Development and progression of microalbuminurina in a clinic sample of patients with insulin dependent diabetes mellitus

C A Jones, G P Leese, S Kerr, K Bestwick, D I Isherwood, J P Vora, D A Hughes, C Smith

Abstract
The evolution of abnormal albumin excretion and its association with suggested risk factors were studied in 233 children with insulin dependent diabetes mellitus (IDDM) attending a single paediatric diabetic clinic over an eight year period. Yearly albumin:creatinine ratios (ACR; measured in mg/mmol) in early morning urine samples, glycated haemoglobin (HbA1c), and blood pressure were recorded. Thirty four (14.5%) children had a persistently raised ACR (ACR > 2.5 mg/mmol on at least three consecutive occasions) and 21 (9%) had an intermittently raised ACR (ACR > 2.5 mg/mmol on at least two occasions). Factors associated with a persistently raised ACR compared with normal albuminuria in IDDM included longer duration of diabetes, raised median HbA1c during the first five years after diagnosis, and final age adjusted systolic and diastolic blood pressure represented as standard deviation scores. The onset of persistently raised ACR in 13 of 34 children was before puberty and in 23 of 34 children it was within the first four years of diagnosis. The cross sectional prevalence of raised ACR was 12.9% at one year, 18.3% at five years, and 33% at 10 years after diagnosis. Raised ACR occurs frequently before puberty and in the early stages of childhood diabetes. (Arch Dis Child 1998;78:518–523)

Keywords: microalbuminuria; insulin dependent diabetes mellitus; screening

Studies in adults with insulin dependent diabetes mellitus (IDDM) have shown that persistent microalbuminuria (albumin excretion rate (AER) 20–200 µg/minute), identified by screening, predicts the development of overt nephropathy,1–3 which can progress to end stage renal failure requiring dialysis and transplantation.4 With increasing albumin excretion there is also an increase in other microvascular complications including proliferative retinopathy.5 6 Estimates of the point prevalence of microalbuminuria in childhood vary between 7% and 20%.7–10 This variation reflects differences in definitions of microalbuminuria and mean age of study populations. Furthermore, these were not longitudinal studies and cannot comment on the cumulative prevalence of persistent microalbuminuria. In a single longitudinal study, following 156 children with IDDM over 15 years, 25% of children below the age of 21 years developed microalbuminuria within 14 years of diagnosis.11

Reported risk factors for the development of diabetic renal disease include a longer duration of IDDM, an earlier age at diagnosis, onset of puberty, poorer glycaemic control during the first five years of diabetes, smoking, and a family history of diabetic nephropathy.11–18 The role of a family history of hypertension as a risk factor remains controversial.19 20

We report a longitudinal evaluation of urinary albumin excretion in children with IDDM, attending a single clinic, over a period of eight years. We assessed associations between raised urinary albumin excretion and age at disease onset, duration of diabetes, pubertal status, glycaemic control, and blood pressure. An understanding of the development and progression of increased albumin excretion in childhood IDDM can guide the development of screening practices, and might have implications for these patients during adulthood.

Methods
PATIENTS
All children and adolescents still attending the diabetic clinic at Alder Hey Children's Hospital, Liverpool between January 1993 and June 1994 were included in the study. All children with IDDM in the Liverpool area are seen in this clinic and are managed by one endocrinologist (CS). Information was collected on this unselected population from January 1986 to June 1994 by case record review.

There were 233 children of whom 114 (49%) were boys. The median duration of IDDM was 3.9 years, (interquartile range (IQR), 1.8–6.7) and the median age at onset of diabetes was 7.7 years, (IQR, 4.1–11.2).

MEASUREMENTS
Urinary albumin:creatinine ratio (ACR), blood pressure, glycated haemoglobin, and pubertal status were recorded as part of the clinical assessment at each appointment and the annual measurements were analysed. An early morning urine sample was collected as part of each routine clinical assessment. Urinary albumin was measured by rate nephelometry (Beckman Instruments UK Ltd, High Wycombe, Bucks, UK). Urinary creatinine was measured using the modified Jaffe reaction. Results are expressed as the ratio of albumin to creatinine (mg/mmol). The overall coefficient of variation for ACR is 2.03%.
Urine microscopy and bacteriological culture were performed on all samples with detectable proteinuria on dipstick, and the ACR result was discarded if the culture was positive. Urine samples were not collected from pubertal girls during menses and any sample with haematuria was excluded from the results.

ACR was used as a measure of urinary albumin excretion and was considered raised if it was ≥ 2.5 mg/mmol. Persistently raised urinary albumin excretion is defined as an ACR ≥ 2.5 mg/mmol on at least three consecutive occasions. Children with diabetes for only two years with an ACR > 2.5 mg/mmol at onset, one year and two years duration of diabetes were also classified as having a persistently raised ACR.

Intermittently raised urinary albumin excretion is defined as an ACR ≥ 2.5 mg/mmol on at least two occasions. Children were considered to have normal albumin excretion if their ACR remained < 2.5 mg/mmol.\(^1\)

Glycated haemoglobin was measured from January 1991 as haemoglobin A1c (HbA1c) using column chromatography. (Bio-Rad Haemoglobin A1c Mini Column Test; Biorad Laboratories Ltd, Hemel Hampstead, Hertfordshire, UK). The normal range for HbA1c in our laboratory in a non-diabetic population is 3.4–6.4%. The imprecision measured as the coefficient of variation was 15.8% at an HbA1c of 4.6% and 4.1% at an HbA1c of 10.2%.

Blood pressure was measured manually by mercury sphygmomanometer with an appropriate paediatric cuff. Diastolic blood pressure was defined as Korotkoff level IV. To correct for age and sex, measurements are expressed as a standard deviation score related to the 1987 Task Force data for blood pressure in normal children.\(^2\) Pubertal status was measured by Tanner staging.

### Results

Throughout the study period, 172 (73.8%) children had normal albumin excretion. Thirty-four (14.6%) children had a persistently raised ACR and 27 (11.6%) children had an intermittently raised ACR.

Figure 1 shows the percentage of children with an ACR < 2.5 mg/mmol at a particular duration of diabetes from which the point prevalence of a raised ACR can be calculated. At the onset of the study, 64 patients already had established IDDM, so that ACR from the onset of their disease is not available. However, clinical details from their subsequent course during the period studied is included, thus extending data available on IDDM to 10 years duration.

There was an increase in the percentage of children who had an ACR > 2.5 mg/mmol at one year following diagnosis (12.9%), at five years after diagnosis (18.3%), and at 10 years (33%) after diagnosis, indicating a progressive increase in the prevalence of a raised ACR with duration of diabetes in our population.

In 24 of the 34 children with a persistently raised ACR, it remained stable throughout the period studied. However, in 10 children it increased progressively. In eight of these, the rise in ACR, over a minimum period of three years, varied from 18% to 78% each year from onset of a persistently raised ACR. In the other two children with IDDM for only two years, ACR increased in one year by 74% and 20–72). Seven patients with a persistently raised ACR had episodes of ACR > 45.5 mg/mmol, indicating macroalbuminuria.\(^3\) One patient had an ACR that was persistently > 45.5 mg/mmol throughout the study period.

The onset of a persistently raised ACR occurred in 13 children before puberty.

A comparison of clinical features in children with normal albumin excretion and those with persistently raised ACR is shown in table 1. There is no significant difference between the two groups in age at diagnosis. There is a significant difference between the two groups with respect to the age of the children at the end of the study (p < 0.001). However, the older age in children with persistently raised ACR at completion of the study reflects the longer duration of diabetes in this group (7.2 vs 3.3 years; p = 0.002). There is a significantly greater proportion of girls with persistently raised ACR (p = 0.0014). The median HbA1c in the first five years after diagnosis was

![Figure 1 Prevalence of children with an albumin:creatinine ratio (ACR) < 2.5 mg/mmol. Numbers at top are numbers of children.](http://adc.bmj.com/content/78/6/518)

Table 1 Median and IQR of children with persistently raised ACR and normal ACR

<table>
<thead>
<tr>
<th></th>
<th>Raised ACR (n = 34)</th>
<th>Normal ACR (n = 172)</th>
<th>p value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age at onset of diabetes (years)</td>
<td>8.9 (4.7, 11.5)</td>
<td>7.1 (3.7, 11.1)</td>
<td>NS</td>
</tr>
<tr>
<td>Age at end of study (years)</td>
<td>16.4 (13.4, 17.9)</td>
<td>12.3 (9.4, 15.3)</td>
<td>0.0001</td>
</tr>
<tr>
<td>Duration of diabetes (years)</td>
<td>7.2 (4.1, 10)</td>
<td>3.3 (2.8, 6.7)</td>
<td>0.002</td>
</tr>
<tr>
<td>HbA1c (%) over first 5 years</td>
<td>9.2 (8.0, 10.2)</td>
<td>8.6 (7.5, 9.6)</td>
<td>0.033</td>
</tr>
<tr>
<td>Sex (M/F)</td>
<td>9/25</td>
<td>96/76</td>
<td>0.0014</td>
</tr>
</tbody>
</table>

IQR, interquartile range; ACR, albumin:creatinine ratio; HbA1c, glycated haemoglobin.

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\(^1\) Children with diabetes for only two years with an ACR > 2.5 mg/mmol at onset, one year and two years duration of diabetes were also classified as having a persistently raised ACR.

\(^2\) Pubertal status was measured by Tanner staging.

\(^3\) Glycated haemoglobin was measured from January 1991 as haemoglobin A1c (HbA1c) using column chromatography. (Bio-Rad Haemoglobin A1c Mini Column Test; Biorad Laboratories Ltd, Hemel Hampstead, Hertfordshire, UK). The normal range for HbA1c in our laboratory in a non-diabetic population is 3.4–6.4%. The imprecision measured as the coefficient of variation was 15.8% at an HbA1c of 4.6% and 4.1% at an HbA1c of 10.2%.

\(^4\) Blood pressure was measured manually by mercury sphygmomanometer with an appropriate paediatric cuff. Diastolic blood pressure was defined as Korotkoff level IV. To correct for age and sex, measurements are expressed as a standard deviation score related to the 1987 Task Force data for blood pressure in normal children.
Table 2  Median and IQR for children with intermittently raised ACR and normal ACR

<table>
<thead>
<tr>
<th></th>
<th>Intermittent ACR (n = 27)</th>
<th>Normal ACR (n = 172)</th>
<th>p value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age at onset of diabetes (years)</td>
<td>10 (7.4, 11.3)</td>
<td>7.1 (3.7, 11.1)</td>
<td>NS</td>
</tr>
<tr>
<td>Age at end of study (years)</td>
<td>15.1 (11.6, 18.8)</td>
<td>12.3 (9.4, 15.3)</td>
<td>&lt; 0.05</td>
</tr>
<tr>
<td>Duration of diabetes (years)</td>
<td>3.8 (1.6, 8.5)</td>
<td>3.3 (1.8, 6.7)</td>
<td>NS</td>
</tr>
<tr>
<td>HbA1c (%) over first 5 years</td>
<td>9.7 (8.2, 10.3)</td>
<td>8.6 (7.5, 9.6)</td>
<td>&lt; 0.01</td>
</tr>
<tr>
<td>Sex (M/F)</td>
<td>9/18</td>
<td>96/76</td>
<td>&lt; 0.05</td>
</tr>
</tbody>
</table>

IQR, interquartile range; ACR, albumin:creatinine ratio; HbA1c, glycated haemoglobin.

Discussion

Persistent microalbuminuria with an AER of 20–200 µg/minute is predictive of nephropathy in adults with IDDM,1–4 and has led to the development of screening programmes and intervention studies.24–27 There is limited information on the evolution of urinary albumin excretion from the onset of IDDM in a childhood population. Most studies have been cross sectional in design, reporting a point prevalence of 7–20% for microalbuminuria, and by their nature unable to evaluate the prevalence of persistent microalbuminuria.7–10

A single longitudinal study of 156 children reported a cumulative prevalence of 25% for microalbuminuria within 14 years of diagnosis of IDDM.10 Variability in the estimates of prevalence can also be affected by differences in measurements used to define microalbuminuria.

The definition of microalbuminuria according to timed AER has been useful in describing accurately the natural history of renal dysfunction in adult IDDM groups. However, in a routine clinic setting and particularly with a paediatric population, the acceptability, practicality, and reliability of regular 24 hour or timed overnight urine collections is reduced. This is especially true if the child is enuretic or where poor compliance is expected. The ACR of early morning urine samples is useful as a surrogate measure for AER, with variation in the sensitivity and specificity according to the defining level of ACR chosen.28–30 An ACR ≥ 2.5 mg/mmol in children had both a sensitivity and specificity of 94% with a positive predictive value of 66% for microalbuminuria using established definitions based on timed collections.31

In selecting a defining value, account should also be taken of the lower urinary albumin excretion seen in normal children compared with normal adults and the increase seen with age.32–34 Using the ACR of early morning urine samples, an upper limit of 1.17 mg/mmol was found in a population of normal British children.35 Using an ACR ≥ 2.5 mg/mmol as the defining value for an increased albumin excretion, and using established definitions for persistence and progression of microalbuminuria,11–21 we describe our observations in a population of 233 children with IDDM followed longitudinally over an eight year period.

A cumulative prevalence of 14.5% with a persistently raised ACR over 8.5 years compares with 11.5% at 7.5 years in the only longitudinal study of urinary albumin excretion in...
Microalbuminuria in a clinic sample of patients with IDDM

within 14 years of diagnosis, one quarter of this group developed microalbuminuria. It is suggested that the progression of microalbuminuria in childhood is low. In 10 children with a persistently raised ACR, we report a progressive increase in their albumin excretion. Adult studies indicate that these patients are at greater risk of overt nephropathy. Seven patients with a persistently raised ACR had, on occasions, an ACR above 45.5 mg/mmol and one patient had an ACR that was persistently greater than 45.5 mg/mmol. This value indicates nephropathy and corresponds to a urinary albumin excretion equivalent to 300 µg/minute, which is albustix positive. Adult patients with diabetic nephropathy have a progressive decline in glomerular filtration rate and ∼ 50% of untreated patients die within seven years from renal failure and cardiovascular disease. The prognosis of patients developing albustix positive proteinuria in childhood is not yet known.

Previous studies have indicated that the onset of microalbuminuria before puberty occurs only rarely and, consequently, screening for microalbuminuria should be recommended for children over 12 years of age. In a longitudinal study, three of 156 children developed persistent microalbuminuria at less than 12 years of age. However, pubertal staging of the children in that study was not discussed. In a further study, in which puberty was defined by age, the duration of diabetes before puberty was found to have little effect on microvascular disease. We have found that in 15 of 34 (44%) children with persistently raised ACR, onset was before puberty, as assessed by Tanner staging. This suggests that children with IDDM are at risk of developing increased albumin excretion before puberty and this would warrant regular screening. Whether these children carry the same risk for the progression of raised albumin excretion to the stage of overt nephropathy will require continued follow up of this group.

The association of increasing urinary albumin excretion in IDDM in childhood and the duration of disease has been demonstrated in some studies, but not in others. We found a clear increase in the cumulative prevalence of affected children with a raised ACR, and in those children with persistently raised ACR, a duration of IDDM of 7.2 years compared with only 3.3 years for normoalbuminuric children with diabetes. It is possible that further follow up might identify children in the normoalbuminuric group who later develop a raised ACR. Although there is increasing urinary albumin excretion with increasing disease duration, we also found that in 56% of children who developed a persistently raised ACR, its onset was within the first four years of their IDDM. These children represent 8% of the total study population, a figure equal to the percentage with early onset of persistent microalbuminuria in the other longitudinal study. If screening is to be undertaken, it would seem appropriate to introduce it from the early stages of disease onset.

**Key messages**

- The cumulative prevalence of persistently raised albumin:creatinine ratios over 8.5 years was 14.5%.
- The onset of persistent microalbuminuria may occur before puberty and during the first four years after diagnosis.
- Persistent microalbuminuria was associated with increasing duration of diabetes, poorer glycaemic control, and increased blood pressure.
- We recommend the routine monitoring of albumin:creatinine ratios in early morning urine in all children with insulin dependent diabetes.

The association between a persistently raised ACR and being a girl might reflect the finding of increased urinary albumin excretion in normal girls at all ages from 4 to 16 years. There was no significant difference in the duration of IDDM for both sexes. Although the girls had more episodes of diabetic ketoacidosis (data not shown), there was no difference in median HbA1c between girls and boys.

We confirm the association of persistently raised urinary albumin excretion with poorer glycaemic control early in disease. Improving glycaemic control might reduce the incidence of microalbuminuria in early IDDM. The youngest patient in the diabetes control and complications trial was only 13 years. The risks of hypoglycaemia are increased with improved glycaemic control, which may be unacceptable in children. The intensive blood glucose monitoring and increased frequency of insulin injections are also impractical. The poorer early metabolic control in our group may explain the increase in prevalence of persistently raised urinary albumin excretion compared with the other longitudinal study (HbA1c values: mean (SD), 9.2% (1.4%) v 8.4% (1.3%), respectively).

An increase in blood pressure is associated with increases in urinary albumin excretion in both adults and children. There is controversy as to whether an increase in blood pressure precedes or is a result of the development of microalbuminuria. Using routine clinic blood pressure measurements, we have found a significant increase in the SDS systolic blood pressure from onset to final blood pressure measurement for children with diabetes who had a persistently or intermittently raised ACR. However, diastolic blood pressure SDS was only raised significantly in children with a persistently raised ACR. In a comparison of children with and without raised ACR, at final measurement, the blood pressure SDS was significantly higher in the persistently raised ACR group for both systolic and diastolic measurements compared with the normoalbuminuric children with diabetes. There was no difference in blood pressure SDS between children with persistently raised ACR and normal albumin excretion at four years duration of diabetes. This time point represents the
median time before the onset of persistent microalbuminuria. This suggests that the increase in blood pressure SDS observed in the children with a persistently raised ACR occurs in parallel with, or as a consequence of, microalbuminuria. This should be considered in decisions surrounding the use of antihypertensive agents in children with microalbuminuria who are still normotensive. Angiotensin converting enzyme (ACE) inhibitors reduce urinary albumin excretion in hypertensive and normotensive adults with diabetes and microalbuminuria, and reduce the incidence of nephropathy in normotensive patients with microalbuminuria. However, there are few data on the use of ACE inhibitors in childhood IDDM, although they have been shown to halt or reverse the progression of microalbuminuria in a small group of normotensive children with microalbuminuria.

The clinical significance of intermittent microalbuminuria in childhood IDDM is unknown. Adults with diabetes who develop persistent microalbuminuria have an increased incidence of intermittent microalbuminuria before the onset of persistent microalbuminuria, and those with a greater number of episodes of intermittent microalbuminuria are more likely to develop persistent microalbuminuria. Twenty-seven children in our population had an intermittently raised ACR. They were similar to the children with persistently raised ACR in that they were older at the end of the study period and had a higher HbA1c compared with the children with normal albumin excretion. They also had a significantly higher SDS for systolic and diastolic blood pressure at the end of the study period compared with children with normal albumin excretion.

We have shown that in an unselected population of children with diabetes, 14.5% developed persistently raised ACR, and of these, 10 increased progressively. Increasing duration of diabetes, poorer glycaemic control, and an increase in blood pressure were associated with this. The routine monitoring of children with IDDM using ACR on an early morning urine sample is a simple and reliable technique in clinical practice. It will enable the identification of groups at high risk of diabetic nephropathy. Monitoring should be extended to those in the early stages of the disease and those who are still prepubertal. Continued follow-up of children with a stable but persistently raised ACR and those with an intermittently raised ACR will clarify whether they will follow a benign course or will progress ultimately to nephropathy. Diabetic children with a progressively increasing ACR should be considered for intervention studies.
Coat of Arms of the Royal College of Paediatrics and Child Health

Archives of Disease in Childhood was first published in 1926. The Journal is owned jointly by the BMJ Publishing Group and the Royal College of Paediatrics and Child Health (formerly the British Paediatric Association). The College was founded in 1996 and the coat of arms of the College has been included on the cover of the Journal since January 1998. An explanation of the symbols in the coat of arms is given below.

The family

RIGHT SUPPORTER (ON THE RIGHT OF THE PAGE)
Father figure of Thomas Phaire who was the author of the first book on paediatrics in English (1545) in contemporary, academic dress. He is holding scales which signify the role of the College in setting standards and professional examinations.

LEFT SUPPORTER
Mother figure with red hair in modern academic dress resembling Baroness Lloyd of Highbury who was the first woman President of the British Paediatric Association. She is holding a rod with a double stranded helix indicating the importance of science and, in particular, genetics, to advances in child health. The device also refers back to the serpent-entwined staff of Aesculapius.

CREST
A child to represent the concept that the aspiration of parents and paediatricians is that the child should reach a level of attainment which is higher than their own. The child has been redrawn from the Coat of Arms of the Foundling Hospital in Coram Fields to show an association with children’s hospitals.

The College

SHIELD
Oak tree on green meadow to represent child development and health. A book to indicate education and scholarship. Two hands shaking to represent friendship, which was one of the first rules of the British Paediatric Association.

BASE
Meadow in the shape of a globe to signify the responsibility of the College towards children throughout the world. At the inaugural meeting of the British Paediatric Association, representatives of different regions of the British Isles were appointed and the College is still constituted in this democratic way, as shown emblematically by the rose, the thistle, the shamrock and the daffodil. Maple leaves commemorate Donald Patterson who was the driving force for the inauguration of the British Paediatric Association and served as a leader for the first 20 years.

Motto
From Psalm 127, Hereditas Domini filii. Children are a heritage from the Lord.