
ABSTRACT

Objectives To describe the characteristics of children and adults with incident type 1 diabetes in contemporary, multiethnic UK, focusing on differences between the islet autoantibody negative and positive.

Design Observational cohort study.

Setting 146 mainly secondary care centres across England and Wales.

Participants 3312 people aged ≥5 years were recruited within 6 months of a clinical diagnosis of type 1 diabetes via the National Institute for Health Research Clinical Research Network. 3021 were of white European ethnicity and 291 (9%) were non-white. There was a small male predominance (57%). Young people <17 years comprised 59%.

Main outcome measures Autoantibody status and characteristics at presentation.

Results The majority presented with classical osmotic symptoms, weight loss and fatigue. Ketoacidosis was common (42%), especially in adults, and irrespective of ethnicity. 35% were overweight or obese. Of the 1778 participants who donated a blood sample, 85% were positive for one or more autoantibodies against glutamate decarboxylase, islet antigen-2 and zinc transporter 8. Presenting symptoms were similar in the autoantibody-positive and autoantibody-negative participants, as was the frequency of ketoacidosis (43% vs 40%, P = 0.3). Autoantibody positivity was less common with increasing age (P = 0.0001), in males compared with females (82% vs 90%, P = 0.0001) and in people of non-white compared with white ethnicity (73% vs 86%, P = 0.0001). Body mass index was higher in autoantibody-negative adults than autoantibody-positive adults (median, IQR 25.5, 23.1–29.2 vs 23.9, 21.4–26.7 kg/m²; P = 0.0001). Autoantibody-negative participants were more likely to have a parent with diabetes (28% vs 16%, P < 0.0001) and less likely to have another autoimmune disease (4% vs 8%, P = 0.01).

Conclusions Most people assigned a diagnosis of type 1 diabetes presented with classical clinical features and islet autoantibodies. Although indistinguishable at an individual level, autoantibody-negative participants as a group demonstrated features more typically associated with other diabetes subtypes.

Strengths and limitations of this study

- We have studied a large multiethnic cohort of adults and children ≥5 years with clinically diagnosed incident type 1 diabetes in whom pancreatic islet autoantibodies were measured in a central laboratory.
- In routine practice, the initial assignment of a diagnosis of type 1 diabetes is a purely clinical one. The lack of further selection before inclusion in this study (eg, based on autoantibody status and/or genetic testing) renders the results of particular relevance to standard clinical care.
- Individual autoantibody-positive and autoantibody-negative patients were indistinguishable clinically, but the size and diversity of the cohort permitted group differences to be detected at high levels of statistical significance, suggesting diagnostic heterogeneity.
- As this was a volunteer study recruiting from mainly secondary care centres, ascertainment bias could have been introduced.
- Provision of a blood sample was optional, and autoantibody status was therefore available in just over half of the patients. Other than having a higher median age, this subgroup was representative of the whole cohort.

Trial registration number ISRCTN66496918; Pre-results.

INTRODUCTION

Type 1 diabetes (T1D) is an autoimmune disease that develops at any age but most frequently in children and young adults. Autoantibodies against islet antigens are typically present before, and for a variable time following, diagnosis. Once initiated, beta cell damage classically leads to progressive loss of insulin secretion and a need for lifelong insulin treatment.
The diagnosis of T1D is a clinical one but may be supported by the presence of one or more of the autoantibodies to islet cell antigens. In routine care, autoantibody status may not be available at diagnosis and may never be checked (management guidelines differ, with some not recommending their routine measurement or restricting measurements to situations where there is clinical doubt). Previous studies suggest that 80%–90% have detectable autoantibodies at disease onset, with a background autoantibody prevalence of around 2% in the young general population. Autoantibody positivity may be lower in some non-white ethnic groups. There is however uncertainty around the clinical and demographic correlates of autoantibody status in incident disease in an unselected multiethnic cohort including children and adults, using well characterised, validated assays. The After Diabetes Diagnosis REsearch Support System, supported by the National Institute for Health Research (NIHR) Clinical Research Network (CRN), recruits people with incident T1D from centres across England and Wales. We aimed to characterise these people with reference to their heterogeneity, focusing on the associations of autoantibody status with variation in presentation characteristics.

METHODS

Ethics approval
The project complies with the recommendations for research on human subjects by the 18th World Medical Assembly, Helsinki 1964 and later revisions and the International Conference on Harmonisation Guideline for Good Clinical Practice (Topic E6–10 June 1996). Protocol details have been reported previously and are therefore described in brief only.

Inclusion and exclusion criteria
People with a clinician-assigned diagnosis of T1D aged ≥5 years were recruited within 6 months of diagnosis. Written informed consent was obtained for all participants.

Data collection
On recruitment: demographic information; medications including insulin(s); medical history, including that of autoimmune disease; family history of diabetes; blood pressure; weight and height; HbA1c; and fasting or random blood glucose. A diagnosis of ketoacidosis was recorded if clinically assigned or if hyperglycaemia was accompanied by acidosis and either ketonaemia or ketonuria. Ethnicity was self-reported as one of 16 categories.

Project-specific blood sampling and measurement of islet autoantibodies
Sample donation was voluntary. Where collected, autoantibodies to glutamate decarboxylase (GADA), islet antigen-2 (IA-2A) and zinc transporter 8 (ZnT8A) were measured in sera using established radiobinding assays in a single central laboratory. Antibodies to both major ZnT8 isoforms, defined by the polymorphic amino acid at position 325 (arginine, ZnT8RA or tryptophan and ZnT8WA), were measured separately. Thresholds for autoantibody positivity were set at the 97th percentile of 974 control samples for GADA, the 98th percentile of 500 control samples for IA-2A and the 97.5th percentile of 523 healthy schoolchildren for both ZnT8RA and ZnT8WA. Positive autoantibody status was defined as positive for one or more of GADA, IA-2A or either form of ZnT8A. In the 2015 Islet Autoantibody Standardization Program Workshop, the assay sensitivities and specificities achieved were 74% and 96.7% for GADA, 72% and 100% for IA-2A, 60% and 100% for ZnT8RA and 46% and 100% for ZnT8WA, respectively.

Data analysis
Children were defined as aged <17 years. Body mass index (BMI) was derived as a z-score for children using WHO (2007) reference data. As a criterion for adiposity shared between children and adults, we applied WHO-recommended definitions of ‘normal’ weight (z-score <1, equivalent percentile <84.13% for children; BMI <25 kg/m² for adults, both including underweight) as distinguished from ‘overweight’ (z-score ≥1, percentile ≥84.13% for children, BMI ≥25 kg/m² for adults, both including obese). Parental and sibling history of diabetes was recorded. No attempt was made to differentiate between diabetes types in the family history. Variables were categorised as ‘individual characteristics’ and ‘diabetes presentation’. We analysed data from participants recruited between 1 September 2011 and 30 April 2016, with data querying and verification completed in November 2016.

Statistical analysis
Statistical analysis was carried out using StataCorp 2013 (Stata Statistical Software: V.13). Median and IQRs were used to summarise continuous variables. Categorical variables were summarised as percentages. The Mann-Whitney U test and Kruskal-Wallis test were used for between-group comparisons of continuous variables. \( \chi^2 \) testing was used for comparisons of categorical variables. Individual characteristics were explored as predictors of diabetes presentation and antibody status in univariate logistic and linear regression analyses. Multiple logistic and linear regression were used to establish the independence of predictors. A significance level of \( P<0.05 \) (two sided) was taken as a guide to interpretation (actual \( p \) values down to \( P<0.0001 \) are reported throughout).

Patient involvement
Patient and public involvement groups within the NIHR CRN representing people with diabetes, and representing children and young people, had input into the design of the patient information sheets, consent forms and recruitment strategies. After the start of recruitment,
RESULTS

Overview of the cohort

Individual characteristics

Data were analysed for 3312 participants recruited with incident T1D (1879 (57%) males and 1946 (59%) children from 146 centres; table 1). The slight male predominance (57%) was more prominent in adults than in children (61% vs 54%, P<0.0001). Islet autoantibodies were measured in the 1778 participants who donated an optional blood sample. For individual characteristics, data recording was >98% complete for all variables except symptom duration (94%). Sample sizes for incomplete data are reported in the tables. Of the total cohort, people of white European origin comprised 91% (n=3021), Asian (not Chinese) 3% (n=107), African-Caribbean 2% (n=63) and other or mixed ethnicity 3% (n=121). Median time from diagnosis to recruitment was 71 days (IQR 40–119) and to blood sampling was 75 days (IQR 42–126). Of those with body weight measured (n=2911), 35% were classified as overweight or obese, more commonly in adults than children (41% vs 31%, P<0.0001). Where records of body weight were available within 28 days of diagnosis (n=554), 35% were also overweight or obese (adults 40% vs children 29%, P<0.005).

Data collection for ketoacidosis at diabetes presentation was based on a record of it being assigned clinically or of hyperglycaemia accompanied by acidosis and either ketonaemia or ketonuria.

The relationships between diabetes presentation and individual characteristics

The main presenting features (table 1) were: osmotic symptoms (polyuria and/or polydipsia): 96%, weight loss: 85%, and fatigue: 82%. Ketoacidosis was identified at clinical presentation in 42%. Another autoimmune disease was present in 6%, 15% had a parent with diabetes and 8% had a sibling with diabetes.

The influence of age

Increasing age was independently associated with an increased prevalence of ketoacidosis, weight loss and fatigue at presentation, decreased prevalence of osmotic symptoms and longer symptom duration (figure 1). In accord with its relationship with age, ketoacidosis was less common in children than adults (39% vs 45%, P=0.0002). Although statistically significant, the differences between children and adults in other presenting symptoms were small (osmotic symptoms: 97% vs 95%, P=0.001; weight loss: 82% vs 89%, P=0.0001; fatigue: 78% vs 88%, P=0.0001, respectively), as were differences in symptom duration (median, 3 vs 4 weeks, P=0.0001).

The influence of gender

Female sex was independently associated with longer symptom duration (figure 1). Median symptom duration in females and males were 4 and 3 weeks, respectively (P=0.0001).

Associations with ethnicity

There were no significant associations between ethnicity and initial clinical presentation, including ketoacidosis, which was equally likely: in white Europeans and non-whites (41% and 44%, respectively; P=0.3).

Family history of diabetes

Having a parent with any diabetes was associated with a lower probability of presenting with ketoacidosis (figure 1; 12% vs 18%, P<0.0001). The same applied to diabetes presentation features was >98% complete for all variables except symptom duration (94%).
those who had a sibling with diabetes (figure 1; 5% vs 10%, P<0.0001).

Other autoimmune disease
The presence or absence of another autoimmune disease had no significant influence on diabetes presentation.

Description of the cohort in whom autoantibodies were measured
Children comprised 38% of the subgroup of 1778 participants who provided a blood sample for autoantibody measurement. The subgroup with blood samples was, accordingly, significantly older than the full cohort (median (IQR) 21.6 (13.4–32.8) vs 14.6 (10.4–26.4) years, P<0.0001). Other parameters were similar. One or more autoantibodies were present in 85% of those who donated a blood sample.

The relationships between autoantibody status and individual characteristics
The influence of age
Autoantibody positivity decreased with increasing age; adults were less likely than children to be positive for one or more antibodies (82% vs 90%, P<0.0001). The decline in autoantibody positivity continued...
throughout adult life (figure 2). Of the individual auto-
antibodies, GADAs were the most frequently observed
in adults, with IA-2A and ZnT8A being relatively more
common in children (figure 2). The autoantibody-pos-
itive adults were of lower BMI than autoantibody-nega-
tive adults (BMI, median (IQR) 23.9 (21.4–26.7) vs 25.5
(23.1–29.2), P<0.0001), and they were less likely to be
overweight or obese (40% vs 55%, P=0.0001). No rela-
tionship to BMI z-score was observed in children, and
there was no independent relationship between over-
weight/obesity and antibody positivity across children
and adults (figure 1).

The influence of gender
Females were more likely than males to be antibody posi-
tive (90% vs 82%, respectively (P<0.0001).

Associations with ethnicity
There was a statistically significant relationship between
ethnicity and autoantibody status (on χ² analysis, P<0.0001).
Among the three major non-white ethnic groups (Asian,
African-Caribbean and other or mixed ethnicity), numbers
with autoantibodies measured were small (n=46, 36 and
51, respectively) and the proportion with autoantibody
positivity did not differ significantly (70%, 64% and 82%,
respectively; P=0.1). People of non-white ethnic origin were
therefore grouped, and comparisons were limited to white
European versus non-white ethnic origin. White Euro-
pean ethnicity was independently associated with a higher
prevalence of autoantibody positivity compared with the
combined non-white group (86% and 73%, respectively;
P<0.0001; figure 1).

Family history of diabetes
People who had a parent with diabetes were less likely
than those without to have autoantibodies (figure 1; 16%
vs 28%, P<0.0001, table 2).

Other autoimmune disease
Another autoimmune disease was present in 204 partic-
ipants and 117 of those in whom autoantibodies were
measured. A history of another autoimmune disease was
positively associated with pancreatic islet autoantibody
positivity (P=0.01, figure 1), being present in 8% of the
autoantibody-positive participants and 4% of the autoan-
tibody-negative participants (table 2).

The relationship between autoantibody status and diabetes
presentation
A statistically significant relationship between posi-
tive autoantibody status and diabetes presentation
was restricted to a very small increase in prevalence of
osmotic symptoms (97% vs 94%, P=0.02, table 2). There
was no significant difference in rates of ketoacidosis at
presentation between autoantibody-positive and autoan-
tibody-negative participants (43% vs 40%, P=0.3, table 2).

DISCUSSION
For the first time, relationships between autoantibody
status (measured centrally in a single reference labo-
ratory) and phenotypic features in incident T1D are
reported from a large, unselected, multiethnic popu-
lation of both children and adults. The study was conducted
with support from the NIHR CRN with most participants
recruited from specialist centres. Based on estimates of
T1D incidence and population demographics,23 24 20%–
25% of eligible incident cases in England and Wales were
recruited.

Male predominance is unusual for an autoimmune
condition but has been reported in adults with incident
T1D.25 In young children, the sex ratio has been reported
to be close to unity.26 In the present study, the male excess
was observed also in children, although the excess was less
marked than in adults. Symptoms at presentation were as

Figure 2 The percentage of participants exhibiting islet autoantibodies (any and individual) in relation to age at diagnosis.
GAD, glutamate decarboxylase; IA2, islet antigen-2; ZnT8, zinc transporter 8.
expected.\textsuperscript{27, 28} Although weight loss was common, average body weight at the time of recruitment was normal, and many participants were overweight or obese, especially adults. This was apparent even in people with body weight measurements obtained within 28 days of diagnosis, bellying the belief that patients presenting with T1D are underweight. An association between increased BMI and increased risk of progression from autoantibody positivity to development of diabetes in at-risk relatives has been reported previously.\textsuperscript{29} Symptom duration was similar to that reported previously by others and was shorter in children than adults.\textsuperscript{30, 31} This may reflect parental vigilance of unwell children or a more insidious onset of clinical disease in older people.

The overall frequency of ketoacidosis at diagnosis (42\%) was high and slightly more so in adults than children. It occurred with similar frequency in white European and non-white ethnic groups. The figure of 42\% is higher than in previous reports from the UK (23\% in a recent national paediatric audit,\textsuperscript{32} 26\%–27\% in regional studies\textsuperscript{33, 34}), and a range of 15\%–80\% in those aged <20 years has been reported internationally.\textsuperscript{35} A very similar figure (40.3\%) has been reported recently for children in Italy.\textsuperscript{36} Ketoacidosis at diagnosis is a quality issue as it reflects lack of awareness of diabetes features among professionals and the general population,\textsuperscript{32} and efforts to increase awareness lead to reductions in ketoacidosis at first presentation.\textsuperscript{37} All methods of estimating the frequency of ketoacidosis at diagnosis have limitations, often leading to under-reporting.\textsuperscript{32} Strengths of the current study include the large number of patients and the ability to confirm or refute the diagnosis where this was in doubt. A limitation is that ascertainment bias could be introduced because recruitment was mainly from secondary care, and those who are the most ill at diagnosis may be the most likely to volunteer or to be referred. The higher ketoacidosis rate in adults versus children in our study appears at variance with the observation that ketoacidosis or severe ketoacidosis is more common in younger than in older children.\textsuperscript{28, 33, 36, 38–40} The current study did not include children <5 years of age, the group in childhood in whom ketoacidosis at diagnosis occurs most frequently,\textsuperscript{41} and this may have contributed to the apparent children to adult difference. Of course, if such younger children had been included, this could have increased the overall rate of ketoacidosis even higher. The lower rate associated with having a parent or a sibling with diabetes could result from a heightened awareness of symptoms leading to earlier clinical referral.\textsuperscript{35} The absence of any significant ethnic influence on ketoacidosis is at variance with

### Table 2

<table>
<thead>
<tr>
<th>Characteristics of pancreatic autoantibody (Ab) positive and negative participants (n=1778 with known antibody status)</th>
<th>Ab positive (n=1510)</th>
<th>Ab negative (n=268)</th>
<th>P values</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Individual characteristics</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Age</td>
<td>20.1 (13.1–31.1)</td>
<td>31.4 (17.7–41.0)</td>
<td>0.0001</td>
</tr>
<tr>
<td>Male</td>
<td>56 (851)</td>
<td>72 (192)</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Children</td>
<td>41 (614)</td>
<td>25 (66)</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td><strong>Body mass index</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Children (z score, n=545, 56)*</td>
<td>0.41 (−0.35 to 1.19)</td>
<td>0.47 (−0.48 to 0.97)</td>
<td>0.4</td>
</tr>
<tr>
<td>Adult (kg/m², n=825, 184)</td>
<td>23.9 (21.4–26.7)</td>
<td>25.5 (23.1–29.2)</td>
<td>0.0001</td>
</tr>
<tr>
<td>Overweight (n=1370, 240)</td>
<td>36 (490)</td>
<td>48 (114)</td>
<td>0.0005</td>
</tr>
<tr>
<td>White European ethnicity</td>
<td>86 (1413)</td>
<td>14 (232)</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Other autoimmune disease (n=1495, 265)</td>
<td>8 (117)</td>
<td>4 (10)</td>
<td>0.01</td>
</tr>
<tr>
<td>Parent(s) with any diabetes (n=1493, 261)</td>
<td>16 (233)</td>
<td>28 (74)</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Siblings with any diabetes (n=1374, 238)</td>
<td>9 (117)</td>
<td>8 (20)</td>
<td>0.9</td>
</tr>
<tr>
<td><strong>Diabetes presentation</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ketoacidosis (n=1483, 260)</td>
<td>43 (639)</td>
<td>40 (104)</td>
<td>0.3</td>
</tr>
<tr>
<td>Osmotic symptoms (n=1495, 267)</td>
<td>97 (1444)</td>
<td>94 (250)</td>
<td>0.02</td>
</tr>
<tr>
<td>Weight loss (n=1480, 267)</td>
<td>87 (1285)</td>
<td>88 (235)</td>
<td>0.5</td>
</tr>
<tr>
<td>Fatigue (n=1490, 265)</td>
<td>86 (1282)</td>
<td>80 (213)</td>
<td>0.01</td>
</tr>
<tr>
<td>Symptom duration (weeks, n=1424, 246)**</td>
<td>6.8 (10.5)</td>
<td>10.4 (32.2)</td>
<td>0.004</td>
</tr>
</tbody>
</table>

Percentages (n) or medians (IQR) are shown.
Sample sizes (n) are given for each variable if data collection was incomplete.
*The children’s BMI z-scores expressed as percentiles (median (IQR)) were 66\% (36\%–88\%) in the autoantibody positive (n=545) and 68\% (32\%–83\%) in the autoantibody negative (n=56).
**Median and IQRs for symptom duration were identical: 4 (2–8); mean and SD is shown to clarify the direction of difference.
some previous reports where higher rates were observed in non-white subgroups.42 43

One or more islet autoantibodies were observed in 85% of participants, more commonly in female than male and in younger compared with older participants. This is compatible with previous literature from the UK and other countries.8 44–46 although assay differences make such comparisons difficult. The positivity rate is higher than reported in people with T1D of non-white ethnic origin,12 13 although with the same caveats and bearing in mind that the previous studies of ethnic influences on antibody status have been limited in size or age range of population studied. The slight female autoantibody preponderance has been observed in other studies in young children, but not in older children and young adults.47 The higher autoantibody frequency in those with a coexistent autoimmune disease reflects the clustering of autoimmune disorders observed in T1D and shared genetic susceptibility.48 Overall, GADAs were the antibodies most commonly present, while IA-2A and ZnT8A were seen most frequently in children, findings compatible with previous studies.49 Insulin itself is considered a potential primary autoantigen as insulin autoantibodies are observed in incident T1D, especially in children.30 31 49 In the current study, most participants had received insulin therapy for weeks before study entry, and as they had developed antibodies to the exogenous insulin, insulin autoantibodies were not measured. In prospective studies of infants at high genetic risk of T1D, insulin autoantibodies were often detected earlier than the other islet autoantibodies,50 51 and in consequence, we may have underestimated the frequency of autoantibody positivity at diagnosis, especially in children.

Although autoantibodies can be present for years in people with diabetes who do not require insulin treatment immediately52–54 and are present in some diagnosed clinically with type 2 diabetes,55 they are generally regarded as a biomarker for T1D. In prospective studies they precede and predict the onset of T1D.56 They typically disappear, or titres drop to very low levels, in the years following diagnosis.57 In the autoantibody-negative participants studied here, several explanations may be proposed. First, insulin autoantibodies were not measured. Second, autoantibodies to as yet unknown antigens may have been present.58 The identification of tetraspanin-7 as an autoantigen could, for example, account for some apparently antibody-negative participants,59 although recent data suggest this is unlikely to account for large numbers.60 Third, autoantibodies may have disappeared or their levels diminished by the time of sampling. As all participants were recruited within 6 months of diagnosis, this is unlikely to be a major factor. Fourth, autoantibodies might develop subsequently, as reported previously for islet cell antibodies in a small proportion of patients.61 Fifth, some people may have autoimmune T1D without a humoral response. Finally, people may actually have another diabetes subtype. The autoantibody-negative participants as a group tended to be older and, if adult, more overweight. These features are compatible with type 2 diabetes. They were more likely to have a parent with diabetes, typical of type 2 or monogenic diabetes. They were more likely to be of non-white ethnicity, more associated with type 2 diabetes. Those with ketosis-prone diabetes (so-called idiopathic diabetes)62 as this is difficult to distinguish from T1D at first presentation. Further studies and follow-up of the cohort are planned to explore the extent to which T1D without detectable autoantibodies describes a subgroup of T1D that is distinct from other diabetes subtypes.

Acknowledgements We are grateful to Nish Chaturvedi and Joanne Holloway, who provided much advice and support at the initiation of the project and to Simon Howell for chairing the Management Committee. We thank the ADDRESS-2 Patient Advocate Group for their input into aspects of study design and conduct. We also thank Kyla Chandler, Claire Williams and antibody laboratory technicians at the University of Bristol for the assay of islet autoantibodies, Flora Christofor for study administrative support at Imperial College London and all the local research teams at participating centres. We are grateful to colleagues, George Alberti, Robert Eikelis, Rochan Agra-Jaffar, Ahmed El-Laboudi, Neil Hill, Monica Reddy and Shivshankar Seechurn for their critical review of early analyses of the results.

Collaborators AbithaKujavambal V; Abouqilah K; Ahmad J; Ahmad K; Ahmad M; Ahmed Z; Ajan R; Akiboye F; Ali A; Ali K; Anand B; Anderson C; Andrews R; Anjum B; Anthony S; Aung T; Ayooluwa O; Baburaj R; Bain S; Baker E; Baksai A; Banerjee K; Banerjeee M; Barnes D; Barnett A; Baynes K; Bellary S; Bennett S; Besser R; Bhardwaj P; Bhattacharyra B; Bhimsaria S; Bickerston A; Bilous R; Bingham E; Bodmer C; Bridges N; Brooks A; Browne D; Buck J; Cackett N; Caldwell G; Campbell F; Carey P; Cavavan D; Chandran S; Chandrasaksaret S; Chang J; Chankramath A; Chavdariani T; Chapman J; Chattington E; Cheng R; Cheng T; Chin Y; Chung C; Clifton R; Clifton J; Clift J; Clift P; Coldwell S; Cooper C; Coppini D; Creely S; Cronin M; Cummings M; Dang C; Darzy K; Dashaura O; Davies J; Davies M; Davies R; Day P; De Silva P; DelRio A; Delrio A; Desilva P; Dornhorst A; Douek I; Dowsett I; Dukhan K; Durkin S; Dutta A; Dutta S; Earle K; Edge J; Edmonds C; Edwards M; Elmati A; El-Reese F; Elrishi M; Evans M; Field B; Flanagan D; Franke B; Gable D; Gallow D; Ganapathi J; Gibbs J; Gibson M; Gnudi L; Gough A; Gough P; Gower K; Greening J; Gupta A; Gupta S; Gurrell E; Hamilton-Shield J; Hammond P; Han T; Hanna F; Hanson P; Haq M; Hardy K; Harms B; Hattersley A; Hayward J; Heald A; Hefferman H; Heller S; Hinde F; Hiltman G; Holt H; Howell S; Huma S; Hyer S; Jan O; Idris I; Ijaz S; Iqbal N; Issa B; Jackson A; Jackson A; Jain N; Jennings P; John K; Johnson R; Jones S; Joseph F; Joseph J; Joseph S; Kalilathi S; Kamulini N; Kapoor R; Karanidic S; Kelly P; Kershaw M; Khetriwal B; Kumar K; Lakdhar A; Lawrence J; Leong K; Levy D; Lowe A; Lyder G; Macklin A; Mallam K; Malik R; Mann N; Mansell P; Margabanthu G; Massoud A; Mathew V; McAulay A; McCowen B; Murphy N; Muzulu S; Mynlit K; Nabi D; Naik S; Narendran B; Natarajan A; Nathan Y; Nayar R; Ng M; Ngwu U; Nikokoom K; O’Connell I; Oelbaum R; O’Hare P; Oliver N; O’Malley B; Oso O; Overend L; Owen C; Owen K; Padinjarkara P; Pali N; Pang T; Paragas N; Patel K; Pathy K; Paul P; Paul Smith J; Peake M; Peet A; Pett A; Piel B; Pope R; Pothina N; Pothina N; Price H; Purewaal T; Puthra P; Puthra R; Puthra T; Rabbi T; Raffeeq P; Rahman S; Rasanhinga A; Randell T; Raith T; Rawl P; Rayman G; Regan F; Rice S; Riddle M; Robertson E; Robinson M; Russell-Jones D; Russell- Taylor M; Rutter M; Sampson M; Samuel J; Sankar S; Sankar V; Saravanathan P; Satish H; Savine R; Scanlon J; Scolibe I; Sennik D; Sharpe R; Shekar S; Sheshkar S; Simon G; Simpson G; Singh B; Singhal P; Smith D; Smith J; Smith P; Solomon H; Srinivasan B; Sriraman S; Stirling H; Strutt J; Surridge A; Sutchfield P; Taitnall S; Thanian K; Timmis A; Trevorley N; Turner B; Valabhi J; Venkatesh U; Vijayaraghavan S; Vijayaraman A; Vithian K; Vora J; Waldron-Lynch F; Walker M; Walsme D; Ward W; Watt A; Weaver J; Webber J; Weerasinghe K; West N; Whitehead P; Whitelaw D; Wickramasuriya N; Wilding J; Williams P; Williams S; Winkley K; Wright N; and Yalliwal C; Yarlagadda S; Zac-Varghese S.

Contributors Literature searches were carried out by VB, HCW, IFG, NSO and DGJ. AK, HCW, IFG, AKJW, PJB, DDB and DGJ were involved in the study design. Data collection was coordinated by AK and HCW. AK, HCW and IFG were responsible for data management. AKJW and PJB were responsible for autoantibody measurements. The statistical analysis plan was developed by IFG. Analysis of the data was by VB, AK and IFG. Data were interpreted by all authors. The manuscript


Open Access

7

Downloaded from http://bmjopen.bmj.com/ on April 12, 2018 - Published by group.bmj.com
was prepared by authors VB, AK, HCW, IFG, AKW and DGL. Critical revisions were made by authors PJDB, DBD, CMD, MP and NSO. Figures were prepared by AK, HCW and IFG. The Management Committee oversaw access to the data and stored biological samples. The Patient Advocate Group had input into aspects of study conduct. Local investigators were responsible for the recruitment of participants and collection of data. HCW is the guarantor of this study.

Funding The National Institute for Health Research (NIHR) provided the funding for the research infrastructure of the national Clinical Research Network (CRN), with project-specific support from Diabetes UK (grant numbers 09/0003919 and 15/0005234) and the Juvenile Diabetes Research Foundation (grant numbers 9-2010-407 and 3-SRA-2015-36-A-N). The funders were consulted about the vision for a UK-based type 1 diabetes ascertainment network. The funders are represented on the management committee that oversees access to the data and stored biological samples. The funding was administered as a grant to Imperial College London. Imperial College London is the study Sponsor.

Provenance and peer review Not commissioned; externally peer reviewed.

Data sharing statement The full anonymous dataset is available to access via a management committee, which includes people living with type 1 diabetes, scientists, clinicians and funder representatives as members. Participants gave informed consent for data sharing subject to conditions described in the access procedure documents available from the study website: www.address2.org.

Open Access This is an Open Access article distributed in accordance with the Creative Commons Attribution Non Commercial (CC BY-NC 4.0) license, which permits others to distribute, remix, adapt, build upon this work non-commercially, and license their derivative works on different terms, provided the original work is properly cited and the use is non-commercial. See: http://creativecommons.org/licenses/by-nc/4.0/.

© Article author(s) (or their employer(s) unless otherwise stated in the text of the article) 2018. All rights reserved. No commercial use is permitted unless otherwise expressly granted.

REFERENCES


Relationship between islet autoantibody status and the clinical characteristics of children and adults with incident type 1 diabetes in a UK cohort

Vassiliki Bravis, Akaal Kaur, Helen C Walkey, Ian F Godsland, Shivani Misra, Polly J Bingley, Alistair J K Williams, David B Dunger, Colin M Dayan, Mark Peakman, Nick S Oliver and Desmond G Johnston

BMJ Open 2018 8:
doi: 10.1136/bmjopen-2017-020904

Updated information and services can be found at:
http://bmjopen.bmj.com/content/8/4/e020904

These include:

References
This article cites 55 articles, 27 of which you can access for free at:
http://bmjopen.bmj.com/content/8/4/e020904#ref-list-1

Open Access
This is an Open Access article distributed in accordance with the Creative Commons Attribution Non Commercial (CC BY-NC 4.0) license, which permits others to distribute, remix, adapt, build upon this work non-commercially, and license their derivative works on different terms, provided the original work is properly cited and the use is non-commercial. See: http://creativecommons.org/licenses/by-nc/4.0/

Email alerting service
Receive free email alerts when new articles cite this article. Sign up in the box at the top right corner of the online article.

Topic Collections
Articles on similar topics can be found in the following collections

Diabetes and Endocrinology (467)

Notes

To request permissions go to:
http://group.bmj.com/group/rights-licensing/permissions

To order reprints go to:
http://journals.bmj.com/cgi/reprintform

To subscribe to BMJ go to:
http://group.bmj.com/subscribe/