An alternative sensor-based method for glucose monitoring in children and young people with diabetes

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ABSTRACT
Objective To determine accuracy, safety and acceptability of the FreeStyle Libre Flash Glucose Monitoring System in the paediatric population.

Design, setting and patients Eighty-nine study participants, aged 4–17 years, with type 1 diabetes were enrolled across 9 diabetes centres in the UK. A factory calibrated sensor was inserted on the back of the upper arm and used for up to 14 days. Sensor glucose measurements were compared with capillary blood glucose (BG) measurements. Sensor results were masked to participants.

Results Clinical accuracy of sensor results versus BG results was demonstrated, with 83.8% of results in zone A and 99.4% of results in zones A and B of the consensus error grid. Overall mean absolute relative difference (MARD) was 13.9%. Sensor accuracy was unaffected by patient factors such as age, body weight, sex, method of insulin administration or time of use (day vs night). Participants were in the target glucose range (3.9–10.0 mmol/L) ~50% of the time (mean 12.1 hours/day), with an average of 2.2 hours/day and 9.5 hours/day in hypoglycaemia and hyperglycaemia, respectively. Sensor application, wear/use of the device and comparison to self-monitoring of blood glucose were rated favourably by most participants/caregivers (84.3–100%). Five device related adverse events were reported across a range of participant ages.

Conclusions Accuracy, safety and user acceptability of the FreeStyle Libre System were demonstrated for the paediatric population. Accuracy of the system was unaffected by subject characteristics, making it suitable for a broad range of children and young people with diabetes.

Trial registration number NCT02388815.

INTRODUCTION
The benefit of good glycaemic control to complications of diabetes is well recognised and self-monitoring of glucose levels is central to improving and maintaining good glycaemic control. This can be challenging in children and young people due to unpredictable and increased frequency of food intake, variable patterns of physical activity, increased rates of intercurrent illness, and psychological and hormonal challenges of adolescence; all of which can lead to greater glycaemic variability, wider glycaemic excursions and more frequent hypoglycaemia than seen in adults. Currently, self-monitoring of glucose for children and young people involves a fingerprick, four or more times daily. The burden of self-monitoring of blood glucose (SMBG) for patients is well known, and includes: pain, inconvenience, disturbance for night-time tests and embarrassment.

Continuous glucose monitoring (CGM) overcomes some of the burden of SMBG and provides a continuum of glucose data, including visibility of night-time glucose, rather than the discrete snapshots obtained by SMBG. A recent review of data from randomised controlled trials on CGM use in the paediatric population concluded that CGM use is safe and efficacious. However, CGM use in paediatrics has yielded mixed results on glycaemic control outcomes. Some studies demonstrated

What is already known on this topic?
- Achieving, improving and maintaining optimal glycaemic control in the paediatric population is challenging because of wide glucose variability, including detected and undetected hypoglycaemia and hyperglycaemia.
- Type 1 diabetes exchange data indicate that the majority (>75%) of children/young people with diabetes do not meet International Society for Paediatric and Adolescent Diabetes/American Diabetes Association guidelines for glycaemic control (glycated haemoglobin <58 mmol/mol (7.5%)).

What this study adds?
- The FreeStyle Libre System is safe and accurate when used by children and young people with diabetes (4–17 years).
- This new sensor-based glucose monitoring technology, requiring no user calibration, was highly acceptable to participants/caregivers. It is anticipated that the system could support enhanced diabetes management.
- Glycaemic variability data for this UK population of 4–17-year-olds show that these children/young people spent on average approximately 50% of the time in euglycaemia.
improved outcomes11–17 and associated increased benefit with increased use of CGM.16–19 Others showed little difference when comparing outcomes following use of CGM and SMBG; in these the lack of clinical benefit was often attributed to poor adherence to CGM use and reduced sensor wear.4 19–22

CGM is underused in children and young people, perhaps in part due to the lack of compelling evidence of the benefit it may provide. Barriers to CGM use include: sensor insertion on a weekly basis; skin reactions, discomfort/pain at the insertion site; frequent alarms especially at night (often turned off due to alarm fatigue); the need for multiple SMBG tests for calibration; interference with activities; poor sensor-receiver connectivity causing a loss of data; high cost and limited reimbursement.4 19–22 It has been proposed that reducing these barriers could lead to increased uptake of continuous sensing by children and young people.25

The FreeStyle Libre System (Abbott Diabetes Care) used in this study potentially resolves many of the issues that affect adherence with CGM. It provides comprehensive glucose data (similar to CGM systems), increased sensor wear duration (up to 14 days, increasing the time between sensor applications), and has no requirement for fingerprick calibrations. The reader, which displays glucose results after scanning over the small sensor worn on the back of the arm, does not need to be kept close to the sensor and has no automatic alarms (but can alert the user to hypo/hyperglycaemia and pending excursions when detected following a scan). Scanning can be performed as often as is needed, after each scan the reader displays current sensor glucose, a trend arrow and 8-hour glucose history. The sensor automatically stores glucose data every 15 min. Device software can be used to generate summary glucose reports for review by the patient at home or in clinic with their healthcare professional (HCP). These features were not available during the study; the device was masked to participants and HCPs. The system is designed to replace SMBG testing for self-management of diabetes, except during rapidly changing glucose, to confirm hypoglycaemia, or if symptoms do not match system results. This study is the first to report accuracy, safety and acceptability of the FreeStyle Libre System in children and young people. Accuracy of the blood glucose (BG) feature of the system is not under evaluation here; regulatory approval/Conformité Européenne (European Conformity marking of the device confirms this feature meets the required industry standards for performance. At the time of the study, the device was approved for use by adults; it has subsequently been approved for use with children (with supervision from a caregiver).

METHODS
This prospective, single arm study was conducted across nine diabetes centres in the UK.

Study participants
Children and young people aged 4–17 years, with type 1 diabetes (T1D) or type 2 diabetes treated with multiple daily injections (MDI) of insulin or continuous subcutaneous insulin infusion (CSII), and currently testing BG at least two times per day were eligible to participate. Exclusion criteria included: concomitant disease/condition that may compromise patient safety; other labelling prohibitions; currently using a CGM device; and known/suspected allergy to medical grade adhesives. Any potentially eligible patient from the general diabetes population at each study site was invited to participate.

Study design
Participants attended clinic three times. Adverse events (AEs) were reviewed at every participant contact. Visit 1 included a physical examination, FreeStyle Libre device training, sensor insertion and sensor application questionnaire. Participants wore the sensor (on the back of their upper arm) for up to 14 days and were asked to perform four capillary BG tests daily using the BG strip-port on the reader (FreeStyle Optium test strips, Abbott Diabetes Care), each immediately followed by an interstitial fluid (ISF) glucose sensor measurement (data masked to participants) to allow comparison of results between sensor and BG. Visit 2 (days 5, 6, 7 or 8) was a safety monitoring check-up, data was also uploaded. During the final clinic visit (days 12, 13, 14 or 15), the device was unmasked for subjects to experience its full functionality, sensors were removed (and the site inspected for any adverse reactions), data were uploaded and user questionnaires completed. Three sensor lots were used. Participants were asked to maintain their pre-existing diabetes self-management plan throughout the study.

The study (NCT02388813) was conducted in compliance with the study protocol and International Conference on Harmonisation Guideline for Good Clinical Practice. Competent authority (MHRA) and ethics approval was obtained and each participant or their caregiver gave written, informed consent prior to participation in the study.

Statistical analyses
For the primary accuracy end point, it was determined that a minimum sample size of 20 participants was needed to obtain a 95% CI around an estimate of the percentage of results in zone A of the consensus error grid (CEG) of ±5%. However, a larger sample size was used in order to compare results to a similar study performed with adult participants.

Sensor values were compared with temporally matched (within ±5 min) BG values using a number of methods to evaluate overall accuracy of the system: mean absolute relative difference (MARD), median absolute relative difference (median ARD), mean relative difference (MRD), proportion of results within ±1.1 mmol/L of the BG value for glucose <5.55 mmol/L and within ±20% of the BG value for glucose ≥5.55 mmol/L (hereafter %within1.1/20). Mean absolute difference (MAD) and MARD were used to evaluate accuracy at low and mid to high glucose ranges, respectively. (MARD, median ARD, MRD and MARD are all measures of the average difference between sensor values and reference values. MARD and median ARD measure the size but not the direction (higher/lower) of the differences compared with the reference (absolute) as a percentage of the reference value (relative). MAD is similar but just reports the size of the difference (it is not reported as a percentage); this is commonly used to assess accuracy at low glucose levels. MRD measures the size and direction of the difference compared with the reference as a percentage of the reference value.)

CEG analysis, a tool developed to determine clinical accuracy of SMBG systems, was used to quantify the clinical accuracy of the device.25 A grid of different zones defines the clinical significance of any difference between test and reference results, as outlined in figure 1A.

Analysis of variance (ANOVA) and simple linear regression were used to assess whether a range of factors affected sensor accuracy.

Hypoglycaemia/hyperglycaemia alert features were evaluated by assessing concordance of sensor alerts obtained within ±15 min of BG results in hypoglycaemia (<3.9 mmol/L) and...
hyperglycaemic (>13.3 mmol/L) ranges. Rates with pending alerts (ie, those with sensor results within ±10% of the BG threshold) were also determined.

Measures of glycaemic variability were considered as a secondary end point of the study. Time in range (TIR) was defined as glucose results between 3.9 mmol/L and 10.0 mmol/L. A hypoglycaemic/hyperglycaemic excursion was defined as ≥2 consecutive results outside the predefined range, corresponding to >15 min.

Safety outcomes were analysed for all participants who enrolled.

All analyses were performed using SAS software, V.9.2 or later (SAS Institute, Cary, North Carolina, USA).

RESULTS

Eighty-nine participants were enrolled in the study, all with T1D; 50.6% were male; 56% used CSII and 44% used MDI; mean age was 10.4 years, with 24 participants aged 4–7 years, 39 aged 8–12 years and 26 aged 13–17 years. Duration of diabetes was 4.0±2.8 (0.3, 12.2) years (mean±SD (min, max)) and HbA1c (glycated haemoglobin) was 60±12 (38, 90) mmol/mol (7.6±1.1 (5.6, 10.4) %).

Paired sensor-BG results were not available for 2 participants (1 withdrawal prior to obtaining sensor data, 1 protocol deviation led to no reader BG data); therefore, 87 evaluable participants were included in the accuracy analyses and 88 were included in the glycaemic variability analysis. All 89 participants were included in the safety analyses.

Accuracy performance

For assessment of sensor accuracy against capillary BG there were 5493 paired sensor-BG results. Overall MARD was 13.9%, median ARD was 10.4%, MRD was 1.7% and the % within1.1/20 was 82.0%. CEG analysis demonstrated 83.8% of results in zone A and 99.4% of results in zones A and B (figure 1A). Sensor results were in good agreement with BG results—Deming regression: slope=1.03, intercept=−0.23 mmol/L, correlation coefficient=0.95. For paired results at lower glucose concentrations, with BG<5.55 mmol/L (n=1468), MAD was 0.75 mmol/L; for paired results at higher glucose concentrations, BG 5.55–10.0 mmol/L (n=2090), MARD was 13.5%; and BG>10.0 mmol/L (n=1935), MARD was 10.6%.

In some sensor systems, a lag is evident between BG and sensor results, such that sensor results are higher than BG results when glucose is decreasing and lower than BG results when glucose is increasing.27 This trend (sensor results higher/lower than BG when glucose is decreasing/increasing) was not evident with the FreeStyle Libre System (figure 1B), demonstrating that there is little ‘lag’ effect.

There were no statistically significant differences in accuracy detected (ANOVA) for sex (p=0.951), method of insulin administration (p=0.640), sensor lot (p=0.133) or daytime versus night-time use (p=0.909), as the percentage of results within CEG zone A were similar for the subgroups. Accuracy was also not significantly influenced by age or body weight—the regression slopes (for percentage within CEG zone A vs participant age and weight) were not significantly different from zero (p=0.133 and p=0.284, respectively).

The sensor detected hypoglycaemia (when capillary BG <3.9 mmol/L) on 70% (438/622) of occasions, increasing to...
84% when pending alerts (sensor results within 10% of the hypoglycaemic threshold) were included. For the 30% of occasions where hypoglycaemia measured in capillary test was not detected by the sensor, further analysis of those results for clinical significance using the CEG indicates that 164 were in zones A and B (clinically acceptable) and 20 were in zone C (altered clinical action—likely to affect clinical outcome). The sensor detected hyperglycaemia (when BG >13.3 mmol/L) on 85% of occasions, increasing to 94% when pending alerts were included (n=999).

**Glycaemic variability**

Sensor data collected during the study were used to retrospectively determine glycaemic variability of participants during their time in the study (table 1). Participants were in euglycaemia (3.9–10.0 mmol/L) ~50% of the time; this was the same through day and night, and for both methods of insulin administration. The relationship of TIR to HbA1c is similar for older and younger participants (p=0.229 and 0.297 for slope and intercept, respectively, figure 2). On average, participants spent 2.2 hours/day and 9.6 hours/day in hypoglycaemia (<3.9 mmol/L) and hyperglycaemia (>10.0 mmol/L), respectively. The breakdown of glucose excursions through daytime and night-time periods and for participants using CSII and MDI is shown in table 1.

Comparison of glycaemic variability observed in this study to that reported for other studies is made in the online supplementary material.

**User acceptability**

Study participants and/or their caregivers rated their experience with the system on a scale of 0 (strongly agree) to 4 (strongly disagree). Statements about sensor application (84.3–92.1%), sensor wear and use (87.2–100%), comparing use to SMBG (85.4–97.5%) and the device itself (68.3–96.3%) were rated favourably (0 or 1) by most respondents. Figure 3 summarises responses by sex, and shows the system was perceived similarly by both sexes.

**Safety**

One participant had a serious AE that was not related to the study or device (pain and lack of feeling in leg). Five device related AEs were reported in total from five (6%) participants, aged 6, 9, 10, 12 and 15 years: allergic reaction, blister, pink mark/scabbing and abrasion on sensor removal (n=2)—four were mild, one was moderate, all were resolved at study completion.

Site exams performed for all sensor insertions checked for anticipated AEs associated with sensor application or insertion sites—moderate erythema was observed on 11.6% of occasions, mild erythema and pain 13.6% and 4.1%, respectively, and mild instances of bleeding, bruising, itching and oedema were each on <3% of occasions. There were no trends in rate of anticipated AEs, including erythema, with age.

**DISCUSSION**

This study was the first to evaluate performance and usability of the FreeStyle Libre System in children and has demonstrated good agreement between sensor and BG results. The system is unique among ISF glucose monitoring sensors in that it can be worn for up to 14 days and does not require user calibration. Comparison of accuracy data from this study to that from other paediatric studies using ISF glucose sensors demonstrated that the factory calibrated system provided similar accuracy to systems requiring daily fingerprick calibrations (table 2). The lack of user calibration eliminates potential

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**Table 1** Average time in key glycaemic ranges

<table>
<thead>
<tr>
<th></th>
<th>Overall</th>
<th>Time of day</th>
<th>Insulin administration</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Overall</td>
<td>CSII</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Daytime (6:00–22:00)*</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Night-time (22:00–6:00)†</td>
<td></td>
</tr>
<tr>
<td>n (participants)</td>
<td>88</td>
<td>88</td>
<td>87</td>
</tr>
<tr>
<td>Sensor glucose, mmol/L</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mean (SD)</td>
<td>9.7 (2.0)</td>
<td>9.9 (2.1)</td>
<td>9.2 (2.4)</td>
</tr>
<tr>
<td>Time in range, hypoglycaemia and hyperglycaemia, hours per day (*per day=16-hour period; †per day=8-hour period)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3.9–10 mmol/L</td>
<td>mean (SD)</td>
<td>12.1 (3.7)</td>
<td>8.1 (2.6)</td>
</tr>
<tr>
<td></td>
<td>min, max</td>
<td>3.6, 21.0</td>
<td>2.7, 14.3</td>
</tr>
<tr>
<td>&lt;3.0 mmol/L</td>
<td>mean (SD)</td>
<td>2.2 (2.2)</td>
<td>1.2 (1.3)</td>
</tr>
<tr>
<td></td>
<td>min, max</td>
<td>0.0, 13.5</td>
<td>0.0, 9.0</td>
</tr>
<tr>
<td>&gt;10.0 mmol/L</td>
<td>mean (SD)</td>
<td>0.9 (1.6)</td>
<td>0.4 (0.9)</td>
</tr>
<tr>
<td></td>
<td>min, max</td>
<td>0.0, 10.5</td>
<td>0.0, 7.0</td>
</tr>
<tr>
<td>&gt;13.3 mmol/L</td>
<td>mean (SD)</td>
<td>9.6 (4.2)</td>
<td>6.7 (2.9)</td>
</tr>
<tr>
<td></td>
<td>min, max</td>
<td>0.0, 19.5</td>
<td>0.0, 13.1</td>
</tr>
<tr>
<td>Frequency of hypoglycaemia and hyperglycaemia events per day (24 hours)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt;3.0 mmol/L</td>
<td>mean (SD)</td>
<td>1.3 (1.0)</td>
<td>1.0 (0.8)</td>
</tr>
<tr>
<td></td>
<td>min, max</td>
<td>0.6, 6.8</td>
<td>0.4 (0.6)</td>
</tr>
<tr>
<td>&gt;10.0 mmol/L</td>
<td>mean (SD)</td>
<td>2.6 (0.8)</td>
<td>2.3 (0.7)</td>
</tr>
<tr>
<td></td>
<td>min, max</td>
<td>1.8 (0.8)</td>
<td>1.5 (0.7)</td>
</tr>
<tr>
<td>Overall glucose variability</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>SD, mmol/L</td>
<td></td>
<td>4.5 (1.1)</td>
<td>4.5 (1.1)</td>
</tr>
<tr>
<td>Coefficient of variation (subject level), %</td>
<td>46.5 (7.8)</td>
<td>45.7 (8.0)</td>
<td>43.6 (9.6)</td>
</tr>
</tbody>
</table>

CSII, continuous subcutaneous insulin infusion; MDI, multiple daily injection.
variations in sensor systems that may be introduced through errors in BG results used for calibration, calibration at inappropriate times (during rapidly changing glucose), missing calibrations or use of sensor rather than BG values for calibration.31–33

Comparison of results from this study to those from a study where adults used the system34 shows: MARD versus capillary BG of 13.9% and 11.4%; percentage in CEG zone A of 83.8% and 86.7%; and percentage in CEG zones A and B of 99.4% and 99.7%, in children and adults, respectively. This trend is similar to that reported for other ISF systems: Dexcom G4 Platinum had MARD versus capillary BG of 15% and 14.0% in children and adults, respectively.43–55 Similarly, Dexcom G4 Platinum with software 505 algorithm had MARD versus capillary BG of 13% and 11.3% in children and adults, respectively.33–35

This study has confirmed that sensor accuracy is robust to patient characteristics such as age, body weight, method of calibration, and age group. Regression statistics:

- For <13 years: TIR = 27.2113 – 1.9369 x HbA1c
- For ≥13 years: TIR = 32.1350 – 2.6536 x HbA1c

Slope and intercept for the different age groups was not significantly different, p=0.229 and 0.297, respectively.
insulin administration and sex. Accuracy of the system along with its robustness to differing patient characteristics, are sufficient to recommend its use to paediatric patients as support in optimising their glycaemic control. The study identified that there were 20/622 occasions where hypoglycaemia measured in capillary testing was not detected by the sensor and CEG analysis identified that there may be altered clinical action based on the sensor result. The product labelling advises patients to use a BG meter to check the sensor result if symptoms do not match the sensor result.

This study, providing the first glucose variability data for a paediatric population in the UK, and others,\textsuperscript{7} 13-14 17 21-23 show there is significant opportunity to improve glycaemic control in children and young people with diabetes. Several studies have associated improved outcomes with increased sensor wear,\textsuperscript{16-19} A sensor with a longer wear period, that does not require finger-prick calibration, may support more frequent sensor use, and thus help improve clinical outcomes in children and young people.

SMBG provides single, intermittent results, which may not capture intervals of extreme variability or nocturnal events. In comparison, the system displays the current glucose result, the last 8 hours of glucose data (including night-time results without any disturbance to the child) and trend information when the sensor is scanned—providing significantly more data to guide treatment for this challenging population than routine SMBG.

This study has demonstrated safety and user acceptability of the system. Device related AEs occurred in 6\% of participants, the same rate was observed in adults.\textsuperscript{14} The AEs were reported by participants aged 6–15 years, and thus were not concentrated in a particular age group. Anticipated sensor application/insertion site symptoms (pain, bleeding, bruising, erythema, itching, oedema) were reported in 44\% of participants (63\% of which were mild in severity), this is similar to the rate observed in adults,\textsuperscript{14} where 36\% of participants experienced skin issues (p=0.337). Erythema was the symptom noted most frequently and, as with other anticipated AEs, was not associated with particular age groups. Despite these symptoms, results of the user questionnaires were similar for boys and girls and were largely favourable, culminating in 96.3\% of respondents stating they would recommend the system to someone else with diabetes.

A limitation of this study is the single body site used. Additional studies are needed to determine suitability of additional body sites for sensor wear and longer-term studies are needed to evaluate whether this system can provide improved adherence with sensor wear and improved glycaemic outcomes with continued use over time. Exploring alternate reference methods and more detailed analysis of the glycemic variability data using subgroups (eg, age) may be of interest.

Table 2 Comparison of accuracy for factory calibrated and user calibrated sensor systems

<table>
<thead>
<tr>
<th>Accuracy measure</th>
<th>Factory calibrated</th>
<th>Require daily fingerpuck calibrations</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean absolute relative difference, %</td>
<td>FreeStyle Libre</td>
<td>Glucoday\textsuperscript{18}</td>
</tr>
<tr>
<td></td>
<td>13.9</td>
<td>18.0, 19.0</td>
</tr>
<tr>
<td>Percentage of results in zone A of CEG</td>
<td>83.8</td>
<td>Medtronic REALTime\textsuperscript{29 30}</td>
</tr>
<tr>
<td></td>
<td>NA</td>
<td>NA, 66.7</td>
</tr>
<tr>
<td>Percentage of results in zones A and B of CEG</td>
<td>99.4</td>
<td>Dexcom G4 Platinum\textsuperscript{4}</td>
</tr>
<tr>
<td></td>
<td>98.6</td>
<td>Dexcom G4 Platinum with software 505 algorithm\textsuperscript{1}</td>
</tr>
<tr>
<td></td>
<td>94.5, 92.3</td>
<td>99</td>
</tr>
<tr>
<td></td>
<td>100</td>
<td></td>
</tr>
</tbody>
</table>

CEG, consensus error grid; NA, not available.

CONCLUSIONS

This prospective study has clearly demonstrated the accuracy, safety and user acceptability of the FreeStyle Libre System for the paediatric population. Accuracy was unaffected by subject characteristics, making the system potentially suitable for a broad range of children and young people with diabetes. It is anticipated that the provision of comprehensive glucose data for up to 14 days, from a system that is easy to use, with reduced pain and burden for the user since there is no requirement for fingerprick calibration, could support enhanced diabetes management. Further studies are needed to determine whether improved outcomes can be achieved with prolonged use of the system.

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Contributors

Abbott Diabetes Care planned and funded (including provision of devices) the study described in the article. Prior to initiation of the study, site clinical staff were trained in use of the device by clinical staff from Abbott Diabetes Care. The study was conducted at the following sites: Oxford Children’s Hospital, principal investigator: JE, Addenbrooke’s Hospital, principal investigator: CA, Leeds Children’s Hospital, principal investigator: FC, Bristol Royal Hospital for Children, principal investigator: JH-S, Royal Devon and Exeter NHS Foundation Trust, principal investigator: CM, Harrogate District Hospital, principal investigator: SP, Nottingham University Hospitals NHS Trust, principal investigator: TR, Northampton General Hospital, principal investigator: AS, Southampton General Hospital, principal investigator: NT. The Paediatric Diabetes and Endocrinology Team, in particular Loraine Bunton and Jane Bowen-Morris, at Oxford Children’s Hospital aided in the design of the participant information sheets prior to submission to the ethics board. The NIHR Young Persons’ Advisory Group, Liverpool Regional Group, critically reviewed the design and content of participant information sheets prior to submission to the ethics board. Study staff at each site contributed to data collection. The principal investigator at each site contributed to the writing of the original article.

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Competing interests All authors' institutions received financial support from Abbott Diabetes Care to conduct the clinical trial. TR received honoraria/fees for consulting from Abbott Diabetes Care outside of the submitted work. The study sponsor (Abbott Diabetes Care) designed the study protocol in collaboration with the principal investigator and provided all study materials. The sponsor was involved in collecting data and reporting results. The sponsor also gave approval to submit for publication. The corresponding author had full access to all the data in the study and, together with all authors, had final responsibility for the decision to submit for publication.

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