Flash Glucose Monitoring System for Diabetes

Image courtesy of Abbott Diabetes Care
Summary

• The FreeStyle Libre Pro is a glucose monitoring system intended to replace finger-stick tests; it uses a sensor implanted in the arm that a health care provider scans with a specialized reader for a record of glucose levels, trends, and patterns in people with diabetes.

• The available evidence suggests that the performance and accuracy of the FreeStyle Libre Pro system is comparable to conventional methods of glucose monitoring, that it may reduce periods of hypoglycemia, or low blood sugar, and that it has high patient satisfaction.

• Although the FreeStyle Libre Pro system is approved for use in Canada, it is not yet commercially available here.

• Canadian pricing is not available, but cost-effectiveness analyses conducted in the UK and other European countries have concluded that it is cost-effective compared with the self-monitoring of blood glucose.

• A consumer version of the FreeStyle Libre Pro system intended for personal use by patients with diabetes is currently awaiting regulatory approval in the US.

The Technology

The FreeStyle Libre Pro system (Abbott Diabetes Care, Alameda, California) is a flash glucose monitoring system that provides a record of an individual’s glucose levels, trends, and patterns for up to 14 days. The FreeStyle system generates an ambulatory glucose profile — a novel way of assessing glucose levels on a continuous 24-hour basis that shows how day-to-day decisions and behaviours impact the control of blood sugar levels.

Similar to continuous glucose monitoring, flash glucose monitoring relies on the measurement of glucose concentrations in the interstitial fluid — the fluid surrounding cells — rather than directly measuring glucose levels in the blood. The FreeStyle Libre Pro system is intended for professional use only, i.e., the person with diabetes does not interact with the system. Rather, the sensor attachment is inserted and scanned by a health care provider to receive the data retroactively. A consumer version of the system, the FreeStyle Libre, is inserted and scanned by the person with diabetes, rather than the health care provider. It is currently under review by the US FDA and is already available in more than 35 countries worldwide (Jessica Sachariason, Abbott Diabetes Care, Alameda, CA: personal communication, 2017 Mar 10).
The FreeStyle Libre Pro system consists of a small, round, disposable, and water-resistant sensor approximately the size of two stacked quarters that is applied to the back of the individual's upper arm by a health care provider in a clinic setting.\(^2\) The sensor continuously measures glucose in the interstitial fluid through a small, 5 mm-long-by-0.4 mm-wide filament inserted just under the skin. The sensor records glucose levels every 15 minutes, capturing up to 1,340 glucose results for up to 14 days, and displays current, eight-hour historic and trend glucose data when scanned.\(^2\) Information about the individual's glucose levels is stored in the sensor which, in turn, is accessed by passing a specialized, hand-held reader over the sensor when the individual returns to the clinic or physician's office.\(^2\)

Flash glucose monitoring differs from continuous glucose monitoring in that information about the person's glucose levels and trends is available from the FreeStyle Libre Pro system only when the sensor is scanned. In comparison, continuous glucose monitoring systems monitor and send information to a device or display monitor, without interruption, throughout the day, and can alert a user if glucose levels are outside a pre-set limit.\(^1\)

An important distinction of the FreeStyle Libre Pro system compared with continuous glucose monitoring is that it is factory-calibrated and does not need daily calibration (or finger sticks) by the user. It also has a longer period of sensor use, 14 days versus 7 days, reads results in seconds, and can be used by people with type 2 diabetes.\(^7,8\) Two other key differences are that the FreeStyle Libre Pro system is not equipped with alarms (or notifications) that will sound in the event of out-of-range glucose values, and it lacks the ability to connect with insulin pump systems that continuously deliver short-acting insulin.\(^7,9\)

In the past, continuous glucose monitoring systems have been used primarily by people with type 1 diabetes; however, complications arising from suboptimal glucose control affects both those with type 1 and type 2 diabetes.\(^10\) The FreeStyle Libre Pro system has the potential to address an unmet need for more intensive glucose monitoring in patients with type 1 or type 2 diabetes who require more vigilant glucose monitoring but who find the currently available continuous glucose monitoring systems too difficult, expensive, or cumbersome to use.

**Availability**

The FreeStyle Libre Pro system received a Class III medical device licence from Health Canada in November of 2016.\(^11\) The approval extends to three components of the system: the reader kit, sensor kit, and software.\(^11\) The FreeStyle Libre Pro system does not appear to be commercially available in Canada yet; however, health care professionals can order it on-line directly from the manufacturer's US website (Jessica Sachariason; personal communication, 2017 Mar).\(^3,7\) The consumer version of the FreeStyle Libre Pro system is not currently licensed by Health Canada.

In September 2016, the FDA approved the FreeStyle Libre Pro system for people with diabetes aged 18 years and older.\(^12,13\) A consumer version of the FreeStyle Libre system intended for personal use is currently under review by the FDA.\(^2,8,14\) In Europe, the consumer version of the FreeStyle Libre system received a CE mark authorizing its use in adults in September of 2014, and a subsequent CE mark in February 2016 authorizing use in children and teens (aged four to 17 years).\(^6,15\)

**Cost**

Canadian pricing information for the FreeStyle Libre Pro system is not available.

In the US, the FreeStyle Libre Pro reader costs US$65 and each individual sensor costs US$60.\(^8\) In the UK, it is reported that the cost for a starter pack of the FreeStyle Libre consumer system, which includes one reader and two sensors, is approximately £170.\(^16\)

**Who Might Benefit?**

To date, continuous glucose monitoring has been used primarily by well-informed people with type 1 diabetes who have been trained on how to use it.\(^7\) In contrast, flash glucose monitoring may be a more accessible technology for people with type 2 diabetes, as it may enable appropriately trained people to monitor their glucose levels regularly after a meal or physical exercise, pending the availability of the consumer version of the system.\(^7\) It has been suggested that the FreeStyle Libre Pro system will be used primarily by people with type 2 diabetes; it could also benefit long-term users of continuous glucose monitors who find the alarms with most of these systems to be bothersome.\(^7,17\)
The FreeStyle Libre Pro system may also be of benefit to people who are unable or unwilling to check their blood sugar using finger sticks because of pain and discomfort, as the system does not need routine finger-stick testing. This may be especially beneficial for children with type 1 diabetes; however, in North America, the system is not currently approved for use in people younger than 18 years of age.

**Current Practice**

Measurement of glycated hemoglobin (A1C) is the preferred standard for assessing long-term blood glucose control in people with both type 1 and type 2 diabetes. A major drawback of A1C is that it does not provide information about fluctuations in glucose levels; i.e., it is possible for A1C to remain within the target range while glucose levels fluctuate between periods of low (hypoglycemia) and high (hyperglycemia) levels throughout the day.

"The FreeStyle Libre Pro system may also be of benefit to people who are unable or unwilling to check their blood sugar using finger sticks because of pain and discomfort..."

Self-monitoring of blood glucose using finger sticks and capillary blood glucose testing provides only discrete "snapshots" of glycemic control at the test times. Self-monitoring is further limited by its time-intensive nature and the discipline required to maintain records of daily blood glucose results over the long-term. The timing and frequency of self-monitoring of blood glucose depends on the type of diabetes and the treatment prescribed. It also depends on the individual's need for information about glucose levels and the ability to use the test information to modify behaviour or adjust treatment.

Continuous glucose monitoring has been used in people with type 1 diabetes to provide comprehensive, continuous glycemic profiles, which can improve glycemic control and reduce hypoglycemia. Notably though, continuous glucose monitoring systems still require self-monitoring for calibration of the system. Self-monitoring must be done at least every 12 hours and also in "real time" to confirm interstitial fluid measurements before making changes to therapy or treating suspected hypoglycemia.

**The Evidence**

Nine published studies of the FreeStyle Libre system, used in adults, were identified. Although it is the FreeStyle Libre Pro system that has been approved for use in Canada, it appears these studies evaluated the consumer version (the FreeStyle Libre system), which is designed for personal use and has been available in Europe since 2014. Four of the studies were conducted on people with type 1 diabetes, four on a mixed population of people with type 1 or type 2 diabetes, and one study was on people undergoing cardiac surgery. Six of the studies were funded by the manufacturer and three were investigator-initiated.

Five of the nine studies evaluated the performance and accuracy of the FreeStyle Libre system compared with other methods of measuring glucose. The other four studies reported clinical outcomes (for example hypoglycemia, A1C, patient satisfaction, and quality of life). Of these, three were observational, single-arm studies and one was a prospective, open-label, multi-centre, randomized controlled trial (RCT), the IMPACT study. The primary outcome in the IMPACT study was the change in the amount of time spent in hypoglycemia from baseline to six months in 241 patients with well-controlled type 1 diabetes who were randomized to either the FreeStyle Libre system or to conventional self-monitoring of blood glucose.

We also identified two published studies of the FreeStyle Libre system used in children with type 1 diabetes. One trial was funded by the manufacturer and the other was independently funded. The first study compared device performance and patient satisfaction between the FreeStyle Libre system and capillary blood glucose testing in the same children (age range: one to 14 years). The second was a pilot study to evaluate the feasibility and acceptability of the ambulatory glucose profile in children aged four to 17 years who were initiated on the FreeStyle Libre system.
Performance and Accuracy

The performance and accuracy of the FreeStyle Libre system in adults and children compared with self-monitoring of blood glucose or capillary blood glucose testing, venous blood glucose analysis, continuous glucose monitoring, or intravascular microdialysis continuous glucose monitoring, are summarized in Table 1. All the studies in Table 1 reported the overall mean absolute relative difference (MARD), expressed as a percentage. A lower MARD value indicates smaller differences, or per cent error, between methods. The overall MARD between the FreeStyle Libre system compared with capillary blood glucose testing ranged from 10.0% to 13.9%; the MARD was 16.6% compared with self-monitored blood glucose, and 18.1% compared with continuous glucose monitoring. When directly compared with reference venous blood glucose results, the FreeStyle Libre system had a MARD of 13.2%, while the continuous glucose monitoring systems evaluated had MARDs of 16.8% (Dexcom G4 Platinum) and 21.4% (Medtronic MiniMed 640G). It should be noted that these MARD results were obtained under controlled conditions and that the accuracy and performance of these continuous glucose monitoring systems may vary considerably during periods of rapidly changing glucose levels, such as during exercise or after meals.

The results of error grid analyses were also reported. This method describes the clinical accuracy of a test method compared with a reference method based on paired glucose samples and the clinical significance of the differences between them. Researchers found that between 62.4% and 94.0% of the paired samples of the FreeStyle Libre system test method and the reference test method were within zone A — meaning that test values deviated less than 20% from the reference value and have no clinical implications. Between 91.6% and 100% of paired samples were within zones A and B — meaning that, although the difference may be more than 20%, no treatment is indicated, or treatment would have little or no effect on clinical outcome).

There was also a high level of agreement between the FreeStyle Libre system and the other test methods, ranging from 76% agreement with continuous glucose monitoring to 95% agreement with venous blood glucose. Two studies reported the mean lag time in glucose levels associated with the FreeStyle Libre system compared with that of venous blood glucose, which ranged from 3.1 minutes to 4.5 minutes. In the study of patients undergoing cardiac surgery, it was reported that, although both systems were reliable and no complications occurred, the intravascular microdialysis continuous glucose monitoring system was more accurate than the FreeStyle Libre system (which repeatedly measured a lower glucose value).

Hypoglycemia

The primary outcome in the IMPACT study was the change in the amount of time in hypoglycemia — defined as hours per day, with a glucose level of less than 3.9 mmol/L (70 mg/dL) from baseline to six months. The average amount of time participants experienced hypoglycemia was statistically significantly lower at six months for people who used the FreeStyle Libre system (1.24 fewer hours per day) compared with those who self-monitored their blood glucose. Analyses of day and night results found that the amount of time spent below all hypoglycemic thresholds and the number of hypoglycemic episodes were also significantly reduced in patients who used the FreeStyle Libre system compared with those who self-monitored their blood glucose.

A single-arm study that prospectively assessed the impact of introducing the FreeStyle Libre system to people with type 1 diabetes over a 16-week period reported that hypoglycemic episodes, recorded and self-reported, were reduced: from 17 episodes in the first two weeks of the study to 12 episodes in the final two weeks of the study. Another single-arm study of people with type 1 or type 2 diabetes that assessed aggregate data from ambulatory glucose profiles obtained using the FreeStyle Libre system found that, despite having similar A1C values, patients with type 2 diabetes had different glycemic characteristics and a different risk of hypoglycemia than those with type 1 diabetes. It was noted that these data, which
have important implications for therapeutic adjustments, were obtained within a short time and with minimal provider and patient input using the FreeStyle Libre system.\textsuperscript{22}

In one of the studies of the FreeStyle Libre system in children with type 1 diabetes, sensor data were used to retrospectively determine glycemic variability in the children during the course of the study.\textsuperscript{27} Overall, the children were in the target glucose range of 3.9 mmol/L to 10.0 mmol/L (70 mg/dL to 180 mg/dL) approximately 50% of the time, or an average of 12.1 hours per day. This range was the same both during the day and at night, regardless of whether insulin was administered by insulin pump or by multiple daily injections.\textsuperscript{27}

**Glycated Hemoglobin**

There are conflicting reports about the effects of the FreeStyle Libre system on A1C. In the IMPACT study, although time spent in hypoglycemia was significantly reduced over the six-month period, A1C values remained essentially unchanged and were not statistically significantly different in participants who used the FreeStyle Libre system compared with those who self-monitored their blood glucose.\textsuperscript{19} In contrast, two single-arm studies without control groups reported statistically significant reductions in A1C values following the introduction of the FreeStyle Libre system.\textsuperscript{9,23}

**Patient Factors**

Two studies, one in adults\textsuperscript{21} and the other in children,\textsuperscript{27} reported that factors such as age, body weight or body mass index, sex, A1C level, method of insulin administration, and time of use (day versus night) did not affect sensor accuracy.

A subgroup analysis of the IMPACT study reported in abstract form evaluated whether younger adults (less than 25 years of age) were less likely than older adults (25 years of age and older) to regularly use glucose monitoring devices.\textsuperscript{32} After six months of use, there were no significant differences between these subgroups of participants in the number of scans, amount of sensor wear, or sensor-derived glucose results.\textsuperscript{32} In addition, there was no significant interaction between participant age and time in hypoglycemia, in keeping with the results for the overall study population.\textsuperscript{32}

The IMPACT study also reported on the average number of daily finger-stick tests performed by all participants.\textsuperscript{19} In users of the FreeStyle Libre system, testing was reduced from an average of 5.5 finger-stick tests per day during the baseline period to 0.5 tests per day (one test every two to five days) during the treatment phase, compared with 5.6 tests per day with self-monitoring of blood glucose.\textsuperscript{19} The mean number of sensor scans per day in users of the FreeStyle Libre system was 15.1 during the treatment phase.\textsuperscript{19} A recent report based on real-world use of the FreeStyle Libre system found that people using the system checked their glucose levels an average of 16 times per day and that average glucose levels, A1C levels, the time spent in hypoglycemia, and instances of hyperglycemia all decreased as scan rates increased.\textsuperscript{33}

**Quality of Life**

Various patient-reported outcomes (using instruments such as the Hypoglycemia Fear Survey, Diabetes Treatment and Satisfaction Questionnaire, Diabetes Distress Scale [DDS], and the Diabetes Quality of Life questionnaire) were reported in the IMPACT study.\textsuperscript{19} The results were inconclusive because some differences between measures were statistically significant, whereas others were not.\textsuperscript{19}

In one single-arm study, quality of life was measured using the DDS; reductions in the mean scale score, and emotional burden and regimen-related distress sub-scores, were reported 16 weeks after introducing the FreeStyle Libre system.\textsuperscript{9}
Cost-Effectiveness

Three cost-effectiveness analyses, all reported in abstract form, were identified that compared the FreeStyle Libre system with conventional self-monitoring of blood glucose, or continuous blood glucose testing, in the UK and other European countries. Overall, compared with self-monitoring or continuous blood glucose testing, the FreeStyle Libre system was cost-effective.

Safety

None of the studies reported serious device-related safety issues or malfunctions associated with the FreeStyle Libre system. In general, most of the adverse events reported for the FreeStyle Libre system were associated with the sensor and/or adhesive (e.g., local irritation, allergy events, itching, redness, rash, and insertion-site symptoms).

“…caution should be taken when the system is used with other implanted medical devices such as pacemakers, and some individuals may be sensitive to the adhesive used on the sensor.”

The clinical study that formed the basis for the FDA approval of the FreeStyle Libre Pro system found that, when the device indicated glucose values at or below 60 mg/dL (3.3 mmol/L), 40% of the time those values were actually in the range of 81 mg/dL to 160 mg/dL (4.5 mmol/L to 8.9 mmol/L). Interpreting readings should therefore be based only on trends and patterns analyzed over time, using the reports available (per the intended use).

Contraindications to the use of the FreeStyle Libre Pro system are that the sensor must be removed before having an MRI, CT scan, or high-frequency electrical heat (diathermy) treatment, as the effects of these interventions on the performance of the FreeStyle Libre Pro system have not been evaluated. Furthermore, caution should be taken when the system is used with other implanted medical devices such as pacemakers, and some individuals may be sensitive to the adhesive used on the sensor.

Concurrent Developments

Various non-invasive glucose monitors are currently in development or are awaiting regulatory approval. The GlucoTrack DF-F (Integrity Applications, Ashkelon, Israel) works in a manner similar to that of a pulse oximeter. It consists of a small sensor that is clipped to the earlobe and connected to a hand-held control and display unit. The NovioSense (Novio Tech Campus, Nijmegen, Netherlands) is a small glucose sensor placed in the eye to continuously measure glucose levels in the tears. It is anticipated that it may enter the market in 2018. The OrSense NBM-200G (OrSense, Petach Tikva, Israel) is a finger sensor that uses changes in the red and near-infrared spectrum of light passed through the finger to correlate with, and detect changes in, glucose concentrations.

Implementation Issues

The FreeStyle Libre Pro system generates a large volume of data that may be overwhelming, although presenting the data as an ambulatory glucose profile may mitigate this issue for some users.

Wearing the sensor could cause skin irritations or it could be perceived as unsightly. Patient satisfaction and sensor wear and tear are important considerations, as one study reported that continuous glucose monitoring sensors were only worn 57% of the time by adults with type 1 diabetes enrolled in an RCT to assess impaired awareness of hypoglycemia. One of the single-arm studies of the FreeStyle Libre system reported that the sensor did not record any results in five patients (6.8%) despite being in place for 14 days. In addition, 12 sensors (16.4%) had less than five days of data — either because they fell off or were removed early.

Because the FreeStyle Libre Pro sensor must be replaced every 14 days by a health care provider, this may impact the number of clinician office visits for the person using the system.

For both continuous and flash glucose monitoring systems, because measurements are taken from interstitial fluid and not blood, there is a lag time in measurement of up to 15 minutes. The lag time is longest if the glucose levels are changing rapidly (e.g., after eating, dosing of insulin, or exercising); therefore, it is still necessary to do a finger-stick and capillary blood glucose test if a change in treatment is being contemplated.
There are no reusable components in the FreeStyle Libre Pro system, other than the reader. The sensor should not be reused and is not suitable for re-sterilization. The sensor reader should be cleaned between patient use.

As there is no patient interaction with the system, patients do not require training on the calibration or use of the system. This would change if the consumer version of the FreeStyle Libre system becomes available in Canada.

With the FreeStyle Libre Pro system, the clinic or physician’s office is required to purchase only one FreeStyle Libre Pro reader, which can be used for multiple patients.

Final Remarks

Flash glucose monitoring has been described as occupying a clinical space between continuous glucose monitoring and the self-monitoring of blood glucose. In principle, flash glucose monitoring using the FreeStyle Libre Pro system could replace self-monitoring with finger sticks and capillary blood glucose testing, as it offers a more convenient and pain-free alternative, especially if frequent testing is required. It is important to note that although the need for routine finger sticks is diminished, they are still required if a change in therapy (such as an adjustment of insulin dose) is being contemplated or if hypoglycemia is suspected.

Because of the pain and discomfort of finger sticks and the need to wake up in the night to test blood glucose levels when nighttime hypoglycemia is a concern, it is especially difficult to have frequent self-monitoring in children. The FreeStyle Libre Pro system may be beneficial for children with type 1 diabetes who require intensive glucose monitoring; however, in North America, it is not currently approved for use in children.

Methods — Literature Search Strategy

A limited literature search was conducted using the following bibliographic databases: PubMed, MEDLINE, Embase, and the Cochrane Library. Grey literature searching included relevant sections of the Grey Matters checklist (https://www.cadth.ca/grey-matters). No methodological filters were applied. The search was limited to English-language documents published between January 1, 2012 and January 18, 2017. Conference abstracts published between January 1, 2015 and January 19, 2017 were included in the search results. Regular alerts updated the search until project completion; only citations retrieved before February 10, 2017 were incorporated into the analysis.
Table 1: Performance and Test Characteristics of the FreeStyle Libre Flash Glucose Monitoring System

<table>
<thead>
<tr>
<th>Study</th>
<th>Design</th>
<th>Comparator</th>
<th>MARD</th>
<th>Error Grid Analysis*</th>
<th>Other Outcomes</th>
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<tbody>
<tr>
<td>Aberer et al. (2017)</td>
<td>N = 12 adult pts with T1DM • Single-arm comparison of FSL with 2 CGM systems in same pts × 12 hours</td>
<td>CGM with Dexcom and Medtronic systems and venous BG (reference)</td>
<td>Overall: • FSL vs. venous BG: 13.2% • Dexcom vs. venous BG: 16.8% • Medtronic vs. venous BG: 21.4%</td>
<td>Percentage of overall paired results for FSL/Dexcom/Medtronic in: • zone A: 85.7%/83.6%/71.1% • zones A and B: 100%/99.3%/98.6%</td>
<td>Reported MARD after exercise, hypoglycemia, euglycemia, and hyperglycemia; and all sensors performed less accurately during hypoglycemia and best during hyperglycemia</td>
<td>• Venous BG levels were measured by laboratory analyzer every 5 minutes × 12 hours • Funding by the European Commission</td>
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<td>Bailey et al. (2015)</td>
<td>N = 72 adult pts with T1DM or T2DM • Single-arm comparison of FSL and CBG in same pts × 14 days</td>
<td>CBG testing and venous BG (reference)</td>
<td>FSL vs. CBG: 11.4% • FSL vs. venous BG: 12%</td>
<td>Percentage of paired results for FSL/CBG in: • zone A: 86.7%/85.5% • zones A and B: 99.7%/99.0%</td>
<td>Mean lag time between FSL and venous BG was 4.5 min. ± 4.8 min. (r² = 0.95)</td>
<td>• Venous BG levels were measured by laboratory analyzer during 3 in-clinic visits • Funding by Abbott Diabetes Care</td>
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<tr>
<td>Bonora et al. (2016)</td>
<td>N = 8 adult pts with T1DM • Single-arm comparison of FSL and CGM in same pts × 14 days</td>
<td>CGM and SMBG</td>
<td>FSL vs. CGM: 18.1% ± 14.8% • FSL vs. SMBG: 16.6% ± 11.6%</td>
<td>Percentage of paired results for FSL/CGM in: • zone A: 62.4% • zones A and B: 91.6%</td>
<td>• FSL and CGM: r² = 0.76 • FSL and SMBG: r² = 0.86</td>
<td>• Funding by the University of Padova, Padua, Italy</td>
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<tr>
<td>Ji et al. (2017)</td>
<td>N = 45 adult pts with T1DM or T2DM • Single-arm comparison of FSL and CBG in same pts × 14 days</td>
<td>CBG testing and venous BG (reference)</td>
<td>FSL vs. CBG: 10.0% • FSL vs. venous BG: 10.7%</td>
<td>Percentage of paired results for FSL/CBG in: • zone A: 87.0%/88.8% • zones A and B: 99.9%/99.5%</td>
<td>Mean lag time between FSL and venous BG was 3.1 min (95% CI: 2.5 to 4.3) • FSL and CBG: r² = 0.94 • FSL and venous BG: r² = 0.95</td>
<td>• Venous BG levels were measured by laboratory analyzer during 3 in-clinic visits • Funding by Abbott Diabetes Care</td>
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<tr>
<td>Schierenbeck et al. (2016)</td>
<td>N = 26 adult pts undergoing cardiac surgery • Single-arm comparison of FSL and MD-CGM in same pts over mean of 23.7 hours</td>
<td>MD-CGM</td>
<td>MD-CGM range: 2.5% to 12.6% • FSL range: 12.0% to 52.1%</td>
<td>Percentage of paired results in: • zone A: 94% • zones A and B: 100%</td>
<td>Both systems followed the trend of true BG (as per arterial blood gas analysis) well, but FSL repeatedly underestimated the BG value</td>
<td>Funding by Mats Kleberg Foundation and the Signe and Olof Wallenius Foundation</td>
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<tr>
<td>Edge et al. (2017)</td>
<td>N = 89 children (4 to 17 years) with T1DM</td>
<td>CBG testing</td>
<td>FSL vs. CBG: 13.9%</td>
<td>Percentage of paired results in: • zone A: 83.8% • zones A and B: 99.4%</td>
<td>Pts in the target BG range approximately 50% of the time (mean 12.1 hours/day)</td>
<td>Funding by Abbott Diabetes Care</td>
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</table>

BG = blood glucose; CBG = capillary blood glucose; CGM = continuous blood glucose monitoring; CI = confidence interval; Dexcom = Dexcom G4 Platinum continuous glucose monitoring system; FSL = FreeStyle Libre system; MARD = mean absolute relative difference; MD-CGM = intravascular microdialysis continuous glucose monitoring; Medtronic = Medtronic MiniMed 640G insulin pump; min. = minutes; pts = patients; SMBG = self-monitoring of blood glucose; T1DM/T2DM = type 1 and type 2 diabetes mellitus; vs. = versus.

* Error grid analysis pairs glucose samples in five zones (A, B, C, D, and E). Values in zone A are within 20% of the reference value and have no clinical implications. Values in zone B exceed a 20% difference from the reference value, but require no treatment or only benign treatment that will not change clinical outcomes. Therefore, the more values in zones A and B, the higher the clinical accuracy of the test method relative to the reference method.
References


