidence of cancer in the Netherlands was that between 1992 and 1996, that incidence remained constant between 1972 and 1992, and that incidence was distributed in a homogeneous way in the 0–44 year age group. The standardised risk in the skin cancer group was calculated with the exclusion of basal cell carcinomas.

SPSS 9.0 and CIA (Confidence Interval Analysis, Prof. M J Gardner and British Medical Journal) were used for statistical calculation. For all factors marked with an asterisk in table 4, persons with the value zero were included in the reference group. Kaplan–Meier analysis was performed to determine cumulative cancer incidence. Cox regression analysis was performed to analyse potential risk factors. Time was defined as the amount of patient years at risk, with risk starting at first RRT (this is the moment one is considered a patient) and ending at diagnosis of malignancy, last recorded hospital visit, or patient death. If potentially oncogenic drugs such as cyclophosphamide were given before start of RRT, these were included.

Results

PATIENT CHARACTERISTICS

Table 1 shows patient characteristics of the whole of the LERIC cohort. From first RRT a total of 3870 patient years was recorded, with data of 81 patient years missing. Mean follow up was 15.5 years per patient. For 82 patients (33%) follow up was more than 20 years. At the end of the study no patients were lost to follow up. Eighteen patients never underwent transplantation; mean follow up time from first RRT was 1.8 years and 17 patients died within a mean time of 1.4 years after start of RRT (range 0.3–4.0). Sixty seven patients were transplanted once, 118 twice, 38 three times, and eight four times. Treatment modality at the
Confidence intervals were calculated with CIA (Confidence Interval Analysis, Prof. MJ Gardner and British Medical Journal group of the general population in the Netherlands between 1992 and 1996.

SR, standardised risk; NMSC, non-melanoma skin cancer; NHL, non-Hodgkin lymphoma.

†Basal cell carcinoma is excluded, because no national registration exists for this tumour.
*Years at risk defined as time between first renal replacement therapy and diagnosis of malignancy, last recorded hospital visit, or patient death.

Table 3  Standardised risk of developing a malignancy for the LERIC cohort

<table>
<thead>
<tr>
<th>Type</th>
<th>No. of tumours</th>
<th>Mean age at diagnosis of malignancy, y (range)</th>
<th>Interval between first transplantation and diagnosis, y (range)</th>
<th>Mean follow up time after diagnosis, y (range)</th>
<th>No. of patients who died due to malignancy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Skin cancer</td>
<td>13</td>
<td>27.8 (14.3–34.5)</td>
<td>15.0 (5.5–20.2)</td>
<td>3.6 (0.4–12.9)</td>
<td>0</td>
</tr>
<tr>
<td>SCC</td>
<td>7</td>
<td></td>
<td></td>
<td></td>
<td>0</td>
</tr>
<tr>
<td>BCC</td>
<td>3</td>
<td></td>
<td></td>
<td></td>
<td>0</td>
</tr>
<tr>
<td>Melanoma</td>
<td>2</td>
<td></td>
<td></td>
<td></td>
<td>0</td>
</tr>
<tr>
<td>M. Bowen</td>
<td>1</td>
<td></td>
<td></td>
<td></td>
<td>0</td>
</tr>
<tr>
<td>Non-Hodgkin lymphoma</td>
<td>5</td>
<td>23.5 (11.1–31.1)</td>
<td>11.7 (0.6–19.9)</td>
<td>0.6 (0.0–1.7)</td>
<td>4</td>
</tr>
<tr>
<td>ALL</td>
<td>1</td>
<td></td>
<td></td>
<td></td>
<td>1</td>
</tr>
<tr>
<td>Fibrosarcoma</td>
<td>1</td>
<td></td>
<td></td>
<td></td>
<td>1</td>
</tr>
<tr>
<td>Grawitz tumour</td>
<td>1</td>
<td></td>
<td></td>
<td></td>
<td>0</td>
</tr>
<tr>
<td>Leiomyosarcoma</td>
<td>1</td>
<td></td>
<td></td>
<td></td>
<td>0</td>
</tr>
<tr>
<td>Total</td>
<td>22</td>
<td>25.9 (11.1–34.5)</td>
<td>12.9 (0.1–21.1)</td>
<td>2.6 (0.0–12.9)</td>
<td>6</td>
</tr>
</tbody>
</table>

*Excluding one patient with malignant melanoma who never underwent transplantation.
SCC, squamous cell carcinoma; BCC, basal cell carcinoma; ALL, acute lymphatic leukaemia.


table 2  Characteristics of malignancies in the LERIC cohort

<table>
<thead>
<tr>
<th>Type</th>
<th>No. of patients</th>
<th>Overall cancer rate LERIC (per 1000 person years)</th>
<th>Overall cancer rate population (per 1000 person years)</th>
<th>NMSC rate LERIC (per 1000 person years)†</th>
<th>NMSC rate population (per 1000 person years)†</th>
<th>NHL rate LERIC (per 1000 person years)</th>
<th>NHL rate population (per 1000 person years)</th>
<th>SR NHL (95% CI)†</th>
</tr>
</thead>
<tbody>
<tr>
<td>Male</td>
<td>136</td>
<td>10 (6.3–15.1)</td>
<td>10 (0.6–14.9)</td>
<td>2.3</td>
<td>0.0087</td>
<td>1.3</td>
<td>0.028</td>
<td>(15–108)</td>
</tr>
<tr>
<td>Female</td>
<td>113</td>
<td>8 (3.6–14.9)</td>
<td>8 (3.6–14.9)</td>
<td>1.8</td>
<td>0.0087</td>
<td>0.59</td>
<td>0.018</td>
<td>(33–124)</td>
</tr>
</tbody>
</table>

Calculations of standardised risks are based on incidence rates per 100 000 person years for invasive tumours among males and females in the range 0–44 years according to the 1996 report of Incidence of Cancer in the Netherlands of the Netherlands Cancer Registry. The LERIC cohort was compared to an age matched group of the general population in the Netherlands between 1992 and 1996. Confidence intervals were calculated with CIA (Confidence Interval Analysis, Prof. MJ Gardner and British Medical Journal).

*Years at risk defined as time between first renal replacement therapy and diagnosis of malignancy, last recorded hospital visit, or patient death.
†Basal cell carcinoma is excluded, because no national registration exists for this tumour.
SR, standardised risk; NMSC, non-melanoma skin cancer; NHL, non-Hodgkin lymphoma.

start of RRT was haemodialysis in 81%, peritoneal dialysis in 16%, and transplantation in 3%.

INCIDENCE AND TYPE OF MALIGNANCY
Twenty two malignancies were found in 21 of the 249 patients (8.4% of those included in the study). Table 2 shows type, frequency, and age at diagnosis of malignancy, as well as interval between first transplantation and diagnosis of malignancy, follow up time recorded after diagnosis of malignancy, and death as a result of malignancy. Except for one patient with malignant melanoma, all patients with malignancies had been transplanted at least once. Thirteen patients had skin cancer (57% of all malignancies) and five had non-Hodgkin lymphoma (24%). The other malignancies found were one case of acute lymphatic leukaemia, one abdominal fibrosarcoma, one Grawitz tumour of the own kidney, and one leiomyosarcoma of the bladder. In the skin cancer group the ratio of basal cell carcinoma versus squamous cell carcinoma was 3:7. Some of the patients with skin cancer had lesions at multiple locations.

Mean age at diagnosis of malignancy was 25.9 years (SD 6.6, range 11.1–34.5), but skin cancer developed at a later mean age than non-Hodgkin lymphoma (at 27.8 and 23.5 years, respectively). The mean interval between first RRT and diagnosis of cancer was 13.8 years (SD 6.5, range 1.3–23.1). With 12 male and nine female patients, sex distribution in the malignancy group was comparable to the whole LERIC cohort as well as to the whole of the Netherlands in 1996 (51% male, 49% female).

STANDARDISED RISKS
Figure 1 shows the probability of developing a malignancy as a function of time since start of RRT (Kaplan–Meier analysis). At 25 years after first RRT the risk of developing a malignancy was 17% (95% CI: 9 to 24%). The increase in risk was most notable at 15 years after first RRT. Table 3 shows standardised risks of developing a malignancy. Compared to the general population the overall risk was ten-fold higher for patients in the LERIC cohort, with a larger risk for males than for females for overall cancer, non-melanoma skin cancer, and non-Hodgkin lymphomas. For non-melanoma skin cancer and non-Hodgkin lymphomas, standardised risks were 222 and 46 respectively. In the skin cancer group basal cell carcinoma was excluded, because no national registration exists for this tumour. This type of skin cancer is often removed without affirmation of diagnosis by pathologist laboratories. Therefore registration of its incidence would not be reliable.

DETERMINANTS
Table 4 shows variables analysed as potential risk factors. Only the use of more than 20 mg/kg cyclophosphamide was associated with an increased risk of malignancy. Standardised risk increased with increasing dosage. Even when the one patient with cancer of the bladder was excluded, malignancy rate increased with increasing dosage.

Table 3  Standardised risk of developing a malignancy for the LERIC cohort

<table>
<thead>
<tr>
<th>No. of patients</th>
<th>Years at risk*</th>
<th>Overall cancer rate LERIC (per 1000 person years)</th>
<th>Overall cancer rate population (per 1000 person years)</th>
<th>NMSC rate LERIC (per 1000 person years)†</th>
<th>NMSC rate population (per 1000 person years)†</th>
<th>NHL rate LERIC (per 1000 person years)</th>
<th>NHL rate population (per 1000 person years)</th>
<th>SR NHL (95% CI)†</th>
</tr>
</thead>
<tbody>
<tr>
<td>Overall</td>
<td>249</td>
<td>5.7</td>
<td>10 (6.3–15.1)</td>
<td>2.1</td>
<td>0.0093</td>
<td>1.3</td>
<td>0.028</td>
<td>(15–108)</td>
</tr>
<tr>
<td>Male</td>
<td>136</td>
<td>5.5</td>
<td>13 (6.9–23.3)</td>
<td>2.3</td>
<td>0.01</td>
<td>1.8</td>
<td>0.038</td>
<td>47</td>
</tr>
<tr>
<td>Female</td>
<td>113</td>
<td>5.3</td>
<td>8 (3.6–14.9)</td>
<td>1.8</td>
<td>0.0087</td>
<td>0.59</td>
<td>0.018</td>
<td>(33–124)</td>
</tr>
</tbody>
</table>
Table 4 Variables analysed as potential risk factors for development of malignancy in the LERIC cohort

<table>
<thead>
<tr>
<th>Variable (index versus reference)</th>
<th>Index group: cancer/ total exposed</th>
<th>Reference group: cancer/ total not exposed</th>
<th>Relative risk</th>
<th>95% confidence interval</th>
</tr>
</thead>
<tbody>
<tr>
<td>Male sex</td>
<td>12/136</td>
<td>9/113</td>
<td>1.04‡</td>
<td>0.44–2.47</td>
</tr>
<tr>
<td>Total transplantation time more than 1 year*</td>
<td>18/204</td>
<td>3/45</td>
<td>1.32‡</td>
<td>0.41–4.30</td>
</tr>
<tr>
<td>Total transplantation time more than 6 years*</td>
<td>16/183</td>
<td>5/66</td>
<td>1.15‡</td>
<td>0.44–3.03</td>
</tr>
<tr>
<td>Use of azathioprine</td>
<td>20/226</td>
<td>1/23</td>
<td>2.04‡</td>
<td>0.29–14.50</td>
</tr>
<tr>
<td>Use of azathioprine in skin cancer group</td>
<td>12/226</td>
<td>1/23</td>
<td>1.22‡</td>
<td>0.17–8.97</td>
</tr>
<tr>
<td>Use of cyclosporine</td>
<td>13/147</td>
<td>8/102</td>
<td>1.13‡</td>
<td>0.49–2.62</td>
</tr>
<tr>
<td>Use of azathioprine and cyclosporine</td>
<td>13/138</td>
<td>8/111</td>
<td>1.31‡</td>
<td>0.56–3.04</td>
</tr>
<tr>
<td>Use of methylprednisolone</td>
<td>12/157</td>
<td>9/92</td>
<td>0.78‡</td>
<td>0.34–1.78</td>
</tr>
<tr>
<td>Use of ATG</td>
<td>6/77</td>
<td>15/172</td>
<td>0.89‡</td>
<td>0.36–2.21</td>
</tr>
<tr>
<td>Use of OKT3</td>
<td>2/28</td>
<td>15/221</td>
<td>0.83‡</td>
<td>0.20–3.38</td>
</tr>
<tr>
<td>Use of cyclosporamide</td>
<td>6/37</td>
<td>15/212</td>
<td>2.92‡</td>
<td>0.95–9.53</td>
</tr>
<tr>
<td>Dose of cyclophosphamide (more than 20 mg/kg compared to less than 20 mg/kg or none)*</td>
<td>6/34</td>
<td>15/215</td>
<td>2.53‡</td>
<td>1.05–6.07</td>
</tr>
<tr>
<td>Dose of cyclophosphamide (more than 100 mg/kg compared to less than 100 mg/kg or none)*</td>
<td>5/17</td>
<td>16/232</td>
<td>4.26‡</td>
<td>1.78–10.20</td>
</tr>
</tbody>
</table>

* Patients with the value 0 were included in the reference group.
‡Relative risks and 95% confidence intervals were calculated with the Cox regression method. Time was defined as the amount of patient years at risk with risk starting at first renal replacement therapy (this is the moment one is considered a patient) and ending at diagnosis of malignancy, last recorded hospital visit, or patient death; a total of 3870 patient years was recorded according to this definition.
§Relative risks were calculated with CIA (Confidence Interval Analysis, Prof. MJ Gardner and British Medical Journal).

Discussion

SUMMARY OF RESULTS

This long term follow up study of late effects of renal insufficiency in children shows a probability of developing malignancies of 17% at 25 years after first RRT, the majority being skin cancers and non-Hodgkin lymphomas. Compared to the general population the risk of developing a malignancy for our patients was tenfold higher. For non-melanoma skin cancer the standardised risk was 222; for non-Hodgkin lymphoma it was 46. Cumulative cyclophosphamide dose >20 mg/kg was associated with an increased risk of malignancy. Finally, malignancies contributed to a significant proportion of overall mortality (9.5%), although the most frequent malignancy, skin cancer, resulted in no deaths.

INCIDENCE AND TYPE OF MALIGNANCY

In a study in paediatric renal transplant recipients comparable to the LERIC study, Offner and colleagues found an incidence of malignancy of 2.7% after a mean of 13.1 years follow up. With a slightly higher mean follow up of 16.6 years we found a fourfold higher incidence. This difference could be explained by a more complete search strategy in the LERIC study; Offner et al contacted patients by telephone to enquire about medical events instead of reviewing medical charts, as was done in the LERIC study. This might have led to a higher incidence.

Figure 1 Probability of cancer as a function of time since start of RRT.
to an underreporting of skin cancer by Offner et al. Furthermore, we found that risk of malignancy strikingly increased at 15 years after start of RRT. The difference in mean follow up, although small, could therefore also have contributed to the lower incidence found by Offner et al.

Similar types of malignancy were found in the LERIC study as in adult studies of renal transplant recipients.\textsuperscript{10-18-20}

The ratio of basal cell carcinoma versus squamous cell carcinoma was 3:7 and thus reversed compared to the general population. In adult studies of renal transplant recipients similar findings have been reported,\textsuperscript{2} but none of these studies have reported an explanation. It has been postulated that squamous cell carcinomas are more susceptible to decreased cellular immunity caused by immunosuppressive agents than basal cell carcinomas. Furthermore, infection with human papillomavirus (HPV), frequently seen in immunosuppressed patients, might increase chances of developing squamous cell carcinomas rather than basal cell carcinomas (personal communication). However, this still has to be investigated.

Non-Hodgkin lymphoma occurred at a younger mean age than skin cancer and had the potential to develop sooner after first transplantation (after a minimum of 0.62 years compared to 5.48 years in skin cancer). Both Opelz and Henderson\textsuperscript{22} and Penn\textsuperscript{11} described this phenomenon in adult studies of renal transplant recipients. Penn\textsuperscript{11} suggested that non-Hodgkin lymphomas develop during the first few months after transplantation because of a high degree of immunosuppression following the use of multiple immunosuppressive agents. Epstein–Barr virus (EBV) plays an important role in development of non-Hodgkin lymphomas.\textsuperscript{11,14-16} EBV infection of B lymphocytes can lead to uncontrolled cell growth and proliferation.\textsuperscript{15} Unfortunately we have no data on exposure of our patients to this oncogenic virus. Gaya et al found that the risk of developing skin cancer increased with time after transplantation.\textsuperscript{1} Impairment of normal DNA repair mechanisms as a result of immunosuppressive drugs, combined with cumulative UV light induced damage may be responsible for the apparent latency period of several years before skin cancer develops. Skin cancer education of patients with RRT, for example, avoidance of excessive sunlight exposure, might lead to a longer latency period or even prevent the development of skin cancer.

**STANDARDISED RISKS**

We found a standardised risk (SR) of 10 for all malignancies compared to the general population. For non-melanoma skin cancer and non-Hodgkin lymphoma, however, standardised risks were 222 and 46. Not many studies have compared cancer incidence among RRT recipients to that of the general population. Opelz and Henderson\textsuperscript{22} found an SR of 37 for non-Hodgkin lymphoma in renal transplant recipients in the first post-transplant year. Gaya et al reported an SR of 6 for overall cancer, 19 for skin cancer, and 45 for non-Hodgkin lymphoma in renal transplant recipients after an average follow up of 9.6 years.\textsuperscript{1} The tenfold higher SR that we found for skin cancer compared to Gaya et al may be related to our longer follow up time.

Standardised risks were based on the incidence rates of malignancies in the Netherlands between 1992 and 1996, whereas malignancies in the LERIC cohort had developed before 1992. Considering the fact that overall and skin cancer incidences in the Netherlands in persons aged 0–44 years have increased since 1989,\textsuperscript{17} the standardised risk of the LERIC cohort could be underestimated.

**DETERMINANTS**

The crude association of cumulative cyclophosphamide dose of >20 mg/kg with increased risk of development of malignancies suggests a potential carcinogenic effect. This analysis was based on small numbers. Warranted use of this immunosuppressive agent is warranted. Development of malignancies secondary to the use of cyclophosphamide has been described: cancer of the bladder as well as acute myeloid leukaemia.\textsuperscript{22-24} The immunosuppressive agent cyclophosphamide can lead to an increased risk of uncontrolled cell growth and proliferation, and invasion or reactivation of oncogenic viruses such as EBV.\textsuperscript{23} Furthermore, acrolein, the major toxic metabolite of cyclophosphamide, has been reported to induce bladder cancer in rats.\textsuperscript{27} Travis et al reported that cyclophosphamide related bladder cancer was dose dependent.\textsuperscript{28} We found evidence of an association between the risk of overall malignancies and dose of cyclophosphamide. Although we found only one case of bladder cancer, analysis of cyclophosphamide as a risk factor with exclusion of this case of bladder cancer still showed an increased risk of malignancy for cumulative doses of 100–200 mg/kg.

**OUTCOME**

Six of 21 patients died as a result of their malignancy, none of which were a result of skin cancer. Although this is the most frequent malignancy associated with renal transplantation, it therefore seems to be mainly contributing to morbidity, but not mortality. However, as the mean follow up time after diagnosis of skin cancer was only 3.6 years, no definitive conclusions can be drawn. On the other hand, as in the general population, mortality caused by non-melanoma skin cancer is extremely low,\textsuperscript{17} the same might be expected for the LERIC cohort as well. Non-Hodgkin lymphomas did contribute in a significant number to both morbidity and mortality: four of five patients died in our cohort. They also caused death more rapidly than other malignancies in our cohort. Cessation or dose reduction of immunosuppressive agents has been reported to lead to tumour regression in non-Hodgkin lymphomas.\textsuperscript{1,26} This is something clinicians need to be aware of in order to try to reduce mortality caused by this type of cancer.
De novo malignancy after paediatric renal replacement therapy

One Grawitz tumour was diagnosed by coincidence in a patient who is currently alive. As this type of cancer leads to death in nearly every case, mortality in the LERIC cohort could easily have been higher.

Malignancies in immunocompromised individuals and possibly also in chronic dialysis patients appear to be more aggressive and more difficult to treat, hence both morbidity and mortality are prone to be higher compared to malignancies in the general population.1-3 The mortality/incidence ratio we found for overall cancer and non-Hodgkin lymphomas supports these findings.

LIMITATIONS OF THE STUDY

Inherent to studies of relatively rare paediatric diseases, our sample size was fairly small. Furthermore, the study design was retrospective and data were collected from multiple centres. We nevertheless believe this long term study provides as complete a picture as possible on frequency, type, determinants, and outcome of malignancies after paediatric RRT. No patients were lost to follow up, and data on important events such as occurrence of malignancy were not missing.

CONCLUSION

In conclusion, the long term risk of certain malignancies is significantly increased in children with RRT. Although the most frequent malignancy, skin cancer, resulted in no deaths, the fact that malignancy related death represented 9.5% of overall mortality warrants further research on this subject.

In patients treated with more than 20 mg/kg of cyclophosphamide extra vigilance towards the development of malignancies is required. Regular medical check ups, such as routine control by dermatologists and skin cancer education of patients, could decrease both morbidity and mortality caused by malignancy as a late effect of paediatric RRT.

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