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Article type : Review

Continuous glucose monitoring: A review of the evidence in type 1 and 2 diabetes mellitus

Short Title: Continuous glucose monitoring: A review

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Abstract word count: 247/250

Text word count: 4948/5000

Tables: 4

Figures: 1

Disclosures: The authors SJ and JJ have type 1 diabetes and have used continuous glucose monitoring technology. The other authors have no conflicting interests to declare.

This is the author manuscript accepted for publication and has undergone full peer review but has not been through the copyediting, typesetting, pagination and proofreading process, which may lead to differences between this version and the [Version of Record](#). Please cite this article as [doi: 10.1111/DME.14528](https://doi.org/10.1111/DME.14528)

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LIST OF STUDY ABBREVIATIONS

- DCCT = Diabetes Control and Complications Trial
- GOLD = Glycaemic Control and Optimisation of Life Quality in Type 1 Diabetes
- DIAMOND = Multiple Daily Injections and Continuous Glucose Monitoring in Diabetes
- SWITCH = Sensing with Insulin Pump Therapy to Control HbA1c
- JDRF = Juvenile Diabetes Research Foundation
- CITY = CGM Intervention in Teens and Young Adults with Type 1 Diabetes
- WISDM = Wireless Innovation for Seniors with Diabetes Mellitus
- HypoDE = Real-Time Continuous Glucose Monitoring in Patients with Type 1 Diabetes at High Risk for Low Glucose Values Using Multiple Daily Injections in Germany
- IN CONTROL = Continuous Glucose Monitoring for Patients with Type 1 Diabetes and Impaired Awareness of Hypoglycaemia
- IMPACT = Novel Glucose-Sensing Technology and Hypoglycaemia in Type 1 Diabetes
- SELFY = FreeStyle Libre Glucose Monitoring System Paediatric Study
- REPLACE = An Evaluation of a Novel Glucose Sensing Technology in Type 2 Diabetes
- GP-OSMOTIC = General Practice Optimising Structured Monitoring to Improve Clinical Outcomes in Type 2 Diabetes

WHAT'S NEW?

What is already known?

- Currently, standard practice in diabetes management involves self-monitoring of blood glucose and measurement of glycated haemoglobin.

What has this study found?

- 70 • Continuous glucose monitoring can offer added benefits to standard care, especially in type 1
71 diabetes, including improvements in:
72 ○ Glycaemic management;
73 ○ Quality of life in people with diabetes and parents of children with diabetes; and
74 ○ Clinician experience.

75 What are the clinical implications of this study?

- 76 • Continuous glucose monitoring is a clinically effective and cost-effective adjunct to diabetes
77 management, and may improve diabetes outcomes.

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80 ACKNOWLEDGMENTS

81 We would like to thank Ms Michele Gaca, Chief Librarian at Health Sciences Library, Austin Health,
82 for her assistance with literature searching.

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92 ABSTRACT

93 **Context and Aim:** Continuous glucose monitoring is becoming widely accepted as an adjunct to
94 diabetes management. Compared to standard care, continuous glucose monitoring can provide detailed
95 information about glycaemic variability in an internationally standardised ambulatory glucose profile,
96 enabling more informed user and clinician decision-making. We aimed to review the evidence, user
97 experience, and cost-effectiveness of continuous glucose monitoring.

98
99 **Methods:** A literature search was conducted by combining subject headings “continuous glucose
100 monitoring” and “flash glucose monitoring”, with key words “type 1 diabetes” and “type 2 diabetes”,
101 limited to “1999 to current”. Further evidence was obtained from relevant references of retrieved articles.

102
103 **Results:** There is strong evidence for continuous glucose monitoring use in people with type 1 diabetes,
104 with benefits of reduced glycated haemoglobin and hypoglycaemia, and increased time in range. While
105 the evidence for continuous glucose monitoring use in type 2 diabetes is less robust, similar benefits
106 have been demonstrated. Continuous glucose monitoring can improve diabetes-related satisfaction in

107 people with diabetes and parents of children with diabetes, as well as the clinician experience. However,
108 continuous glucose monitoring does have limitations including cost, accuracy and perceived
109 inconvenience. Cost-effectiveness analyses have indicated that continuous glucose monitoring is a cost-
110 effective adjunct to type 1 diabetes management that is associated with reduced diabetes-related
111 complications and hospitalisation.

112

113 **Conclusions:** Continuous glucose monitoring is revolutionising diabetes management. It is a cost-
114 effective adjunct to diabetes management that has the potential to improve glycaemic outcomes and
115 quality of life in people with diabetes, especially type 1 diabetes.

116

117 **Key Words:** blood glucose self-monitoring, health technology, type 1 diabetes, type 2 diabetes

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120 **INTRODUCTION**

121 It has been three decades since the landmark DCCT trial demonstrated the importance of glycaemic
122 management in the prevention or delay of microvascular complications (1). Glycated haemoglobin
123 (HbA1c) has long been the gold standard for assessing long-term glycaemic management (2). However,
124 HbA1c does not provide information about glucose trends and excursions; individuals with the same
125 HbA1c can have significantly different diurnal glucose variation (3). Therefore, self-monitoring of
126 blood glucose (SMBG) is also recommended in select individuals, such as those receiving insulin
127 therapy (2). Structured SMBG can distinguish between fasting and post-prandial hyperglycaemia,
128 uncover glycaemic variability, and allow people with diabetes (PWD) to intervene promptly to
129 minimise dysglycaemia (3,4). In individuals receiving insulin, international guidelines recommend
130 SMBG several times per day (e.g. before meals, at bedtime, before exercise, when hypoglycaemia is
131 suspected) (2,5,6), although factors including pain, cost, inconvenience, and fear of needles often lead
132 to non-adherence. It has been estimated that only 44% of people with type 1 diabetes (T1DM) and 24%
133 of people with type 2 diabetes (T2DM) routinely perform SMBG as per guidelines (7).

134

135 Continuous glucose monitoring (CGM) is a minimally invasive modality of monitoring glucose levels.
136 Unlike SMBG, which provides discrete snapshots of glucose levels, CGM records glucose levels
137 continuously (8). Despite first becoming available in 1999, CGM use has only really increased during
138 recent years, due to wider availability, evidence and subsidised funding (9-11). For example,
139 recognising the increasing demand for CGM in PWD, the Australian Government recently announced
140 an AUD\$300 million CGM initiative. Since 1 March 2020, eligible Australians have been able to
141 receive fully subsidised CGM products (12). Global CGM usage is expected to further increase, with
142 recent international guidelines recommending that CGM be considered in all children and adolescents
143 with T1DM (5).

144

145 This review aims to review the literature on CGM in terms of efficacy, user experience, and cost-
146 effectiveness. Medline and Embase searches were conducted by combining subject headings
147 “continuous glucose monitoring” and “flash glucose monitoring”, with key words “type 1 diabetes” and
148 “type 2 diabetes”, limited to “1999 to current”. Clinical trials and systematic reviews describing CGM
149 use in T1DM and T2DM were included. Articles describing CGM use in gestational diabetes, use of
150 CGM as a tool for assessing other diabetes interventions, or comparing CGM with interventions other
151 than standard care were excluded. Further evidence was obtained from relevant references of retrieved
152 articles.

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155 **WHAT IS CONTINUOUS GLUCOSE MONITORING?**

156 CGM allows continuous monitoring of glucose levels via a tiny electrochemical sensor electrode
157 inserted under the skin (8). The sensor is connected to a transmitter, which sends this information to a
158 detector, such as a smartphone, or a continuous subcutaneous insulin infusion (CSII) device. CGM
159 devices are of two types: traditional CGM and flash glucose monitoring (FGM), also known as
160 intermittently scanned CGM. Both systems provide information about current and previous glucose
161 levels, glucose trends, and anticipated future glycaemic status (13). A main difference is that CGM
162 passively transmits this information continuously to the reader without user engagement, whereas FGM
163 only provides this information when the user scans the sensor (14). A summary of available CGM
164 devices is provided in Table 1.

165

166 **Traditional continuous glucose monitoring**

167 CGM can be further categorised into personal or “real-time” use, and professional or “retrospective”
168 use (15). Personal CGM allows PWD to access continuous glucose data and intervene as required in
169 real-time, whereas with the less commonly used professional CGM the data are blinded to PWD and
170 only accessible retrospectively by clinicians. Some personal CGM devices, such as the Dexcom G5™
171 Mobile, Dexcom G6™ Mobile and Medtronic Guardian™ Connect, allow remote monitoring, where up
172 to 5 individuals are able to view CGM readings and receive alerts in real-time (10,16). Most
173 transcutaneous sensors have a wear life of 7-10 days. An alternative is Eversense™, an implantable
174 real-time CGM device which is approved for up to 6 months use in Europe and 3 months use in America
175 (10).

176

177 CGM devices measure interstitial glucose levels. Although generally comparable, blood and interstitial
178 glucose readings can differ by 10-20% (8). To maintain accuracy of readings, older CGM devices
179 require calibration multiple times per day. Calibration is not required with newer devices such as

180 Dexcom G6™, unless the difference between CGM readings and blood sugar levels (BSL) are
181 consistently >20% (17). SMBG is also recommended before treating hypo- or hyperglycaemia (15).
182 Studies have shown that using CGM to guide insulin dosing without confirmatory SMBG is safe (18),
183 although currently only Dexcom G5™ Mobile and Dexcom G6™ are approved for insulin dosing by
184 the Therapeutic Goods Administration (TGA) and Food and Drug Administration (FDA) (5,19).
185 Eversense™ is also FDA-approved for insulin dosing (20). CGM devices can also provide alarms during
186 times of actual or impending dysglycaemia, which can be particularly useful in PWD prone to
187 hypoglycaemia, and those with hypoglycaemia unawareness (11).

188
189 Most CGM devices are compatible with CSII pumps. Some devices can also lead to alteration of insulin
190 administration. The Medtronic Guardian™ □ 2 Link system (Enlite™ □ sensor, Guardian™ Link 2
191 transmitter and MiniMed™ □ 640G pump), can suspend insulin delivery either when glucose levels
192 reach a pre-specified low-glucose threshold, or when glucose trends predict hypoglycaemia beyond the
193 threshold within the next 30 minutes (21). The Medtronic MiniMed™ □ 670G system (Guardian™ □
194 Sensor 3, Guardian™ □ Link 3 transmitter and MiniMed™ □ 670G pump) utilises hybrid closed-loop
195 technology to automatically increase, decrease or suspend insulin delivery to maintain a target glucose
196 level of 6.7mmol/L (21). Real-world data indicate this technology results in increased time in range
197 (TIR), reduced hypoglycaemia, and increased user quality of life (QOL) (10,22).

198 199 **Flash glucose monitoring**

200 FGM is a relatively newer CGM technology, first becoming available in 2016. In contrast to traditional
201 CGM, FGM information is only available when the user accesses it. PWD can flash a reader over the
202 sensor to obtain glucose information including: current glucose level (within the last minute), glucose
203 trend, and results from the past 8 hours or up to the last scan, if scanned <8 hours ago (3). To maintain
204 adequate data collection, PWD must scan the sensor at least every 8 hours, otherwise data will be lost
205 and the report will show data gaps (13). The FreeStyle™ Libre Pro is a retrospective FGM device,
206 which does not require any action by the user, including sensor scanning. The data provided for
207 retrospective analysis are similar to that provided by traditional CGM.

208
209 FGM is pre-calibrated in the factory, requiring no further calibration throughout use (23). One FGM
210 device is approved by the TGA, the FreeStyle™ Libre. The FreeStyle™ Libre 2 is currently approved
211 for use in America and Europe (24). Both devices are FDA-approved for insulin dosing without SMBG
212 (21, 25). The only situations where SMBG is recommended to make therapeutic decisions when using
213 FGM include: when symptoms do not match the glucose level detected, hypoglycaemia needs to be
214 confirmed, and when glucose levels are changing rapidly (13).

215

216 The FreeStyle Libre™ does not provide alarms (3). The FreeStyle Libre™ 2 offers optional
217 customisable alarms for hypo- and hyperglycaemia (26). In contrast to traditional CGM which alarm
218 every minute if dysglycaemia is sustained, the FreeStyle Libre™ 2 does not alarm again once the alarm
219 is confirmed, until the system resets when euglycaemia is re-established. FGM may be more suited to
220 PWD who find alarms intrusive, people who do not perform SMBG often, and people who are unable
221 or unwilling to perform regular SMBG for calibration. FGM is also cheaper than CGM, probably related
222 to a longer wear life of 14 days (14).

223
224

225 **AMBULATORY GLUCOSE PROFILE**

226 Traditionally, different CGM devices have required use of unique software to report glycaemic control,
227 rendering analysis and interpretation time-consuming and difficult. Nowadays, an internationally
228 standardised report, called the ambulatory glucose profile (AGP), is commonly utilised (Fig. 1). The
229 AGP is a single-page report with statistical and graphic information organised into 5 major components:
230 1) data completeness captured by sensor; 2) glucose level statistics (e.g. hypoglycaemia, TIR); 3)
231 glucose profile based on a “model day” (also called the AGP); 4) glucose management indicator; and
232 5) daily glucose profiles (13,27,28). CGM data can be provided from the previous 5 days to 3 months.
233 Data are collated to produce a glucose profile as if all the readings had occurred in a single 24-hour
234 period (the “model day”) (13). The AGP enables retrospective analysis of CGM data, allowing PWD
235 and clinicians to characterise diurnal glucose patterns to inform management (11,29).

236

237 International guidelines have been developed to aid interpretation of the AGP. The primary goal of
238 diabetes management is to increase TIR while reducing hypoglycaemia (5,11). Increased TIR is
239 associated with slowed progression of complications such as diabetic retinopathy (30,31) and
240 microalbuminuria (31), and reductions in HbA1c (32). For individuals with T1DM and T2DM, targets
241 of >70% TIR and <4% hypoglycaemia are recommended (11). Guidelines also recommend a glycaemic
242 variability target (% coefficient of variation [CV]) of ≤36% (11), although studies have shown that a
243 lower %CV target of <33% provides additional protection against hypoglycaemia for PWD on insulin
244 or sulfonylureas (33). The AGP facilitates easy identification of these parameters, allowing safe and
245 effective glucose management.

246

247 There are several practical challenges associated with the AGP. Healthcare practices must be equipped
248 with adequate technology systems, and be able to link these systems to the appropriate software at the
249 time of the consultation to access CGM data (27). Furthermore, data need to be near-complete to allow
250 accurate interpretation. The timing of activities such as meals and exercise may vary between days, and
251 glucose variability between days may be diluted in summary data, which combined with the time

252 constraints in clinical settings, can make interpretation challenging (28). However, with apt education
253 on AGP interpretation, clinicians are able to make informed treatment decisions to improve diabetes
254 management (34).

255

256

257 **EVIDENCE FOR CONTINUOUS GLUCOSE MONITORING**

258 Recently, there has been an increasing number of trials demonstrating the efficacy of CGM and FGM
259 in improving diabetes-related outcomes. Notably, many have demonstrated a HbA1c reduction,
260 increased TIR (3.9-10.0mmol/L) and reduced hypoglycaemia (<3.9mmol/L). Currently, the majority of
261 the evidence supports CGM and FGM use in T1DM. Findings involving people with T2DM on insulin
262 are also available in the literature, while studies involving people on oral hypoglycaemics are scarce.
263 Tables 2-3 contain summaries of randomised controlled trials (RCTs) on CGM and FGM.

264

265 Maiorino et al conducted a meta-analysis of RCTs assessing the efficacy of CGM or FGM in people
266 with T1DM or T2DM. They found an overall weighted mean reduction in HbA1c of 2mmol/mol (0.2%;
267 $p=0.003$), and an overall increase in TIR of 70.74 min/day (95%CI 46.73–94.76; $p<0.001$) (S3). Time
268 in level 1 hypoglycaemia (3.0mmol/L–3.9mmol/L) reduced by 27.16min/day (95%CI -42.08 to -12.25;
269 $p<0.001$), whereas time in level 2 hypoglycaemia (<3.0mmol/L) reduced by 13.58min/day (95%CI -
270 20.63 to -6.53; $p<0.001$) (S3).

271

272

273 **Traditional Continuous Glucose Monitoring**

274

275 Type 1 Diabetes

276

277 *HbA1c*

278 There is strong evidence for CGM use in people with T1DM. RCTs including the GOLD, DIAMOND,
279 SWITCH, CITY and WISDM studies all reported significant HbA1c reductions with CGM use in
280 children and adults with T1DM (37-39,41,42). A meta-analysis reported that CGM was associated with
281 a significantly lower HbA1c at end-point compared to SMBG, with a between-group difference of -
282 2mmol/mol (-0.2%) (S4). The JDRF study stratified the study population according to age and reported
283 a mean between-group difference in HbA1c of -6mmol/mol (-0.5%; $p<0.001$) in adults (≥ 25 years), but
284 no significant change in children (8–14 years) or adolescents (15–24 years). In another meta-analysis,
285 CGM was associated with a HbA1c reduction of 3mmol/mol (0.3%; $p=0.004$), although these changes
286 were only significant in people aged >15 years (S5). The benefits of CGM are primarily seen in PWD
287 with near daily use (39,S6). This age-related phenomenon could potentially be explained by reduced
288 sensor use in children and adolescents, as observed in the JDRF study (40). A RCT by Cosson et al

289 which involved two discrete 48-hour CGM wears 3 months apart, reported no significant difference in
290 HbA1c change between the CGM and control groups (45), which further emphasises the importance of
291 daily CGM use.

292

293 *Time in Range and Hypoglycaemia*

294 The DIAMOND, SWITCH, CITY and WISDM RCTs all reported an increased mean TIR of 1.3–2.4
295 hours/day (38,39,41,42). When stratified according to age, the JDRF study demonstrated a significant
296 increase in TIR in adults, but no significant difference in children and adolescents (40). Reduced time
297 in hypoglycaemia and reduced incidence of severe hypoglycaemic events were reported in the GOLD,
298 DIAMOND, SWITCH, CITY and WISDM studies (37-39,41,42). The DIAMOND study specifically
299 noted reduced nocturnal hypoglycaemia in the CGM group (38).

300

301 CGM is beneficial in people with hypoglycaemia unawareness. The HypoDE and IN CONTROL
302 studies involved people with T1DM and hypoglycaemia unawareness, with the primary outcome of
303 time spent in hypoglycaemia. The HypoDE RCT reported a median hypoglycaemia duration of 23.9
304 mins/day (95%CI 12.9–54.5) in the CGM group, compared to 92.2 mins/day (95%CI 51.8–172.6) in
305 the control group ($p<0.0001$) (35). The IN CONTROL crossover study showed a reduction in mean
306 time in hypoglycaemia of 1.1 hours/day (95%CI -1.4 to -0.8; $p<0.0001$) in the CGM phase, compared
307 to the SMBG phase (36). The study also reported a between-group difference in mean TIR of 2.3
308 hours/day (95%CI 1.9–2.7; $p<0.0001$), favouring the CGM group (36). No significant difference in TIR
309 was reported in the HypoDE study (35). No significant HbA1c change was reported in either of these
310 studies (35,36).

311

312 Type 2 Diabetes

313 There are a handful of RCTs involving CGM use in T2DM, all of which have demonstrated a HbA1c
314 reduction (43-45). A meta-analysis comparing CGM to SMBG found a pooled mean HbA1c reduction
315 of 3mmol/mol (0.3%; $p=0.01$) (S7). Similarly, another meta-analysis reported a HbA1c reduction of
316 2mmol/mol (0.2%) (S8). Ehrhardt et al observed that a high baseline HbA1c was associated with a
317 greater HbA1c reduction (44). When considering glycaemic variability, Beck et al reported an increase
318 in median TIR from 802 min/day to 882 min/day in the CGM group, compared to an increase from 794
319 min/day to 836 min/day in the control group (43). Cosson et al reported no significant difference (45).
320 No significant effect on hypoglycaemia was observed in any of these studies (43,45,S7), probably due
321 to low levels of hypoglycaemia at baseline in people with T2DM (43).

322

323 **Flash Glucose Monitoring**

324

325 Type 1 Diabetes

326

327 *HbA1c*

328 The efficacy of FGM in T1DM has been assessed in three studies: IMPACT, SELFY and Tyndall et al,
329 all of which compared FGM to SMBG. The IMPACT RCT recruited adults with T1DM and HbA1c
330 ≤ 59 mmol/mol (7.5%), who reported SMBG ≥ 3 times/day. The investigators found that FGM had no
331 significant effect on HbA1c, although it is important to note that this study was designed for the effect
332 of FGM on hypoglycaemia as the primary endpoint (46). In contrast, the SELFY single arm pre-post
333 study in children and adolescents (mean baseline HbA1c 63mmol/mol [7.9%]) reported a HbA1c
334 reduction of 4mmol/mol (0.4%; $p < 0.0001$) from baseline to study end (47), with similar results seen in
335 Tyndall et al, a real-world prospective observational study (48). Tyndall et al noted that individuals
336 with a higher baseline HbA1c were more likely to achieve HbA1c reductions following FGM
337 commencement (48), which could explain why no HbA1c reduction was seen in the IMPACT trial.

338

339 *Time in Range and Hypoglycaemia*

340 The IMPACT and SELFY studies reported increased TIR of 1.0 hours/day (95%CI 0.41–1.59;
341 $p = 0.0006$) and 0.9 hours/day (95%CI 0.27–1.53; $p = 0.005$) respectively, in the FGM group compared
342 to control (46,47). The impact of FGM on hypoglycaemia was variable. The IMPACT trial reported
343 reduced mean time in hypoglycaemia by 1.24 hours/day (95%CI -1.71 to -0.77; $p < 0.0001$), as well as
344 a 39.8% reduction in nocturnal hypoglycaemia ($p < 0.0001$) in the FGM group (46). The SELFY study
345 observed that FGM had no impact on hypoglycaemia, although baseline time in hypoglycaemia was
346 low in this population (47). In contrast, based on questionnaire data, Tyndall et al found that
347 hypoglycaemia (< 3.5 mmol/L) was increased in individuals following FGM commencement (48). This
348 was attributed to the revealing of previously unrecognised hypoglycaemia by FGM data, rather than the
349 development of increased hypoglycaemia (48).

350

351 Type 2 Diabetes

352

353 *HbA1c*

354 Several studies have assessed FGM use in people with T2DM on insulin therapy. The REPLACE RCT
355 analysed the effect of FGM compared to SMBG for 6 months in people with T2DM on intensive insulin
356 therapy, and found no between-group difference in HbA1c change (49). However, subgroup analysis
357 found that in people < 65 years, FGM led to a greater HbA1c reduction compared to controls ($p = 0.03$),
358 whereas in people ≥ 65 years, the HbA1c reduction was greater in controls compared to the FGM group
359 ($p = 0.008$) (49). They hypothesised that the convenience of sensor scanning resulted in more frequent
360 glucose readings in younger participants (49), a population which is often less adherent to SMBG (S9).
361 Yaron et al analysed the effect of FGM compared to SMBG for 10 weeks in people with T2DM on
362 MDI, and reported a significant HbA1c reduction in the FGM group (S1). In the REPLACE trial, the

363 mean self-reported SMBG frequency was 3.6 times/day in the intervention group and 3.9 times/day in
364 the control group, whereas this was not specified in Yaron et al. These contrasting findings may indicate
365 that FGM is most effective in PWD who do not perform SMBG regularly.

366

367 Wada et al compared FGM with SMBG in people with non-insulin treated T2DM over 24 weeks in a
368 RCT. They reported a HbA1c reduction of 3mmol/mol (0.3%; $p=0.022$) favouring the FGM group at
369 study end, although there was no significant change in HbA1c at 12 weeks (S2). This suggests that the
370 benefits of FGM are less obvious in people with non-insulin treated T2DM, and are only realised with
371 a prolonged period of regular use.

372

373 *Time in Range and Hypoglycaemia*

374 The REPLACE RCT found that time in hypoglycaemia reduced by 0.47 hours/day (95%CI 0.21–0.72;
375 $p=0.0006$) in the FGM group compared to control, where nocturnal hypoglycaemia reduced by 0.29
376 hours (95%CI -0.45 to -0.13; $p=0.0001$), although there was no significant difference in TIR (49). Haak
377 et al analysed the effects of FGM in the intervention group from the REPLACE trial for a further 6
378 months, and yielded similar results (50). In contrast, Wada et al reported increased TIR of 2.36
379 hours/day (95%CI 1.21–3.51; $p<0.001$) in the FGM group compared to SMBG, with no significant
380 difference in time in hypoglycaemia (S2). Yaron et al found that there was no difference in frequency
381 of hypoglycaemic events in the FGM group compared to control (S1).

382

383 *Retrospective FGM*

384 Ajjan et al involved people on insulin therapy, and the GP-OSMOTIC RCT involved people on insulin
385 therapy or ≥ 2 oral hypoglycaemic agents (S10,S11). Both studies reported HbA1c reductions. Ajjan et
386 al reported a between-group difference in HbA1c change of -5mmol/mol (-0.5%; $p=0.004$) favouring
387 FGM, with no significant effect of professional FGM on hypoglycaemia (S10). In contrast, the GP-
388 OSMOTIC trial found a between-group difference in HbA1c change of -5mmol/mol (-0.5%; $p=0.0001$)
389 at 6 months, although these observations were not sustained at 12 months, with no significant between-
390 group difference (S11). In terms of glucose variability, the GP-OSMOTIC trial reported that the mean
391 percentage TIR at 12 months was 7.9% (95%CI 2.3–13.5; $p=0.006$) higher in the FGM group compared
392 to control, whereas Ajjan et al reported no significant difference (S10,S11).

393

394

395 **USER EXPERIENCE**

396

397 **People with Diabetes**

398 Many studies have demonstrated that CGM user satisfaction is high, and that CGM contributed to
399 improved diabetes-related QOL compared to SMBG (35-38,43,46,S1,S10,S12,S13). In a systematic

400 review of FGM, all RCTs found improvements in diabetes treatment satisfaction, diabetes QOL, and
401 diabetes-related stress (S12). Reduced fear of hypoglycaemia was reported in multiple studies
402 (36,37,S14,S15), perhaps because CGM allows PWD to easily and quickly identify and respond to
403 dysglycaemia, thereby allowing individuals to regain a sense of empowerment over glucose
404 management, and more broadly, their diabetes (S16). CGM can also allow PWD to better understand
405 the impact of food and exercise on glucose management, enabling short-term lifestyle planning such as
406 changing the timing of a meal or reducing the length of a run to avoid dysglycaemia (S7,S15,S17). A
407 recent position statement recommended specific target glucose levels and corrective actions before,
408 during and after exercise for PWD using CGM, depending on their risk of hypoglycaemia (S18). CGM
409 readings and BSL may differ during exercise, therefore the CGM reading needs to be interpreted
410 together with the trend arrow (S18). Lawton et al reported that while participants expressed confidence
411 in adjusting insulin and lifestyle to address impending dysglycaemia, most needed clinicians to interpret
412 retrospective CGM data and determine changes to diabetes treatment (S15).

413

414 Driving

415 CGM can potentially increase safety when driving for PWD. Driving simulator studies have
416 demonstrated that driving performance deteriorates during hypoglycaemia (S19). In Australia and
417 Europe, guidelines require that glucose level is checked ≤ 2 hours before driving to ensure glucose level
418 $>5\text{mmol/L}$, then every 2 hours thereafter to ensure levels remain $>5\text{mmol/L}$ (S20-S22). Similarly,
419 American guidelines recommend that an extended drive not be initiated with low-normal glucose level
420 (3.9-5.0mmol/L) without prophylactic carbohydrate consumption, and that glucose levels be checked
421 at regular intervals when driving for >1 hour (S23). CGM allows PWD to effortlessly determine current
422 glucose level and predictive trends, to ensure normoglycaemia is maintained. Glucose data can even be
423 shared to the car dashboard via Apple CarPlay or Android Auto, although there are no studies evaluating
424 this technology. The alarm capacity for current or impending hypoglycaemia can add another layer of
425 safety.

426

427 **Parents**

428 Interviews with parents of children with diabetes revealed that CGM with remote monitoring has
429 enabled parents to pre-empt and prevent dysglycaemia in a timely manner (S15). Some parents noted
430 the capacity to examine night-time readings and readings when children were at school offered peace
431 of mind (S15). One study reported that parental hypoglycaemia fear scores were lower when their
432 children were using CGM with remote monitoring, and that parental health-related QOL, family
433 functioning, anxiety and parental sleep measures also improved significantly (16).

434

435 **Clinicians**

436 Healthcare professionals are vital in helping PWD adopt new technology (S24). Clinicians have
437 reported that CGM data supported effective communication with PWD (29,S10), and that the AGPs
438 were easy to read and understand, enabling them to make informed decisions on therapy adjustments
439 (S10,S17). However, some clinicians find CGM adoption difficult. Barriers include limited time in
440 clinic to download data and explain glycaemic trends to PWD, and limited time to learn about different
441 CGM devices and new developments in CGM, with inconsistent resources for training (3,13,S25).
442 Moreover, guidelines dictate that PWD using CGM must receive adequate education, training and
443 support to ensure adherence and help achieve glycaemic goals (5,S26). Tanenbaum et al found that
444 clinicians who had more time with PWD, were younger, and found it easy to keep up with technological
445 advances were more likely to recommend CGM to PWD (S25).

446

447 CGM in Telemedicine

448 Following the outbreak of COVID-19, telemedicine has replaced majority of face-to-face appointments.
449 There is evidence that supports the use of telemedicine in diabetes with benefits in glycaemic
450 management (S27,S28) and satisfaction in PWD (S29) compared to conventional practice. CGM can
451 potentially further improve this experience for clinicians and PWD, by remotely providing a wealth of
452 data. However, studies in this area are lacking.

453

454

455 DISADVANTAGES

456

457 **Cost**

458 CGM devices are costly, with inconsistent reimbursement across government bodies (3,8,14). Many
459 countries, including Australia and America, offer reimbursement for people with T1DM, with limited
460 subsidisation for people with T2DM (S30,S31). Germany reimburses real-time CGM for all types of
461 diabetes, whereas Spain offers no reimbursement at all (S30). CGM systems require sensor changes
462 every 6-14 days, which costs AU\$3000-\$6000 per year without subsidisation (21). Table 4 contains a
463 summary of the cost of CGM devices in Australia.

464

465 **Accuracy**

466 Multiple factors affect CGM device accuracy. CGM measures interstitial glucose levels, which
467 compared to BSLs have a mean absolute relative difference of 10-20%. Accuracy reduces at the
468 extremes of glycaemia and during times of rapid glucose change, especially during exercise (8,14,S32-
469 S34). Therefore, many CGM devices require calibrations to maintain reliable correlation between CGM
470 readings and BSL, and infrequent or incorrect calibration could potentially reduce the accuracy of CGM
471 (13). FGM and newer CGM devices are factory-calibrated so do not have the same issue. There is also

472 a physiological lag of 5-10 minutes between blood and interstitial glucose concentrations, which could
473 result in inappropriate or excessive correction of dysglycaemia (8,13).

474

475 Certain chemicals can interfere with the accuracy of readings, including medications such as
476 paracetamol, ibuprofen, lisinopril, and vitamin C; and endogenous substances such as bilirubin,
477 cholesterol and creatinine (8,S35,S36). Paracetamol in particular is known to falsely elevate interstitial
478 glucose readings, due to interference with the electrochemical reaction that occurs during CGM sensing
479 (S37,S38). One study reported that, following oral administration of 1g paracetamol, glucose reading
480 increased by 1.6mmol/L with the Dexcom G4™ Platinum compared to BSL. In comparison, The
481 Dexcom Seven Plus™, an older sensor no longer available, saw a 10.0mmol/L increase (S39). With
482 technological improvements, the issue of chemical interference has improved dramatically, with a
483 recent study showing no significant paracetamol interference with the Dexcom G6™ (S40). FGM
484 devices also avoid paracetamol interference (S39).

485

486 **Inconvenience**

487 PWD are prone to “alarm fatigue”. Alarms can result in interrupted sleep and/or unwelcomed
488 distractions at school or work, which can lead to silencing of the alarm function and disengagement
489 with CGM (13,S15). Allergic reactions to adhesive materials used to keep CGM sensors in place can
490 also occur. In the IMPACT trial, 10 out of 328 participants reported local allergic reactions or insertion
491 site symptoms (46). Further, CGM devices are not compatible with magnetic resonance imaging and
492 need to be removed prior.

493

494 **COST-EFFECTIVENESS**

495 CGM systems are expensive, although with the potential benefits of improved glycaemic management
496 and reduced diabetes-related complications, CGM may be a cost-effective adjunct to diabetes
497 management. Studies have demonstrated that CGM use is associated with reduced hospitalisation for
498 hypoglycaemia and diabetic ketoacidosis (DKA), as well as reduced diabetes-related work absenteeism
499 (48,S14,S41,S42). In a study by Charleer et al, initiation of reimbursed CGM in 515 adults with T1DM
500 resulted in a 12% reduction in hospitalisation with reduced admission days compared to the year prior,
501 equating to a nationwide saving of EUR€345,509 (AU\$563,460) over the study period of 27 months
502 (S14). Although cost-effectiveness analyses of CGM have yielded varying results, studies have
503 commented that the incremental cost-effectiveness ratio (ICER) of CGM can be significantly improved
504 with reduced CGM cost, increased HbA1c reduction, and increased adherence to CGM use (S31,S43).

505

506 **Type 1 Diabetes**

507 Wan et al performed cost-effectiveness analysis on the DIAMOND trial population, and found that
508 during the 6-month trial, compared with SMBG, CGM increased costs (US\$11,032 [AU\$15,882] vs.

509 US\$7,236 [AU\$10,417]) without immediately improving QOL (S43). However, in predictive lifetime
510 analysis, CGM emerged as a cost-effective intervention with PWD gaining 0.54 quality-adjusted life
511 years (QALYs) and an ICER of US\$98,108 (AU\$141,242) per QALY compared to SMBG, assuming
512 a willingness-to-pay of US\$100,000 per QALY (S43). Similarly, the JDRF study comparing usual care
513 to CGM in PWD with HbA1c \geq 53mmol/mol (7.0%), reported a reduction in projected lifetime risk of
514 microvascular complications, with a gain of 0.60 QALYs and an ICER of US\$98,679 (AU\$142,064)
515 per QALY (S44). The cost-effectiveness of CGM was even greater in the HbA1c <53mmol/mol (7.0%)
516 cohort, with an average gain in 1.11 QALYs and an ICER of US\$78,943 (AU\$113,651) per QALY
517 (S44). Another study comparing CGM versus SMBG in people on intensive insulin therapy reported an
518 improvement in cost-effectiveness of 0.52 QALYs and an ICER of US\$45,033 (AU\$64,832) per QALY
519 (S45). However, a Spanish study by Garcia-Lorenzo et al comparing CGM to SMBG in a model
520 population showed that CGM was not cost-effective, with a gain of 0.05 QALYs and an ICER of
521 EUR€2,554,723 (AU\$4,166,268) per QALY, assuming a willingness-to-pay of EUR€25,000. However,
522 this study did not include costs such as emergency visits and diabetes-related hospital admissions in
523 their economic evaluation (S31). Overall, CGM seems to be a cost-effective intervention in the
524 management of T1DM, especially with long term regular use.

525

526 **Type 2 Diabetes**

527 An American study of people with T2DM not on prandial insulin, using CGM as a tool to inform
528 behavioural choices without clinician guidance, reported a gain of 0.07 QALYs and an ICER of
529 US\$8,898 (AU\$12,810) per QALY with CGM compared to SMBG, deeming it a cost-effective
530 intervention (S46). However, Garcia-Lorenzo et al reported that CGM was not cost-effective, with a
531 gain of 0.27 QALYs and an ICER of EUR€180,533 (AU\$294,415) per QALY (S31). These findings
532 suggest that CGM may be cost-effective in people with T2DM on insulin treatment.

533

534

535 **CONCLUSION**

536 As CGM continues to evolve, a new era of diabetes management has come. CGM offers added benefits
537 compared to traditional HbA1c measurement and SMBG, including detailed information on glycaemic
538 trends and excursions. CGM can improve glycaemic outcomes including HbA1c, TIR, and reduced
539 hypoglycaemia, with study findings consistently proving either non-inferiority or superiority compared
540 to SMBG, except for one study. Many studies have also shown high user satisfaction with CGM. CGM
541 can significantly improve long-term diabetes management and QOL in PWD. Further studies around
542 the impact of CGM on driving and telemedicine are required.

543

544 There is strong evidence for CGM use in T1DM management. Studies suggest that CGM is most
545 beneficial in adults with poorly managed T1DM, with a high HbA1c and/or hypoglycaemia

546 unawareness. Studies have also demonstrated that CGM is a cost-effective intervention in the
547 management of T1DM. Long term, regular daily use of CGM is required to maximise its benefits and
548 value. Further study is required to assess the efficacy and cost-effectiveness of CGM in people with
549 well managed T1DM.

550
551 The evidence for CGM use in T2DM management is less robust. There is more evidence for FGM
552 rather than traditional CGM use in T2DM, especially in people <65 years on insulin therapy with a high
553 HbA1c, and those who seldom perform SMBG. The main benefit of FGM in this population is HbA1c
554 reduction without increased hypoglycaemia. Further study around the cost-effectiveness of CGM in
555 T2DM management is required.

556

557

558

559 **REFERENCES**

- 560 1. Diabetes C, Complications Trial Research G, Nathan DM, Genuth S, Lachin J, Cleary P, et al.
561 The effect of intensive treatment of diabetes on the development and progression of long-term
562 complications in insulin-dependent diabetes mellitus. *N Engl J Med.* 1993;329(14):977-86.
- 563 2. The Royal Australian College of General Practitioners. General practice management of type
564 2 diabetes: 2016-18. East Melbourne, Australia 2016.
- 565 3. Hirsch IB. Professional flash continuous glucose monitoring as a supplement to A1C in
566 primary care. *Postgrad Med.* 2017;129(8):781-90.
- 567 4. Polonsky WH, Fisher L, Schikman CH, Hinnen DA, Parkin CG, Jelsovsky Z, et al. Structured
568 self-monitoring of blood glucose significantly reduces A1C levels in poorly controlled, noninsulin-
569 treated type 2 diabetes: results from the Structured Testing Program study. *Diabetes Care.*
570 2011;34(2):262-7.
- 571 5. American Diabetes Association. 7. Diabetes technology: standards of medical care in
572 diabetes-2020. *Diabetes Care.* 2020;43(Suppl 1):S77-S88.
- 573 6. Riddell MC, Gallen IW, Smart CE, Taplin CE, Adolfsson P, Lumb AN, et al. Exercise
574 management in type 1 diabetes: a consensus statement. *Lancet Diabetes Endocrinol.* 2017;5(5):377-
575 90.
- 576 7. Patton SR. Adherence to glycemic monitoring in diabetes. *J Diabetes Sci Technol.*
577 2015;9(3):668-75.
- 578 8. Mian Z, Hermyer KL, Jenkins A. Continuous glucose monitoring: review of an innovation in
579 diabetes management. *Am J Med Sci.* 2019;358(5):332-9.
- 580 9. Jones TW, Chee M, Haurat J, Holmes –Walker DJ, Johnson S, Makin J, et al., editors. Impact
581 on glycaemic outcomes of funding continuous glucose monitoring for youth in Australia. Australasian
582 Diabetes Congress; 2019 Sydney, Australia.

- 583 10. Sherr JL, Tauschmann M, Battelino T, de Bock M, Forlenza G, Roman R, et al. ISPAD
584 clinical practice consensus guidelines 2018: diabetes technologies. *Pediatr Diabetes*. 2018;19 Suppl
585 27:302-25.
- 586 11. Battelino T, Danne T, Bergenstal RM, Amiel SA, Beck R, Biester T, et al. Clinical targets for
587 continuous glucose monitoring data interpretation: recommendations from the international consensus
588 on time in range. *Diabetes Care*. 2019;42(8):1593-603.
- 589 12. Department of Health. 58,000 type 1 diabetics to have free access to new glucose monitoring
590 device Australia2020 [Available from: [https://www.health.gov.au/ministers/the-hon-greg-hunt-
591 mp/media/58000-type-1-diabetics-to-have-free-access-to-new-glucose-monitoring-device](https://www.health.gov.au/ministers/the-hon-greg-hunt-mp/media/58000-type-1-diabetics-to-have-free-access-to-new-glucose-monitoring-device).
- 592 13. Bruttomesso D, Laviola L, Avogaro A, Bonora E, Del Prato S, Frontoni S, et al. The use of
593 real time continuous glucose monitoring or flash glucose monitoring in the management of diabetes:
594 A consensus view of Italian diabetes experts using the Delphi method. *Nutr Metab Cardiovasc Dis*.
595 2019;29(5):421-31.
- 596 14. Ang E, Lee ZX, Moore S, Nana M. Flash glucose monitoring (FGM): A clinical review on
597 glycaemic outcomes and impact on quality of life. *J Diabetes Complications*. 2020;34(6):107559.
- 598 15. Wood A, O'Neal D, Furler J, Ekinci EI. Continuous glucose monitoring: a review of the
599 evidence, opportunities for future use and ongoing challenges. *Intern Med J*. 2018;48(5):499-508.
- 600 16. Burckhardt MA, Roberts A, Smith GJ, Abraham MB, Davis EA, Jones TW. The use of
601 continuous glucose monitoring with remote monitoring improves psychosocial measures in parents of
602 children with type 1 diabetes: a randomized crossover trial. *Diabetes Care*. 2018;41(12):2641-3.
- 603 17. Dexcom. Dexcom G6 continuous glucose monitoring system (Decom G6) reading and meter
604 value United States of America 2020 [Available from: [https://www.dexcom.com/faqs/is-my-dexcom-
605 sensor-accurate](https://www.dexcom.com/faqs/is-my-dexcom-sensor-accurate).
- 606 18. Aleppo G, Ruedy KJ, Riddlesworth TD, Kruger DF, Peters AL, Hirsch I, et al. REPLACE-
607 BG: A randomized trial comparing continuous glucose monitoring with and without routine blood
608 glucose monitoring in adults with well-controlled type 1 diabetes. *Diabetes Care*. 2017;40(4):538-45.
- 609 19. FDA news release: FDA expands indication for continuous glucose monitoring system, first
610 to replace fingerstick testing for diabetes treatment decisions [press release]. United States of
611 America2016.
- 612 20. FDA news release: FDA approves first continuous glucose monitoring system with a fully
613 implantable glucose sensor and compatible mobile app for adults with diabetes [press release]. United
614 States of America2018.
- 615 21. Pyrlis F, Brown F, Ekinci EI. Recent advances in management of type 1 diabetes. *Aust J Gen*
616 *Pract*. 2019;48(5):256-61.
- 617 22. Medtronic. MiniMed® 670G system real-world data show improved time in range and
618 reduced lows and highs across all patient groups including a 41 per cent time in range improvement
619 for previous MDI patients. Dublin: Medtronic; 2018.

- 620 23. Garg SK, Akturk HK. Flash glucose monitoring: the future is here. *Diabetes Technol Ther.*
621 2017;19(S2):S1-S3.
- 622 24. U.S. Food and Drug Administration. Freestyle Libre 14 day flash glucose monitoring system -
623 P160030/S017. United States of America2018.
- 624 25. FDA news release: FDA approves first continuous glucose monitoring system for adults not
625 requiring blood sample calibration [press release]. United States of America2017.
- 626 26. Abbott's FreeStyle Libre 2, with optional real-time alarms, secures CE Mark for use in Europe
627 [press release]. Illinois: Abbott2018.
- 628 27. Johnson ML, Martens TW, Criego AB, Carlson AL, Simonson GD, Bergenstal RM. Utilizing
629 the ambulatory glucose profile to standardize and implement continuous glucose monitoring in
630 clinical practice. *Diabetes Technol Ther.* 2019;21(S2):S217-S25.
- 631 28. Twigg S, Cohen N, Wischer N, Andrikopoulos S. Consensus position statement on: utilising
632 the ambulatory glucose profile (AGP) combined with the glucose pattern summary to support clinical
633 decision making in diabetes care. Australian Diabetes Society; 2019.
- 634 29. Mazze R. Advances in glucose monitoring: Improving diabetes management through
635 evidence-based medicine. *Prim Care Diabetes.* 2020;S1751-9918(20):30164-9.
- 636 30. Lu J, Ma X, Zhou J, Zhang L, Mo Y, Ying L, et al. Association of time in range, as assessed
637 by continuous glucose monitoring, with diabetic retinopathy in type 2 diabetes. *Diabetes Care.*
638 2018;41(11):2370-6.
- 639 31. Beck RW, Bergenstal RM, Riddlesworth TD, Kollman C, Li Z, Brown AS, et al. Validation
640 of time in range as an outcome measure for diabetes clinical trials. *Diabetes Care.* 2019;42(3):400-5.
- 641 32. Vigersky RA, McMahon C. The relationship of hemoglobin A1C to time-in-range in patients
642 with diabetes. *Diabetes Technol Ther.* 2019;21(2):81-5.
- 643 33. Monnier L, Colette C, Wojtusciszyn A, Dejager S, Renard E, Molinari N, et al. Toward
644 defining the threshold between low and high glucose variability in diabetes. *Diabetes Care.*
645 2017;40(7):832-8.
- 646 34. Akturk HK, Garg S. Technological advances shaping diabetes care. *Curr Opin Endocrinol*
647 *Diabetes Obes.* 2019;26(2):84-9.
- 648 35. Heinemann L, Freckmann G, Ehrmann D, Faber-Heinemann G, Guerra S, Waldenmaier D, et
649 al. Real-time continuous glucose monitoring in adults with type 1 diabetes and impaired
650 hypoglycaemia awareness or severe hypoglycaemia treated with multiple daily insulin injections
651 (HypoDE): a multicentre, randomised controlled trial. *Lancet.* 2018;391(10128):1367-77.
- 652 36. van Beers CA, DeVries JH, Kleijer SJ, Smits MM, Geelhoed-Duijvestijn PH, Kramer MH, et
653 al. Continuous glucose monitoring for patients with type 1 diabetes and impaired awareness of
654 hypoglycaemia (IN CONTROL): a randomised, open-label, crossover trial. *Lancet Diabetes*
655 *Endocrinol.* 2016;4(11):893-902.

- 656 37. Lind M, Polonsky W, Hirsch IB, Heise T, Bolinder J, Dahlqvist S, et al. Continuous glucose
657 monitoring vs conventional therapy for glycemic control in adults with type 1 diabetes treated with
658 multiple daily insulin injections: the GOLD randomized clinical trial. *JAMA*. 2017;317(4):379-87.
- 659 38. Beck RW, Riddlesworth T, Ruedy K, Ahmann A, Bergenstal R, Haller S, et al. Effect of
660 continuous glucose monitoring on glycemic control in adults with type 1 diabetes using insulin
661 injections: the DIAMOND randomized clinical trial. *JAMA*. 2017;317(4):371-8.
- 662 39. Battelino T, Conget I, Olsen B, Schutz-Fuhrmann I, Hommel E, Hoogma R, et al. The use and
663 efficacy of continuous glucose monitoring in type 1 diabetes treated with insulin pump therapy: a
664 randomised controlled trial. *Diabetologia*. 2012;55(12):3155-62.
- 665 40. Juvenile Diabetes Research Foundation Continuous Glucose Monitoring Study G,
666 Tamborlane WV, Beck RW, Bode BW, Buckingham B, Chase HP, et al. Continuous glucose
667 monitoring and intensive treatment of type 1 diabetes. *N Engl J Med*. 2008;359(14):1464-76.
- 668 41. Laffel LM, Kanapka LG, Beck RW, Bergamo K, Clements MA, Criego A, et al. Effect of
669 continuous glucose monitoring on glycemic control in adolescents and young adults with type 1
670 diabetes: a randomized clinical trial. *JAMA*. 2020;323(23):2388-96.
- 671 42. Pratley RE, Kanapka LG, Rickels MR, Ahmann A, Aleppo G, Beck R, et al. Effect of
672 continuous glucose monitoring on hypoglycemia in older adults with type 1 diabetes: a randomized
673 clinical trial. *JAMA*. 2020;323(23):2397-406.
- 674 43. Beck RW, Riddlesworth TD, Ruedy K, Ahmann A, Haller S, Kruger D, et al. Continuous
675 glucose monitoring versus usual care in patients with type 2 diabetes receiving multiple daily insulin
676 injections: a randomized trial. *Ann Intern Med*. 2017;167(6):365-74.
- 677 44. Ehrhardt NM, Chellappa M, Walker MS, Fonda SJ, Vigersky RA. The effect of real-time
678 continuous glucose monitoring on glycemic control in patients with type 2 diabetes mellitus. *J*
679 *Diabetes Sci Technol*. 2011;5(3):668-75.
- 680 45. Cosson E, Hamo-Tchatchouang E, Dufaitre-Patouraux L, Attali JR, Paries J, Schaepelynck-
681 Belicar P. Multicentre, randomised, controlled study of the impact of continuous sub-cutaneous
682 glucose monitoring (GlucoDay) on glycaemic control in type 1 and type 2 diabetes patients. *Diabetes*
683 *Metab*. 2009;35(4):312-8.
- 684 46. Bolinder J, Antuna R, Geelhoed-Duijvestijn P, Kroger J, Weitgasser R. Novel glucose-
685 sensing technology and hypoglycaemia in type 1 diabetes: a multicentre, non-masked, randomised
686 controlled trial. *Lancet*. 2016;388(10057):2254-63.
- 687 47. Campbell FM, Murphy NP, Stewart C, Biester T, Kordonouri O. Outcomes of using flash
688 glucose monitoring technology by children and young people with type 1 diabetes in a single arm
689 study. *Pediatr Diabetes*. 2018;19(7):1294-301.
- 690 48. Tyndall V, Stimson RH, Zammitt NN, Ritchie SA, McKnight JA, Dover AR, et al. Marked
691 improvement in HbA1c following commencement of flash glucose monitoring in people with type 1
692 diabetes. *Diabetologia*. 2019;62(8):1349-56.

- 693 49. Haak T, Hanaire H, Ajjan R, Hermanns N, Riveline JP, Rayman G. Flash glucose-sensing
694 technology as a replacement for blood glucose monitoring for the management of insulin-treated type
695 2 diabetes: a multicenter, open-label randomized controlled trial. *Diabetes Ther.* 2017;8(1):55-73.
- 696 50. Haak T, Hanaire H, Ajjan R, Hermanns N, Riveline JP, Rayman G. Use of flash glucose-
697 sensing technology for 12 months as a replacement for blood glucose monitoring in insulin-treated
698 type 2 diabetes. *Diabetes Ther.* 2017;8(3):573-86.
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700 **TABLES**701 **Table 1: List of continuous glucose monitoring devices available**

	Accuracy (MARD %)	Sensor life (Days)	Calibrations required (times/day)	Attachment site	Alert capability	How CGM is used	Smart phone compatibility	Insulin pump compatibility	Automated Insulin Adjustment	Approved for insulin dosing
<u>CONTINUOUS GLUCOSE MONITORING</u>										
Real-Time										
Dexcom G4 Platinum™□	9	7	2	Abdomen	Yes	Standalone or insulin pump	No	Animas Vibe	No	No
Dexcom G5 Mobile™□	9	7	2	Abdomen	Yes	Standalone or insulin pump	Yes	Tandem X2™□	T-Slim No	Yes
Dexcom G6™□	9.3	10	None	Arm# or Abdomen	Yes	Standalone or insulin pump	Yes	Tandem X2™□, Diabeloop™□ DBLG1	T-Slim Yes	Yes
Medtronic Enlite™□ sensor and Guardian™□ Link 2 transmitter	11	6	2	Abdomen	Yes	Insulin pump	No	MiniMed™□ 640G	Yes	No
Medtronic Enlite™□ Sensor and MiniLink™□ Transmitter	14	6	2	Abdomen	Yes	Insulin pump	No	MiniMed™□ Veo	No	No
Medtronic Guardian™□ Sensor 3 and Guardian™□ Connect transmitter	9.4	7	2	Arm or abdomen	Yes	Standalone	Yes	No	-	No

Medtronic Guardian™□ Sensor 3 and Guardian™□ Link 3 transmitter	9.6	7	3-4	Arm or Abdomen^	No	Insulin pump	No	MiniMed™□ 630G, MiniMed™□ 640G, MiniMed™□ 670G, MiniMed™□ 770G, MiniMed™□ 780G**	Yes	No
Senseonics Eversense™□	8.5	90-180*	2	Arm	Yes	Standalone	Yes	No	-	Yes
Retrospective										
Medtronic Enlite™□ Sensor and iPro2™□ Recorder	11	6	2	Abdomen	N/A	N/A	No	N/A	N/A	N/A
<u>FLASH GLUCOSE MONITORING</u>										
Real-Time										
Abbott™□ FreeStyle Libre	9.3	14	None	Arm	No	Standalone or insulin pump	Yes	Omnipod Horizon™□	Yes	Yes
Abbott™□ FreeStyle Libre 2	9.3	14	None	Arm	Yes	Standalone or insulin pump	Yes^^	Omnipod Horizon™□	Yes	Yes
Retrospective										
Abbott™□ FreeStyle Libre Pro	12.3	14	None	Arm	N/A	N/A	No	N/A	N/A	N/A

702

703

*Implantable sensor approved for 3 months use in America, and 6 months use in Europe

704

#Upper arm approved for use in Europe only, not approved by the FDA

705

^ Abdomen only if used with MiniMed™□ 670G

706

**Smartguard feature which automatically gives insulin boluses and adjusts basal insulin rate, CE-approved for use in Europe only

707

^^Designed for use with the FreeStyle Libre 2 app, which is currently under FDA review

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709 MARD – mean absolute relative difference; FDA = Food and Drug Administration

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Table 2: Summary of continuous glucose monitoring studies in people with type 1 or type 2 diabetes

Year	Inclusion Criteria	Duration	Change in HbA1c	Hypoglycaemia (<3.9 mmol/L)	Time in Range (3.9-10 mmol/L)
TYPE 1 DIABETES					
Heinemann et al (35) (HypoDE Study)	2018 Adults Hypoglycaemia unawareness	6 months	No significant difference	Reduction in mean number of hypoglycaemia events per 28 days by 0.28 events (95% CI -0.39 to -0.20; p<0.0001) in CGM group compared to control. Median time in hypoglycaemia in the CGM group was reduced from 70.9 mins/day (95% CI 38.8 to 130.2) to 23.9 mins/day (95% CI 12.9 to 54.5), compared to a reduction from 99.5 mins/day (95% CI 52.3 to 178.1) to 92.2 mins/day (95% CI 51.8 to 172.6) in the control group, p<0.0001.	No significant difference
Van Beers et al (36) (IN CONTROL study)	2016 Adults Hypoglycaemia unawareness CSII or MDI	44 weeks	No significant difference	Reduction in mean time in hypoglycaemia by 1.1 hours/day (95% CI -1.4 to -0.8; p<0.0001) in CGM group compared to control.	Increase in mean time in range by 2.3 hours/day (95% CI 1.9 to 2.7; p<0.0001) in CGM group compared to control.

Lind et al (37) (GOLD Study)	2017	Adults MDI	26 weeks	Reduction in mean HbA1c by 5mmol/mol (0.4%) (95% CI -6 to -3mmol/mol [-0.6 to -0.3%]; p<0.001) in CGM group compared to control.	Mean percentage of time in hypoglycaemia was 2.97% (95% CI -2.85% to 8.79%) in the CGM group compared to 4.79% (95% CI -3.11% to 12.69%) in the control group.	-
Beck et al (38) (DIAMOND Study)	2016	Adults MDI	24 weeks	Reduction in mean HbA1c by 7mmol/mol (0.6%) (95% CI -9 to -3mmol/mol [-0.8 to -0.3%]; p<0.001) in CGM group compared to control.	Median time in hypoglycaemia was 43 min/day (IQR 27-69) in the CGM group compared to 80 min/day (IQR 36-111) in the control group (p=0.002).	Increase in mean time in range by 77 mins/day (99% CI 6-147; p=0.005) in CGM group compared to control.
Battelino et al (39) (SWITCH Study)	2012	Children and adults CSII	6 months	Reduction in mean HbA1c by 5mmol/mol (0.4%) (95% CI -6 to -3mmol/mol [-0.6 to -0.3%]; p<0.001) during sensor on (CGM) compared to sensor off (control).	Median time in hypoglycaemia was 19 min/day (IQR 7.9-38) during sensor on (CGM) compared to 31 min/day (IQR 10-57) during sensor off (control; p=0.009).	Mean time in range was 774 min/day (95% CI 737 to 812) during sensor on (CGM) compared to 669 min/day (95% CI 635 to 703) during sensor off (control; p<0.001)
Tamborlane et al (40) (JDRF CGM Study)	2008	Children and adults	26 weeks	Age ≥25 years: Reduction in mean HbA1c by 6mmol/mol (0.5%) (95% CI -8 to -4mmol/mol [-0.7 to -0.4%]; p>0.001) in CGM group compared to control. Age 15-24 years: No significant difference Age 8-14 years: No significant difference	No significant difference	Age ≥25 years: Mean time in range was 986 min/day in the CGM group compared to 840 min/day in the control group (p<0.001). Age 15-24 years: No significant difference Age 8-14 years: No significant difference
Laffel et al (41) (CITY Study)	2020	Young Adults (14-24 years) CSII or MDI	17 months	Reduction in mean HbA1c by 4mmol/mol (0.37%) (95% CI -7 to -1mmol/mol [-0.7 to -0.1%]; p=0.01) in CGM group compared to control	Reduction in median percentage time in hypoglycaemia by 0.7% (95% CI -1.5 to -0.1; p=0.02) in CGM group compared to control.	Increase in mean percentage time in range by 6.9% (95% CI 3.1 to 10.7; p<0.001) in CGM group compared to control.

Pratley et al (42) (WISDM Study)	2020	Adults (≥60 years) CSII or MDI HbA1c <86mmol/mol (10.0%)	26 weeks	Reduction in mean HbA1c by 3mmol/mol (0.3%) (95% CI -4 to -1mmol/mol [-0.4 to -0.1%]; p<0.001) in the CGM group compared to control.	Reduction in median percentage time in hypoglycaemia by 1.9% (95% CI -2.8 to -1.1; p<0.001) in the CGM group compared to control.	Increase in mean percentage time in range by 8.8% (95% CI 6.0 to 11.5; p<0.001) in the CGM group compared to control.
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TYPE 2 DIABETES

Beck et al (43)	2017	Adults MDI HbA1c 59-85mmol/mol (7.5-9.9%)	24 weeks	Reduction in mean HbA1c by 3mmol/mol (0.3%) (95% CI -5 to 0mmol/mol [-0.5 to 0.0%]; p=0.022) in the CGM group compared to control.	No significant difference	Median time in range increased from 802 min/day to 882 min/day in the CGM group compared to an increase from 794 min/day to 836 min/day in the control group.
Ehrhardt et al (44)	2011	Adults HbA1c 53-108mmol/mol (7.0-12.0%) Not on prandial insulin	12 weeks	Mean reduction in HbA1c was 11mmol/mol (1.0%) (95% CI -14.2 to -8mmol/mol [-1.3 to -0.7%]) in the CGM group compared to 5mmol/mol (0.5%) (95% CI -8 to -3mmol/mol [-0.7 to -0.3%]) in the control group (p=0.006).	-	-

TYPE 1 AND TYPE 2 DIABETES

Cosson et al (45)	2009	Adults HbA1c 64-91mmol/mol (8.0-10.5%) T1DM: CSII or MDI T2DM: MDI or oral hypoglycaemics	48 hours	Overall: Reduction in mean HbA1c by 7mmol/mol (0.6%) (95% CI -8 to -5mmol/mol [-0.8 to -0.5%]; p=0.023) in the CGM group, compared to no significant HbA1c reduction in the control group. T2DM: Reduction in mean HbA1c by 7mmol/mol (0.6%) (95% CI -8 to -5mmol/mol [-0.8 to -0.5%]; p=0.05) in the CGM group compared to no significant HbA1c reduction in the control group. T1DM: No significant difference	No significant difference	No significant difference
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722 HbA1c = glycated haemoglobin; T1DM = type 1 diabetes mellitus; CSII = continuous subcutaneous insulin infusion; MDI = multiple daily injections; CI = confidence interval; CGM = continuous glucose monitoring;
 723 IQR = interquartile range; T2DM = type 2 diabetes mellitus

724 **Table 3: Summary of Flash glucose monitoring studies in people with type 1 or type 2 diabetes**

Year	Inclusion Criteria	Duration	Change in HbA1c	Hypoglycaemia (<3.9 mmol/L)	Time in Range (3.9-10 mmol/L)
<u>TYPE 1 DIABETES</u>					
Bolinder et al (46) (IMPACT Study)	2016 Adults HbA1c ≤59mmol/mol (7.5%) CSII or MDI Exclusion: Hypoglycaemia unawareness	6 months	No significant difference	Reduction in mean time in hypoglycaemia by 1.24 hours/day (95% CI -1.71 to -0.77; p<0.0001) in the FGM group compared to control (38.0% reduction). Reduction in mean time in nocturnal hypoglycaemia (11pm to 6am) by 0.47 hours (95% CI -0.70 to -0.24; p<0.0001) in the FGM group compared to control (39.8% reduction).	Increase in mean time in range by 1.0 hour/day (95% CI 0.41 to 1.59; p=0.0006) in FGM group compared to control.
Campbell et al (47) (SELFY Single-Arm Study)	2018 Children and Teenagers (4-17 years) CSII or MDI	8 weeks	Mean HbA1c reduced by 4mmol/mol (0.4%) (95% CI -6 to -3mmol/mol [-0.5 to -0.3%]) from 63mmol/mol (7.9%) (95% CI 60 to 65mmol/mol [7.7 to 8.1%]) at baseline to 59mmol/mol (7.5%) (95% CI 56 to 61mmol/mol [7.3 to 7.7%]) at study end (p<0.0001).	No significant difference	Mean time in range increased by 0.9 hours/day (95% CI 0.27 to 1.53) from 10.1 hours/day (95% CI 9.42 to 10.8) at baseline to 11.1 hours/day (95% CI 10.4 to 11.8) at study end (p=0.005).
Tyndall et al (48) (Prospective observational Study)	2019 Adults CSII or MDI	-	Median reduction in HbA1c by 4mmol/mol (0.4%) (IQR -10 to 0mmol/mol [-0.9 to 0.0%]); p<0.001 between the last value prior to FGM use and the most recent value.	Frequency of symptomatic hypoglycaemia (<3.5mmol/L) increased, with those reporting more than 2-3 episodes per week increasing from 25.8% to 48.8% (p<0.001) following commencement of FGM. The proportion of people experiencing any asymptomatic hypoglycaemia (<3.5 mmol/L) increased from 20.4% to 29.5% (p<0.001) following commencement of FGM.	-
<u>TYPE 2 DIABETES</u>					

Haak et al (49) (REPLACE Study)	2017	Adults HbA1c 59- 108mmol/mol (7.5- 12.0%) CSII or MDI	6 months	Overall: No significant difference Age <65 years: Mean HbA1c reduced by 6mmol/mol (0.5%) (95% CI -8 to -4mmol/mol [-0.7 to -0.4%]) in the FGM group, compared to a reduction of 2mmol/mol (0.2%) (95% CI -5 to 0.4mmol/mol [-0.4 to 0.04%]) in the control group (p=0.03). Age ≥65 years: Mean HbA1c reduced by 5mmol/mol (0.5%) (95% CI -8 to -3mmol/mol [-0.7 to -0.2%]) in the control group, compared to a reduction of 1mmol/mol (0.1%) (95% CI -3 to 2mmol/mol [-0.3 to 0.2%]) in the FGM group (p=0.008).	Reduction in mean time in hypoglycaemia by 0.47 hours/day (95% CI -0.72 to -0.22; p=0.0006) in the FGM group compared to control (43.1% reduction). Reduction in mean time in nocturnal hypoglycaemia (11pm to 6am) by 0.29 hours (95% CI -0.45 to -0.13; p=0.0001) in the FGM group compared to control (54.3% reduction).	No significant difference
Haak et al (50)	2017	Adults HbA1c 59- 108mmol/mol (7.5- 12.0%) CSII or MDI	12 months	-	Mean time in hypoglycaemia reduced by 0.70 hours/day (95% CI -1.01 to -0.39; p=0.0002) in the FGM group from baseline to study end (49.4% reduction). Mean time in nocturnal hypoglycaemia (11pm to 6am) reduced by 0.31 hours (95% CI -0.45 to -0.17; p=0.0002) in the FGM group from baseline to study end (52.3% reduction).	No significant difference
Yaron et al (S1)	2019	Adults HbA1c 59- 86mmol/mol (7.5- 10.0%) MDI	10 weeks	Mean HbA1c reduced by 9mmol/mol (0.9%) (95% CI -10 to -8mmol/mol [-0.9 to -0.8%]) in the FGM group, compared to a reduction of 3mmol/mol (0.3%) (95% CI -4 to -3mmol/mol [-0.4 to -0.2%]) in the control group (p=0.005).	No significant difference	-
Wada et al (S2)	2020	Adults HbA1c 59- 69mmol/mol (7.5- 8.5%) Non-insulin treated	24 weeks	At 12 weeks: No significant difference At 24 weeks: Reduction in mean HbA1c by 3mmol/mol (0.3%) (95% CI -6 to -1mmol/mol [-0.5 to -0.1%]); p=0.022) in the FGM group compared to control.	No significant difference	Increase in mean time in range by 2.36 hours/day (95% CI 1.21 to 3.51; p<0.001) in the FGM group compared to control.

726 T1DM = type 1 diabetes mellitus ; HbA1c = glycated haemoglobin; CSII = continuous subcutaneous insulin infusion; MDI = multiple daily injections; CI = confidence interval; FGM = Flash glucose monitoring; IQR =
727 interquartile range; T2DM = type 2 diabetes mellitus

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Author Manuscript

	Sensor Wear Life (Days)	Cost per Sensor	Transmitter Wear Life	Cost per Transmitter	Cost per Receiver/Reader	Annual Cost	Annual Cost with Subscription
CONTINUOUS GLUCOSE MONITORING							
Real-Time							
Dexcom Platinum™	G4 7	\$92.50	12 months	\$580	\$810	\$6,195	-
Dexcom Mobile™	G5 7	\$92.50	3 months	\$540	\$650 (Optional)	\$6,970*	-
Dexcom G6™	10	\$110	3 months	\$400	\$650 (Optional)	\$5,560*	\$3,960
Medtronic Enlite™ sensor and Guardian™ Link transmitter	6	\$75	12 months	\$699	N/A	\$5,199	\$3,000
Medtronic Enlite™ Sensor and MiniLink™ Transmitter	6	\$75	12 months	\$699	N/A	\$5,199	\$3,000
Medtronic™ Guardian Sensor 3 and Guardian™ Connect transmitter	7	\$75	12 months	\$699	N/A	\$5,199	\$3,300
Medtronic Guardian™ Sensor 3 and Guardian™ Link 3 transmitter	7	\$75	12 months	\$699	N/A	\$5,199	\$3,000
Retrospective							
Medtronic Enlite™ Sensor and iPro2™ Recorder	6	\$75	12 months	\$995	N/A	\$5,495	N/A
FLASH GLUCOSE MONITORING							
Abbott™	14	\$92.50	N/A	N/A	\$95 (Optional)	\$2,405*	N/A

733 Table 4: Cost of continuous glucose monitoring devices available in Australia

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735 *Annual cost without receiver

736 ^ Approved for 3 months use in America, and 6 months use in Europe

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745 FIGURE LEGENDS

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747 **Figure 1:** The ambulatory glucose profile is an internationally standardised report of continuous
748 glucose monitoring information. The report is organised into five major components: ①

749 Sensor capture data completeness; ② Glucose level statistics including hypoglycaemic events and time

750 in range; ③ Glucose profile based on a “model day”; ④ Glucose management indicator; and ⑤ Daily

751 glucose profiles.

AGP Report

21 April 2020 - 4 May 2020 (14 Days)

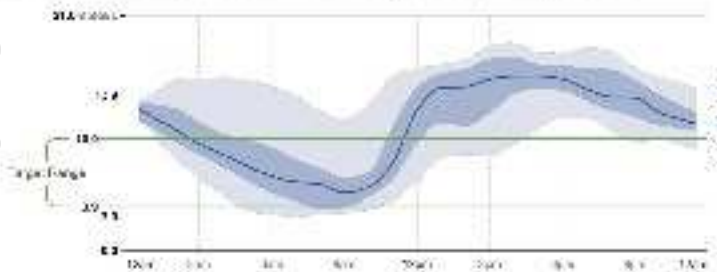
GLUCOSE STATISTICS AND TARGETS

① % Time CGM is Active	75%
②	
③	
④ Glucose Management Indicator (GMI)	8.1% or 85 mmol/mol
Glucose Variability	38.4%

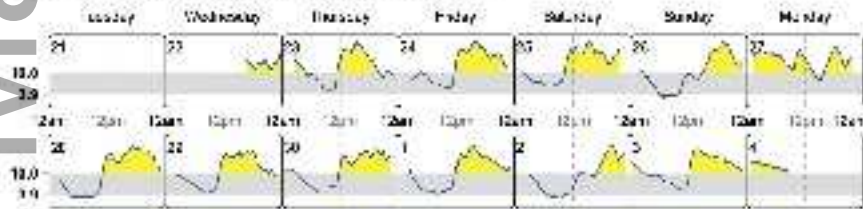
TIME IN RANGES



AMBULATORY GLUCOSE PROFILE (AGP)



DAILY GLUCOSE PROFILES



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