Beneficial effect of real-time continuous glucose monitoring system on glycemic control in type 1 diabetic patients: systematic review and meta-analysis of randomized trials

A Szypowska, A Ramotowska, K Dżygoła and D Golicki

Department of Pediatrics, Medical University of Warsaw, Dzialdowska 1, 01-184 Warsaw, Poland and 1Department of Pharmacoeconomics, Medical University of Warsaw, Warsaw, Poland

(Correspondence should be addressed to A Szypowska; Email: agnieszka.szypowska@gmail.com)

Abstract

Objective: Real-time continuous glucose monitoring (RT-CGM) provides detailed information on glucose patterns and trends, thus allowing the patients to manage their diabetes more effectively.

Design: The aim of this study was to explore the potential beneficial effects of the use of RT-CGM on diabetes management compared with self blood glucose measurement (SBGM) in patients with type 1 diabetes mellitus (T1DM), by means of a systematic review and meta-analysis of randomized controlled trials (RCTs).

Methods: MEDLINE, EMBASE, and the Cochrane Library were searched through by two independent investigators for RCTs concerning the use of RT-CGM in patients with T1DM. Only studies with a similar insulin regimen in the experimental and control groups were included in the analysis.

Results: Seven RCTs (n = 948) met the inclusion criteria. Combined data from all studies showed better HbA1c reduction in subjects using RT-CGM compared with those using SBGM (mean difference (MD) -0.25; 95% confidence interval (95% CI): from -0.34 to -0.17; P < 0.001). Patients treated with insulin pump and RT-CGM had a lower HbA1c level compared with subjects managed with insulin pump and SBGM (four RCTs, n = 497; MD -0.26; 95% CI: from -0.43 to -0.10; P = 0.002). The benefits of applying RT-CGM were not associated with an increasing rate of major hypoglycemic episodes. The use of RT-CGM for over 60–70% of time was associated with a significant lowering of HbA1c.

Conclusions: RT-CGM is more beneficial than SBGM in reducing HbA1c in patients with type 1 diabetes. Further studies are needed to evaluate the efficacy of this system in the pediatric population, especially in very young children.

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Background

The Diabetes Control and Complications Trial confirmed that tight metabolic control is regarded as crucial to prevent microvascular and macrovascular complications in type 1 diabetic patients (1). Both HbA1c and glucose variability play important roles in the evaluation of the risk of long-term diabetic complications (2). Intensive insulin therapy prevents or at least delays long-term diabetic complications. Aggressive diabetes management with continuous subcutaneous insulin infusion (CSII) or multiple daily injections (MDI) using insulin analogs and frequent blood glucose monitoring are the recommended methods to achieve therapeutic targets in type 1 diabetic patients.

The main factor limiting insulin management of type 1 diabetes mellitus (T1DM) subjects in the achievement of a strict glycemic goal is hypoglycemia (3). Unfortunately, despite active education, it is quite difficult to avoid hypoglycemia. Even the most frequent self blood glucose measurement (SBGM) gives insufficient information. Usually, T1DM patients carry out between four and eight finger-prick measurements per day, or less, and rarely monitor their blood glucose level at night. This is the cause of overlooking blood glucose excursion, and especially postprandial hyperglycemia, asymptomatic hypoglycemia, and glucose fluctuation during the night.

Continuous glucose monitoring (CGM) provides detailed information on glucose patterns and trends, thus allowing patients to manage their diabetes more effectively. Several continuous monitoring systems are commercially available. Some of them use CGM in a retrospective way and others are real-time glucose monitors. There are different types of real-time glucose monitors: the DexCom Seven (DexCom, San Diego, CA, USA), the MiniMed Paradigm Real-time Insulin Pump and CGM System (Medtronic, Medtronic MiniMed, Northridge, CA, USA), and the FreeStyle Navigator.
Real-time CGM (RT-CGM) provides a new dimension to diabetes management. Several studies, many of them observational, have assessed the effect of RT-CGM on metabolic control in type 1 diabetic patients (7). A number of trials have demonstrated a reduction in HbA1c with RT-CGM. Other studies have not confirmed any benefits or have found that the benefit associated with CGM was strongly related to age.

In this study, we sought to explore the potential beneficial effects of the use of RT-CGM on diabetes management compared with SBGM in patients with type 1 diabetes, by conducting a systematic review and meta-analysis of RCTs.

Inclusion and exclusion criteria

The systematic review and meta-analysis were conducted according to standards of the Cochrane Collaboration (8). Studies included in the review had to be RCTs with parallel or crossover design in which RT-CGM and self-monitoring of blood glucose were compared with self-monitoring of blood glucose alone in the management of type 1 diabetes. We included studies that used commercially available real-time glucose monitors: the DexCom Seven (DexCom), the MiniMed Paradigm Real-time Insulin Pump and CGM System (Medtronic), or the FreeStyle Navigator (Abbott Diabetes Care) and Guardian RT (Medtronic MiniMed, Northridge, CA, USA). Each system consists of a glucose oxidase-based electrochemical sensor, which is placed subcutaneously and along with a receiver to which interstitial glucose measurements are sent wirelessly and stored. A significant benefit of CSII over MDI for HbA1c reduction had been previously confirmed by some authors. Therefore, only studies with the same insulin regimen or studies with a similar proportion of patients using CSII and MDI in both experimental and control groups were included in the analysis. The studies had to be of at least 3 months’ duration and had to have a follow-up rate of over 80%. We excluded unpublished studies, letters to the editor, abstracts, and proceedings of scientific meetings. We also excluded studies in which patients used both CSII and MDI, but in which authors gave no information about the structure of usage in the experimental and control groups or the groups were not balanced in terms of the usage structure. We also excluded trials involving patients with type 2 diabetes, pregnant women with T1DM, and pancreas/islet-cell transplant patients. Studies using the Gluco-Watch G2 Biographer (Cygnus, Redwood City, CA, US) were not included in this analysis due to a different method of glucose measurement. This device takes non-invasive glucose measurements using low electric current to pull glucose through the skin. It caused a lot of skin irritations that led to very low compliance rates. Moreover, because of its side effects, the Gluco-Watch G2 Biographer has been withdrawn from the market. Trials that used other RT-CGM devices that are not available on the market anymore, or evaluated the use of blinded, retrospective CGM were excluded. Studies performed in settings such as pre- and post-surgical or cardiac care unit were excluded as well.

Outcomes

The primary end point was the change in HbA1c between the RT-CGM and the SBGM groups. The secondary end points were: major and minor hypoglycemic episodes (as defined by the investigators), mean daily area under the CGM curve for glucose < 3.89 mmol/l, mean daily area over the CGM curve for glucose > 9.99 mmol/l, local adverse effects and quality of life (QoL).

Search strategy

The following electronic databases were systematically searched through for relevant studies: MEDLINE (PubMed), EMBASE (Ovid), and the Cochrane Central Register of Controlled Trials. The search was conducted from 1996 to March 2011. The search strategy included the use of a validated filter for identifying RCTs (9). Keywords included a constellation of different phrases centered around CGM system (‘CGMS’ or ‘CGM’ or ‘Continuous Glucose Monitoring’ or ‘continuous glucose monitoring’ or ‘RT-CGM’ or ‘continuous subcutaneous glucose monitoring’ or ‘DexCom’ or ‘real-time system’ or ‘FreeStyle Navigator’ or ‘guardian’ or ‘sensor-augmented insulin pump’) and type 1 diabetes (‘diabetes type 1’ or ‘diabetes t. 1’ or ‘diabetes mellitus’ or ‘juvenile onset’ or ‘type 1 diabetes’ or ‘IDDM’ or ‘autoimmune diabetes’ or ‘DM1’ or ‘DM type 1’ or ‘insulin-dependent’ or ‘T1DM’ or ‘brittle diabetes’ or ‘T1D’). Subsequently, reference lists based on original studies and review articles were identified.

Data extraction

Two independent reviewers (A Ramotowska and K Dźygalo) screened the abstracts from the clinical trials according to the search strategy. Full texts of all potentially relevant articles were examined to determine whether they met the inclusion criteria. Both reviewers (A Ramotowska and K Dźygalo) extracted data independently, using standard data extraction forms. Extracted data were compared to

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Study quality
The methodological quality of the included studies was assessed by independent reviewers, without blinding to authorship or journal. The application of the following strategies associated with good-quality studies was examined: i) allocation concealment; ii) blinding of participants, investigators, outcome assessors and data analysts (yes/no); iii) intention-to-treat (ITT) analysis (yes/no); and iv) comprehensive follow-up. The allocation concealment was considered adequate when the randomization method used did not allow the investigator or the participant to identify or influence the intervention group before the entry of eligible participants into the study. The quality of allocation concealment was regarded as unclear when randomization was used, but no information about the method of randomization was available. It was regarded as inadequate when inappropriate methods of randomization (e.g. alternate medical record numbers, unsealed envelopes and tossing the coin) were used. In ITT analysis, a ‘yes’ answer meant that the authors had specifically reported undertaking this type of analysis and/or that our own study confirmed this finding. Conversely, ‘no’ meant that the authors had not reported the use of ITT analysis and/or that we could not confirm its use in the study assessment. The completeness of patient follow-up was evaluated by ascertaining the percentage of participants excluded or lost to follow-up. Completeness of follow-up was considered to be adequate if ≥80% of participants were included in the final analysis.

Statistical analysis
We used data from the end of each trial included in the systematic review. Data were analyzed using Comprehensive Meta-analysis Software (version 2.2.057; Biostat, Englewood, NJ, USA) (10). The mean difference (MD) was selected to determine differences in continuous outcomes between the experimental and control groups. The binary measure for individual studies and pooled statistics was calculated as the risk ratio (RR) between the experimental and the control groups, with 95% confidence interval (95% CI). The difference between the study groups was considered significant when the P value was <0.05 or when the 95% CI for RR did not exceed 1.0 and that for MD did not exceed 0. Heterogeneity was determined by I². Substantial heterogeneity was represented by I² of 50% or more (11). A fixed-effect model was used as baseline and a random-effect model was used in case of substantial heterogeneity.

Results
Study description
Based on the search strategy, 744 abstracts from clinical trials regarding CGMS were identified. The diagram of data extraction is illustrated in Fig. 1. We identified 38 articles that underwent further analysis. Finally, we included seven RCTs (n=948) to both qualitative and quantitative analyses (12, 13, 14, 15, 16, 17, 18). Table 1 summarizes the characteristics of the included trials. In five studies, insulin pump therapy was used in both experimental and control groups (13, 14, 16, 17, 18); in the next two studies, the number of patients treated with CSII or MDI was comparable for the experimental and control groups (12, 15). All trials included in the review, except one (17), were multicentered. All trials contained a sufficient proportion (≥80%) of participants in the final analysis. One of them included only a pediatric population (16), one regarded only adults (17), and the rest assessed mixed populations. The follow-up period ranged from 3 to 12 months. In four studies, randomization sequences were described and were adequate (12, 15, 16, 18). Allocation concealment was well reported and suitable in two studies (16, 18). Investigators of two studies conducted ITT analyses (12, 15). Withdrawals and dropouts were described in two studies (13, 18). Table 2 summarizes the quality assessment of the included studies.

HbA1c
Meta-analysis of seven RCTs (948 subjects) showed a significant reduction in HbA1c (MD -0.25; 95% CI: from -0.34 to -0.17; P<0.001) for patients managed with RT-CGMS compared with patients monitored with SBGM (Fig. 2). Moreover, patients treated with insulin pump combined with RT-CGMS had a lower HbA1c level (four RCTs, n=497; MD -0.26; 95% CI: from -0.43 to -0.10; P=0.002) compared with subjects managed with conventional insulin pump combined with SBGM (Fig. 3). The reduction
in HbA1c in adults (three RCTs, n = 224, MD = −0.37; 95% CI: from −0.76 to 0.02; P = 0.06, I² = 77%) and in children (three RCTs, n = 308, MD = −0.19; 95% CI: from −0.42 to −0.03; P = 0.09) using RT-CGM compared with SBGM groups was close to statistical significance. An additional analysis in subgroups divided according to glycemic control showed lower HbA1c in patients managed with RT-CGM compared with those managed with SBGM in both subgroups: with good metabolic control (one RCT, n = 129, MD = −0.31; 95% CI: from −0.46 to −0.16; P < 0.001) and poor glycemic control (four RCTs, n = 603, MD = −0.21; 95% CI: from −0.32 to −0.09; P < 0.001) at baseline.

There was a significant inverse correlation between the HbA1c level and the frequency of sensor use (13, 14, 16, 18). In the JDRF study (12), RT-CGM effectively lowered HbA1c only in adults aged ≥25 years. In four studies, more subjects in the RT-CGM group achieved the level of HbA1c of ≤7% (53 mmol/mol) than in the SBGM group (12, 13, 15, 18).

Major hypoglycemic episodes

RT-CGM usage had no influence on the incidence of major hypoglycemic episodes (six RCTs, n = 864, RR = 0.69; 95% CI: 0.41–1.14; P = 0.15). The data are shown in Fig. 4. None of the included studies confirmed that RT-CGM decreased the rate of major hypoglycemia. In two studies, authors excluded patients with a history of major hypoglycemia (12, 18).

Minor hypoglycemic episodes

Minor hypoglycemia, defined as glucose level below 3.89 mmol/l (70 mg%), was presented in five studies in two ways: as a number of episodes and as time spent in hypoglycemia (12, 13, 14, 15, 18). In one of them, authors did not find any difference in hypoglycemic episodes between patients using RT-CGM and control groups (13). There was no significant reduction in time spent in hypoglycemia in RT-CGM subjects, compared with that in SBGM group (12, 14, 15, 18).

Mean daily time and daily area under the CGM curve for glucose level of < 3.89 mmol/l

The area under the curve (AUC) calculated from CGM for glucose < 3.89 mmol/l (70 mg%) was significantly reduced in RT-CGM groups compared with patients monitored with SBGM in two studies (13, 15). Other authors did not show any differences between RT-CGM and control groups (14).

Hyperglycemia > 9.99 mmol/l (180 mg%)

A significant difference in favor of the RT-CGM group was observed with respect to time spent in
hyperglycemia in two studies (12, 14), which was not confirmed by other authors (15, 18). In two studies, there was no difference between RT-CGM groups and controls in the number of hyperglycemic events (12, 13). A significantly lower AUC in the RT-CGM groups compared with controls was noted by some authors (14) and not by others (13). In addition, there was a significant reduction in episodes of glucose above 250 mg%, in the RT-CGM group compared with controls in one study (12), which was not noted by other authors (15).

Mean amplitude of glycemic excursions

In two studies (14, 16), glycemic variability was significantly lower in the sensor group. The difference between the groups was not observed by other authors (15).

Ketoacidosis and local adverse events

Ketoacidosis was infrequent and without any significant difference between experimental and control groups. Local adverse events were uncommon and included mainly skin problems at the sensor or insulin infusion site.

Compliance

The sensor use was consistently high but declined over time in some trials (12, 14, 15, 16). An increased frequency of sensor use was associated with a greater reduction in HbA1c (13, 14, 15, 16, 18). The compliance with the sensor wear was age related and lower in children and the lowest in adolescents (12, 15). Self-reported pre-study daily blood glucose measurements were associated with successful use of RT-CGM (15). An association between sensor use and baseline HbA1c was not noted (12). No significant effect of age, duration of diabetes or duration of insulin pump therapy on the frequency of sensor use was noted by other authors (18).

Quality of life

Two studies (16, 17) estimated QoL as their secondary end points. We did not include this in our meta-analysis because of the different forms of evaluation used. In the trial by Kordonouri et al. (16), children aged 8–18 years and their primary caregivers were asked at the start of the study and at 24 and 52 weeks to complete the DISABKIDS and KIDSCREEN-27 questionnaires for evaluation of the patient’s health-related QoL and their caregiver’s impression of the patient’s QoL. Own well-being was assessed with the WHO-5 questionnaire. For physical, psychological, social support, and school, the scores were significantly lower at baseline compared with European norm data, reached normal values after 6 months and remained normal after 1 year, with no differences between experimental and control groups.

In the study by Peyrot et al. (17) all participants completed the User Acceptance Questionnaire, Insulin Delivery System Rating Questionnaire, and Blood Glucose Monitoring System Rating Questionnaire, which was developed for this study. In this trial, the investigators found that several patient-reported outcomes were significantly more positive in the RT-CGM arm than the control arm, including satisfaction measures, particularly the burden of blood glucose monitoring and convenience, as well as measures of health-related QoL, including social burden and diabetes-related worries.

Table 2

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References

Hirsch 2008 (13)
JDRF 2008 (12)
Raccah 2009 (14)
O’Connell 2009 (18)
Kordonouri 2010 (16)
Peyrot 2009 (17)
JDRF 2009 (15)

Figure 2

Mean difference and 95% CI of change in HbA1c (%) of patients treated with CSII or MDI in whom the RT-CGM and SBGM were compared with SBGM alone in the management of T1DM. Fixed-effect model. Heterogeneity $I^2 = 0%$.
carried out for a period of 3 or 6 months. The short duration of the follow-up made it difficult to predict whether the decreased HbA1c level would be maintained for a longer period. In view of a marked heterogeneity in the definition and assessment of hypoglycemia, a pooled analysis of this end point was not performed. Some studies reported a positive association between the primary end point and the degree of compliance. However, QoL was only assessed in two studies (16, 17). These studies were conducted using different questionnaires. The lack of standard QoL questionnaires prevented execution of the analysis. We observed a substantial clinical heterogeneity of the analyzed studies performed in adults. To deal with the statistical heterogeneity, we used the random-effect model.

Discussion

This meta-analysis of seven RCTs showed that the RT-CGM provides a superior benefit over self-monitoring of blood glucose with regard to HbA1c reduction in type 1 diabetic patients. The improvement in HbA1c in patients using the RT-CGM was achieved without an increase in severe hypoglycemia.

The recently published systematic review of nine RCTs indicated that RT-CGM has a beneficial effect on glycemic control in adult patients with T1DM, without an increase in the incidence of hypoglycemia. Less convincing evidence was available for children and type 2 diabetes patients (19). The authors of this review could not perform a meta-analysis because of an extensive clinical heterogeneity of trials. They included in their analysis patients using different methods of insulin administration (MDI or CSII), with different types of diabetes (type 1 and/or type 2 diabetes), as well as subjects monitored with Gluco-Watch G2 Biographer. Our meta-analysis differs from the study by Hoeks et al. (19) due to different inclusion criteria. In our meta-analysis, we included only trials with a similar method of insulin administration in both control and experimental groups. Previous meta-analyses had already shown that CSII compared with MDI was a more effective form of metabolic control (20, 21). Therefore, assessment of the efficacy of RT-CGM is not possible if the insulin delivery method is different in experimental and control groups.

Limitations at study and outcome level

In all included trials, medical devices for real-time glucose measurement were used, therefore blinding was not possible. Some of the analyzed trials revealed methodological limitations, including the lack of ITT analysis, unclear or inadequate allocation concealments and no data describing randomization. In one study, the sample size was limited (17). Moreover, the trials were conducted for up to 12 months; most of them were

Clinical implications

The previous meta-analysis comparing blinded CGM with SBGM showed no superiority of CGM over SBGM in lowering HbA1c in type 1 diabetic patients (5). However, those devices were clinician oriented and allowed only for a retrospective evaluation of data. A new generation of CGM devices offers real-time interstitial glucose monitoring and allows for advanced decisions made by patients. The results of our meta-analysis support the notion that the use of RT-CGM is associated with a significant lowering of HbA1c as well as glycemic variability. Both components – chronic sustained hyperglycemia and acute glycemic fluctuations – lead to diabetes complications through two main mechanisms – excessive protein glycation and activation of oxidative stress (22). Tight glycemic control is therefore of great importance in diabetes management. According to ISPAD guidelines, a target range of HbA1c for all age groups with type 1 diabetes of <7.5% (58 mmol/mol) is recommended (23). However, lowering HbA1c to below or around 7% (53 mmol/mol) has been shown to reduce microvascular and neuropathic complications of diabetes. Therefore, in American Diabetes Association (ADA) recommendations, a reasonable HbA1c goal for many
Continuous glucose monitoring in T1DM patients

Implications for further research

The use of RT-CGM provides a better insight into glycemic profiles, which may have a beneficial effect on patients with frequent severe hypoglycemia. Therefore, further studies are needed in subjects selected specifically for that problem. There are no randomized studies evaluating whether RT-CGM is beneficial in the management of toddlers and preschool children with T1DM. Although frequent SBGM is an integral part of intensive diabetes management, there are difficulties in minimizing glucose fluctuations in this age group. Parents and caregivers of young children experience a high level of stress related to fear of hypoglycemia, which can interfere with normal developmental and psychosocial interaction among diabetic children. Therefore, further studies are important not only for assessing the effectiveness, safety, and tolerance of RT-CGM device, but also to evaluate the impact of RT-CGM on the QoL. A decrease in compliance during the course of a trial was reported by some authors. Therefore, further research evaluating the lack or decreasing compliance in the follow-up is needed.

Conclusions

Our meta-analysis confirmed that the use of RT-CGM compared with SBGM effectively lowered HbA1c in type 1 diabetes. The benefit of applying RT-CGM was not associated with an increasing rate of acute hypoglycemia. The reduction in HbA1c was noted not only in patients with poorly controlled type 1 diabetes, but also in well-controlled subjects. The superiority of RT-CGM over SBGM in lowering HbA1c was also confirmed in pump users. Further age-related studies are needed to evaluate the efficacy of this system in the pediatric population, especially in very young children.

Declaration of interest

A Spyrowska and A Ramotowska co-authored educational materials for patients with diabetes, whose edition was sponsored by Abbott. Medtronic MiniMed sponsored the lectures as well as participation in medical conferences. K Dżygalo and D Golicki have no conflict of interest.

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References


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