



Glucose metrics and device satisfaction in adults with type 1 diabetes using different treatment modalities: a multicenter, real-world observational study

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Abstract

Aims To evaluate glucose metrics, device satisfaction and diabetes impact in adults with type 1 diabetes using different treatment modalities in a real-life setting in Italy.

Methods This was a multicentre, nationwide, cross-sectional study. Candidates were consecutively evaluated for eligibility during their routine medical visit at the diabetes centre. Researchers collected comprehensive demographic, socioeconomic, anamnestic and clinical data, and administered the Diabetes Impact and Device Satisfaction scale.

Results From 2021 to 2022, a total of 428 subjects, 45% males, with a median age of 32 years (IQR 23–47) were recruited in 11 participating centres from all over Italy. No differences in age, physical activity, and diabetes impact were found for the different treatment modalities. HCL/AHCL and SAP groups reported higher device satisfaction vs. MDI + SMBG and MDI + CGM ($p < 0.001$). Subjects treated with HCL/AHCL exhibited significantly higher TIR and significantly lower time spent in hypoglycemia level 1, time spent in hyperglycemia, CV and GMI compared to MDI + CGM, and significantly higher TIR and significantly lower time spent in hypoglycemia level 2, time spent in hyperglycemia, and CV compared to SAP. Significant reduction in hypoglycemia level 2 was also found with PLGM compared to SAP. High education attainment was associated with optimal metabolic control.

Conclusion Real-life use of advanced technologies for type 1 diabetes is associated with improved glucose metrics and device satisfaction. Education level also contributes to success of treatment.

Keywords Type 1 diabetes · Adults · Treatment · Continuous subcutaneous insulin infusion · Hybrid closed loop · Time in range · Device satisfaction

Background

Type 1 diabetes is a lifelong disease requiring intensive insulin treatment and daily monitoring of blood glucose levels. Despite efforts to maintain the glucose levels as close as possible to the recommended target, the majority of patients do not achieve this goal, leading to an increased risk of acute and chronic complications and adverse effects on quality of life [1].

However, in the last decades there have been many technological advances that have positively impacted the

management of type 1 diabetes. First developed in the late 1970s, insulin pumps providing continuous subcutaneous insulin infusion (CSII) have shown to reduce both HbA1c and the rate of hypoglycemic events when compared with multiple daily injections (MDI) of insulin [2]. More recently, continuous glucose monitoring (CGM) with minimally invasive devices has further revolutionized diabetes care, with meaningful improvements in glycemic control, risk of hypoglycemia, and quality of life as compared with self-monitoring of blood glucose (SMBG) [3–7]. CGM devices provide actionable information that is updated every

few minutes, including historic and current glucose values, rate of change of glucose, and alarms/alerts for high or low glucose fluctuations. However, they differ from each other by configuration (all-in-one vs. multicomponent devices), type of sensor (transcutaneous vs. fully implantable sensors), visualization tools (handheld receiver and/or smartphone apps), sensor lifetime, data update cycle (real-time vs. intermittently scanned devices), type of glucose alerts (only threshold alerts vs. threshold, predictive, and rate-of-change alerts), possibility of integration with other devices, and other features.

Importantly, the combination of CSII and CGM technologies has resulted in increasing level of automation of insulin delivery in response to sensor glucose readings, ranging from no automation (sensor-augmented pump [SAP] therapy) to algorithm-driven suspension of basal insulin for actual and/or impending hypoglycemia (predictive low glucose management, PLGM), algorithm-driven infusion of basal insulin (hybrid closed loop [HCL] insulin delivery) or algorithm-driven infusion of both basal insulin and correction boluses (advanced hybrid closed loop [AHCL] insulin delivery). In randomized clinical trials, SAP therapy has been associated with significant HbA_{1c} lowering as compared with MDI+ SMBG [8], and PLGM and HCL/AHCL systems with reduced hypoglycemia measures and increased time spent within the target glucose range of 70–180 mg/dL (TIR) together with reduced time spent in hypoglycemia, respectively, as compared with SAP [9, 10]. Importantly, HCL/AHCL systems have obtained more favorable psychological outcomes than the comparators in the majority of published trials [11].

Ultimately, several alternative opportunities are nowadays available for the treatment of type 1 diabetes, possibly with different effects on glycemic control and patient-reported outcomes (PROMs). A multicentre, real-world observational study conducted at 22 pediatric diabetes centers in Italy has recently confirmed that patients treated with HCL/AHCL systems achieve the highest TIR and the lowest time spent in hyperglycemia as compared with other therapeutic modalities [12]. Moreover, SAP, PLGM and HCL/AHCL, but not MDI+ SMBG, were associated with increased device satisfaction and lower diabetes impact than MDI+CGM as measured by the Diabetes Impact and Device Satisfaction (DIDS) scale.

The current study was designed to evaluate glycemic control and PROMs in a cohort of adult patients with type 1 diabetes using different treatment modalities, including traditional strategies for glucose monitoring and insulin administration and more advanced technological approaches.

Materials, and methods

This was a multicentre, nationwide, cross-sectional study. Inclusion criteria were: 18 to 60 years of age, being diagnosed with type 1 diabetes for at least six months, being on a MDI- or CSII-based treatment for at least three months, adequate understanding of Italian language. Major exclusion criteria were personal history of psychiatric disease and use of open source automated insulin delivery systems. Candidates were consecutively evaluated for eligibility during their routine medical visit at the diabetes centre and enrolled after giving informed consent.

The study was submitted to local institutional ethics committees (protocol no. 2020 439, approved on March 25, 2021 by the Regional Ethics Committee of Marche, University Hospital “Ospedali Riuniti”, Ancona, Italy, as the coordinating center) and carried out in adherence to Good Clinical Practice, ICH Harmonized Tripartite Guidelines for Good Clinical Practice and Declaration of Helsinki.

Procedures

Upon obtaining informed consent, researchers collected comprehensive demographic, anamnestic and clinical data, and administered the 11-item DIDS scale. Importantly, collection of demographic data involved socio-economic indicators such as educational attainment (classified as follows: low: lower secondary school or less, medium: upper secondary school, or high: university degree or more), employment status, household annual income (classified as follows: low: <26,000 €, medium: 26,000–54,999 €, or high > 54,999 €) and housing tenure. Time spent doing physical activity was recorded from self-reporting and expressed as hours per week. Number of diabetic ketoacidosis (DKA) and severe hypoglycemia (i.e., requiring third-party assistance) episodes in the past 12 months was collected from both self-reporting and medical records. Finally, for CGM users, the following metrics were obtained from the last 30 days before enrolment: TIR, time spent in hypoglycemia level 1 (< 70–54 mg/dL), time spent in hypoglycemia level 2 (< 54 mg/dL), time spent in hyperglycemia level 1 (> 180–250 mg/dL), time spent in hyperglycemia level 2 (> 250 mg/dL), coefficient of variation of glucose (CV), and Glucose Management Indicator (GMI). Devices data sources included Dexcom Clarity (Dexcom, Inc., San Diego, CA, USA), Glooko-Diasend (Glooko, Inc., Mountain View, CA, USA), Medtronic Carelink System (Medtronic, Inc., Minneapolis, MN, USA), and LibreView (Abbott Diabetes Care, Inc., Alameda, CA, USA) platforms.

Diabetes impact and device satisfaction scale questionnaire

The DIDS questionnaire consists of 11 items, each rated using a 10-point Likert scale, assessing two domains [13]. The first domain comprises seven items measuring satisfaction related to insulin delivery devices, while the second domain includes the remaining four items assessing the impact of diabetes on daily activities, concerns over hypoglycemia, and sleep disturbances. Higher scores in the two domains indicate higher device satisfaction and higher diabetes impact, respectively.

Importantly, the DIDS scale has been recently translated into Italian and validated in a pediatric population [12]. For the purposes of this study, validation assessment was carried out in a subsample of adult participants using CGM devices. Briefly, structural integrity was assessed through confirmatory factor analysis, internal consistency reliability by calculating the Cronbach's alpha coefficient, and discriminant ability by comparing people with $TIR \geq 75\%$ and $TIR < 50\%$ through the Wilcoxon rank-sum test.

Treatment modalities

For the purposes of our study, the following treatment modalities were compared: MDI + SMBG, MDI + CGM, SAP, PLGM, and HCL/AHCL. All devices were provided by the Italian National Health System with no charge for the patients, regardless of their income, age, or gender.

Statistical analysis

Continuous variables were expressed as mean $\pm$ standard deviation or median (interquartile range), while discrete variables as absolute and percentage frequencies. Normal distribution of continuous variables was assessed through the Shapiro-Wilks test. Demographic, anamnestic and clinical data, treatment satisfaction and impact of diabetes as measured through the DIDS, and CGM-derived glucose metrics were evaluated according to treatment modalities, and the Kruskal-Wallis test was used to compare groups.

The probability of achieving optimal glycemic control ($TIR \geq 70\%$) with the different treatment modalities was assessed via a logistic regression model; specifically, achieving a $TIR \geq 70\%$ (yes vs. no) was the dependent variable, the therapeutic modality was the explicative factor, and gender, age, disease duration, physical activity, educational attainment, and family income were entered as controlling covariates. Since TIR values were not available for subjects treated with MDI + SMBG, they were not included in this analysis.

Factors associated with device satisfaction and diabetes impact were analysed using quantile regression models. In this analysis, DIDS scores were treated as outcome variables of interest, while treatment modalities were examined as primary factors. To ensure a comprehensive understanding, adjustments were made for a number of variables, including gender, age, disease duration, physical activity, educational attainment, and family income.

Results

From 2021 to 2022, a total of 428 subjects, 45% males, with median age of 32 years (IQR 23–47) and median diabetes duration of 17 years (IQR 11–25), were recruited in 11 participating centres from all over Italy (Table S1). Main patients' characteristics are reported in Table 1. Information on treatment modality was available for 427 out of 428 participants. Specifically, 39 (9.1%) subjects were on MDI + SMBG, 155 (36.3%) on MDI + CGM, 99 (23.2%) on SAP therapy, 33 (7.7%) on PLGM, and 101 (23.7%) on HCL/AHCL. All SAP users were using tubeless pumps.

Figure 1 and Table S2 show the demographic and clinical characteristics of the subjects by treatment modalities. No statistically significant differences in age, physical activity, and diabetes impact were found among treatment modalities. Patients treated with SAP and HCL/AHCL had a significantly longer diabetes duration [19 (11–26) years and 18 (13–28) years, respectively] than those treated with MDI + CGM [14 (8–22) years], and reported higher device satisfaction vs. both MDI + SMBG and MDI + CGM. Patients treated with PLGM also exhibited a significantly higher device satisfaction than MDI + SMBG.

Subjects treated with HCL/AHCL exhibited significantly higher TIR [73% (64–80)] and significantly lower time spent in hypoglycemia level 1 [1% (1–2)], time spent in hyperglycemia [20% (16–26)], CV [32% (29.7–36)] and GMI [6.9% (6.7–7.2)] compared to patients treated with MDI + CGM, and significantly higher TIR and significantly lower time spent in hypoglycemia level 2 [0% (0–1)], time spent in hyperglycemia, and CV compared to SAP therapy (Fig. 2, Table S2). Significant reduction in hypoglycemia level 2 was also found with PLGM as compared with SAP therapy. Number of self-reported episodes of DKA and severe hypoglycemia was generally low (Table S3).

The use of HCL/AHCL systems increased the probability of being at TIR target ($\geq 70\%$) by 5.1 times compared to MDI + CGM ($p < 0.001$), independently from demographic, clinical and anamnestic variables. Furthermore, completing a “high” level of education increased the probability of being at $TIR \geq 70\%$ by 3.5 times compared to a low level ($p = 0.036$). The probability of having $TIR \geq 70\%$ also

Table 1 Main demographic and clinical patients' characteristics

Characteristics	<i>n</i>	
Age, years [median (IQR)]	428	32 (23; 47)
Diabetes duration, years [median (IQR)]	428	17 (11; 25)
Male gender, n (%)	428	193 (45.09)
Glucose Management Indicator, % [mean (SD)]	376	7.1 (0.7)
Therapeutic Strategy, n (%)	427	
MDI + SMBG		39 (9.1)
MDI + CGM		155 (36.3)
SAP		99 (23.2)
PLGM		33 (7.7)
HCL/AHCL		101 (23.7)
Time with glucose level below 54 mg/dL, % [median (IQR)]	386	0.2 (0; 1)
Time with glucose level between 54–69 mg/dL, % [median (IQR)]	386	2 (1; 4)
Time in glucose range 70–180 mg/dL, % [median (IQR)]	386	64 (51.3; 75)
Time with glucose level between 181–250 mg/dL, % [median (IQR)]	386	24 (18; 29.5)
Time with glucose level above 250 mg/dL, % [median (IQR)]	386	7 (3; 14)
Subjects with at least one episode of hypoglycemia in the previous year, n, %	426	33 (7.7)
Subjects with at least one episode of DKA in the previous year, n, %	427	6 (1.4)
Frequency of SMBG, tests/day [median (IQR)]	427	2 (0; 4)
Physical activity, hours / week [median (IQR)]	427	2 (0; 4)
Geographical area, n (%)	428	
North		126 (29.4)
Centre		63 (14.7)
Sud		239 (55.8)
Educational level, n (%)	412	
Low		39 (9.4)
Medium		280 (67.8)
High		93 (22.8)
Family gross annual income, n (%)	363	
Low		148 (40.8)
Medium		167 (46.0)
High		48 (13.2)

IQR: interquartile range; SD: standard deviation; MDI: multiple daily injections of insulin; SMBG: self-monitoring of blood glucose; CGM: continuous glucose monitoring; SAP: sensor-augmented pump; PLGM: predictive low glucose management; HCL: hybrid closed loop; AHCL: advanced hybrid closed loop

increased by 2% and 9% for each year of age and each hour of physical activity added, respectively (Table 2).

Table 3 reports the predictors of device satisfaction and diabetes impact as assessed with quantile regression analysis. Compared to MDI + CGM, SAP and HCL/AHCL treatments were significantly and independently associated with higher device satisfaction, MDI + SMBG with lower device satisfaction, and PLGM with higher diabetes impact. With regards to demographic and socio-economic characteristics, higher diabetes duration was associated with a higher device satisfaction, while increasing age and having a “medium” vs. “low” family income were both associated with lower diabetes impact.

The validation assessment of the Italian version of the DIDS scale was conducted in a subsample of 389 CGM

users, 45% male, with a median age of 32 years (23–47), and showed moderate level of structural integrity (Table S4, Figure S1) and good internal consistency (Tables S5).

In detail, internal consistency estimates were 0.71 (95% CI: 0.66; 0.75) and 0.74 (95% CI: 0.70; 0.78) for Device Satisfaction and Diabetes Impact domains, respectively. When evaluating the contribution of each single item to overall internal consistency, the Cronbach's α coefficient ranged from 0.63 to 0.74. The discriminant validity assessment was carried out comparing 75 participants with TIR < 50% with 100 participants with TIR $\geq$ 75%. Significant differences in both DIDS domains were observed, with higher median device satisfaction and lower diabetes impact being reported for participants with TIR $\geq$ 75% (Figure S2).

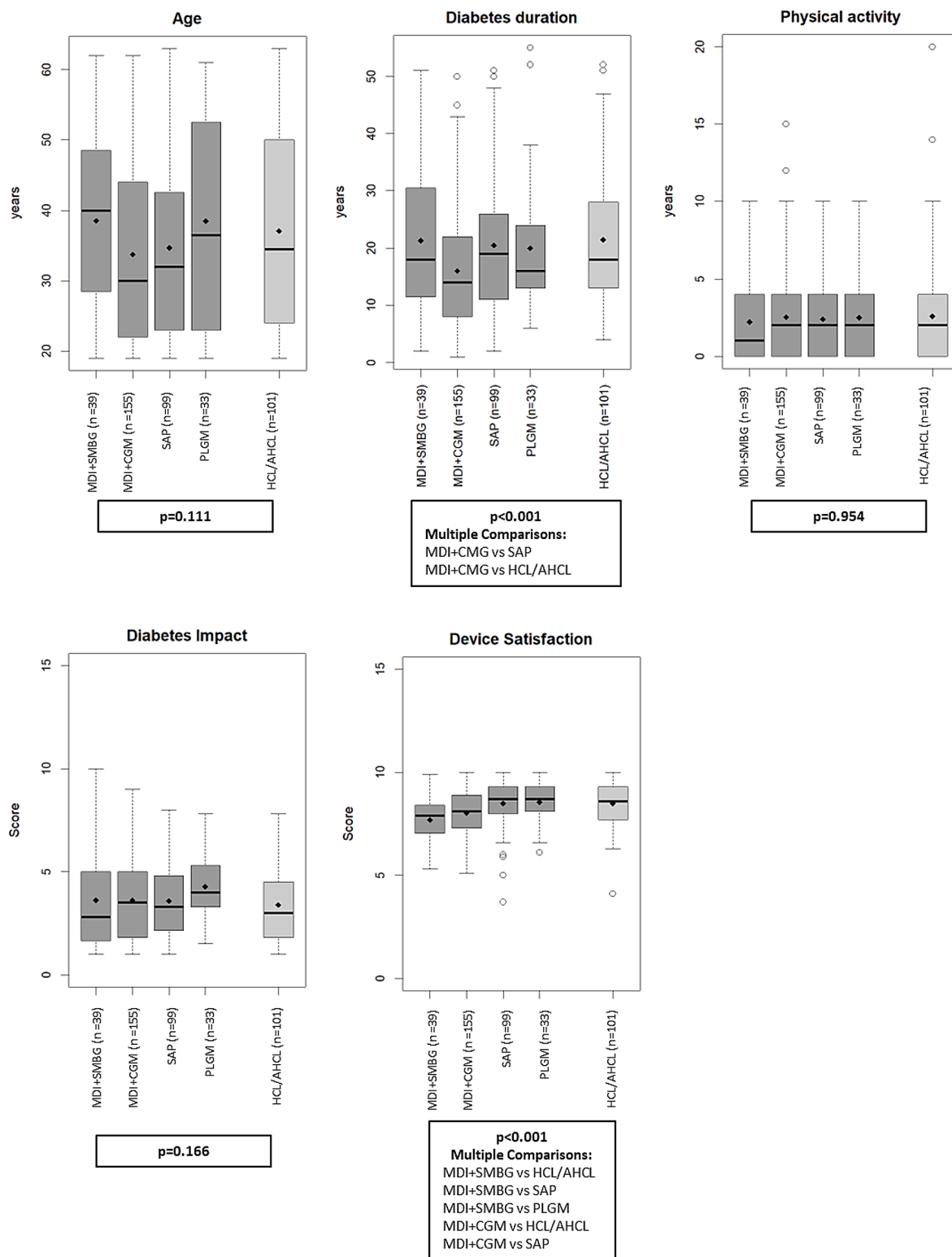


Fig. 1 Subjects' demographic and clinical characteristics by treatment modalities. MDI: multiple daily injections of insulin; SMBG: self-monitoring of blood glucose; CGM: continuous glucose monitoring;

PLGM: predictive low glucose management; HCL: hybrid closed loop; AHCL: advanced hybrid closed loop

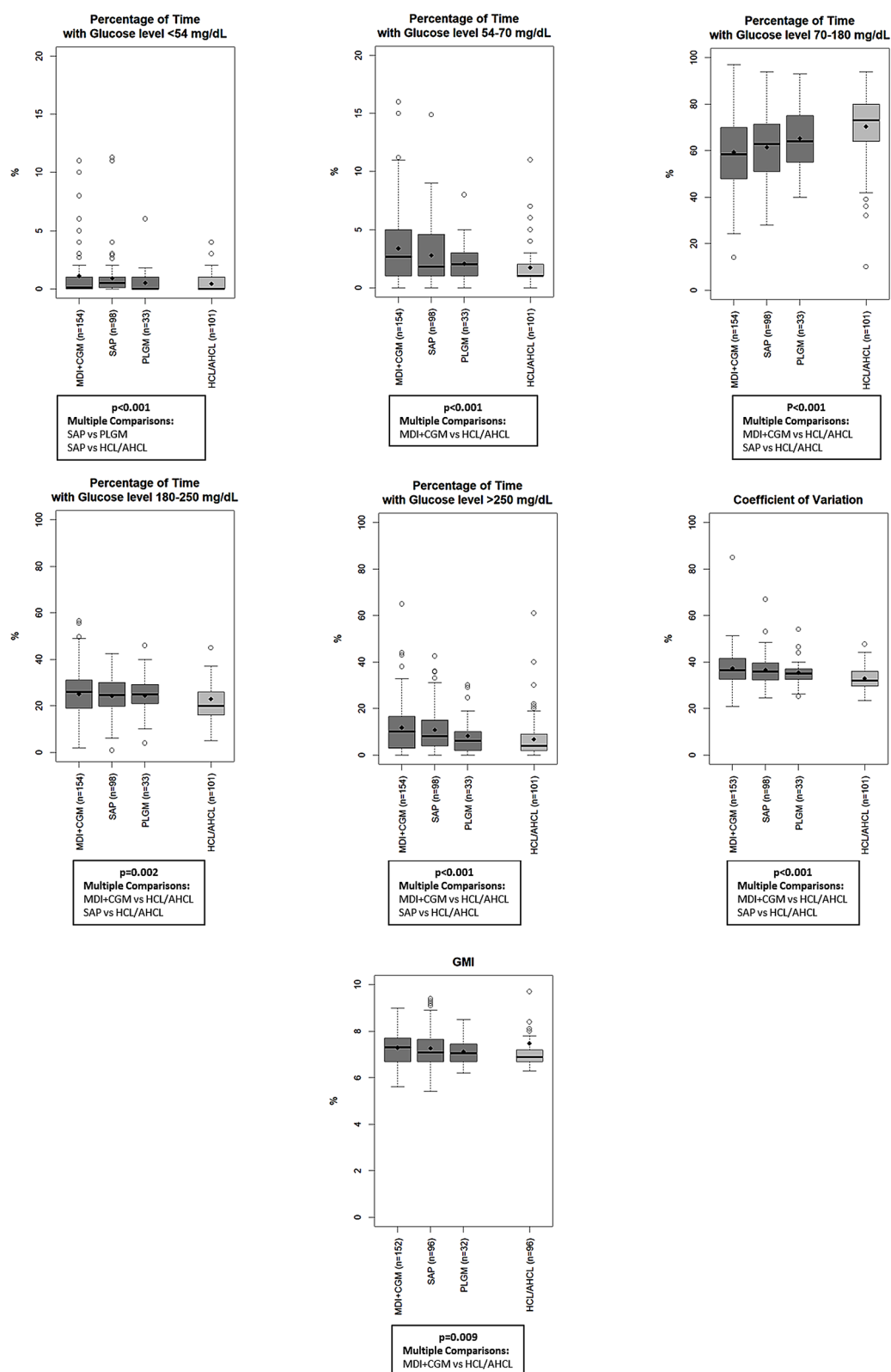


Fig. 2 Glucose metrics by treatment modalities. MDI: multiple daily injections of insulin; SMBG: self-monitoring of blood glucose; CGM: continuous glucose monitoring; PLGM: predictive low glucose management; HCL: hybrid closed loop; AHCL: advanced hybrid closed loop

Table 2 Variables associated to optimal metabolic control (TIR $\geq 70\%$, $n = 143$)

	OR	95%CI	<i>p</i>
Gender: female vs. male	0.76	0.45; 1.26	0.282
Age (years)	1.02	1.01; 1.06	0.004
Diabetes duration (years)	0.99	0.96; 1.02	0.410
Therapy: SAP vs. MDI + CGM	1.25	0.66; 2.38	0.498
Therapy: PLGM vs. MDI + CGM	1.27	0.48; 3.18	0.615
Therapy: HCL/AHCL vs. MDI + CGM	5.06	2.68; 9.79	<0.001
Physical activity (hours/week)	1.09	1.01; 1.19	0.035
Education level: medium vs. low	1.98	0.74; 6.02	0.196
Education level: high vs. low	3.46	1.17; 11.39	0.031
Family annual income: medium vs. low	0.78	0.45; 1.36	0.383
Family annual income: high vs. Low	0.91	0.4; 2.05	0.823

TIR: time in range; SAP: sensor-augmented pump; MDI: multiple daily injections of insulin; SMBG: self-monitoring of blood glucose; CGM: continuous glucose monitoring; PLGM: predictive low glucose management; HCL: hybrid closed loop; AHCL: advanced hybrid closed loop; OR: Odds Ratio; 95% CI: 95% Confidence Interval. Statistically significant results are in bold

Discussion

In the last few years, there have been many technological advances in glucose monitoring and insulin delivery, which have resulted in new opportunities for the treatment of type 1 diabetes.

The results of our study show that adult HCL/AHCL users with type 1 diabetes achieve the highest TIR, the lowest time spent in hyperglycemia, and the lowest time spent in hypoglycemia compared to other CGM-enhanced treatment modalities, with statistically significant differences being reported vs. MDI and SAP therapies. What's more, HCL/AHCL users exhibited median TIR values that met the recommended target of $> 70\%$ for non-fragile non-pregnant adults with type 1 diabetes [14] with negligible time spent in hypoglycemia, and use of HCL/AHCL systems was

identified as the single best predictor of achieving optimal metabolic control.

Randomized clinical trials and other observational studies have already shown the superiority of such systems in providing favourable glycemic outcomes as compared with other treatment modalities [10, 15], however our report is the first in the literature focusing on the Italian scenario and evaluating also the socioeconomic status of users, with more than 400 adult participants enrolled in 11 diabetes centres from all over the country.

In our analysis, PLGM was associated with lower time spent < 54 mg/dL than SAP therapy. Reduction of hypoglycemia measures with PLGM systems has also been replicated in randomized clinical trials and real-world studies [9, 16]. In line with these findings, international guidelines recommend use of integrated CGM and insulin pump systems proving automated insulin suspension/dosing over non-integrated systems in persons with type 1 diabetes [17, 18].

Time spent in hypoglycemia was numerically similar between HCL/AHCL and PLGM users. However, this is not surprising; in fact, according to the results of different RCT and real-world studies, superiority of HCL/AHCL systems vs. PLGM for hypoglycemia reduction has yet to be proven with certainty [19].

Interestingly, occurrence of severe hypoglycemic episodes was infrequent with any treatment modality, the proportion of patients experiencing at least one episode being lower than that reported in the Study of Adults' GlycEmia in T1DM (SAGE), therefore confirming a high level of commitment in the management of hypoglycemia among the Italian patients [1, 20].

Use of technological devices for glucose monitoring and/or insulin administration was generally associated with higher device satisfaction without increased disease burden compared to the traditional approach, except for PLGM. In

Table 3 Variables associated with DIDS domains. Results of quantile regression analysis

Variables	Device Satisfaction		Diabetes Impact	
	B	95%CI	b	95%CI
Gender: female vs. male	0.03	-0.12; 0.34	0.36	-0.12; 0.81
Age (years)	-0.01	-0.02; 0.01	-0.02	-0.04; -0.01
Diabetes duration (years)	0.02	0.01; 0.04	-0.01	-0.04; 0.01
Therapy: MDI + SMBG vs. MDI + CGM	-0.71	-1.2; -0.01	0.42	-0.5; 2.08
Therapy: SAP vs. MDI + CGM	0.30	0.02; 0.59	0.16	-0.45; 0.88
Therapy: PLGM vs. MDI + CGM	0.29	-0.1; 0.65	1.18	0.03; 1.71
Therapy: HCL/AHCL vs. MDI + CGM	0.31	-0.08; 0.62	0	-0.9; 0.58
Physical activity (hours/week)	-0.03	-0.06; 0.03	0.04	-0.01; 0.1
Education level: medium vs. low	-0.17	-0.49; 0.26	-0.23	-0.97; 0.34
Education level: high vs. low	-0.11	-0.51; 0.36	0.13	-0.95; 0.79
Family annual income: medium vs. low	0.05	-0.28; 0.29	-0.74	-1.11; -0.21
Family annual income: high vs. low	0.41	-0.04; 0.72	0.11	-0.38; 0.81

b: quantile regression coefficient; 95%CI: 95% Confidence Interval; MDI: multiple daily injections of insulin; SMBG: self-monitoring of blood glucose; CGM: continuous glucose monitoring; PLGM: predictive low glucose management; HCL: hybrid closed loop; AHCL: advanced hybrid closed loop. Statistically significant results are in bold

recent years, the assessment of PROMs, including quality of life and satisfaction with treatments and technologies, has progressively emerged as a critical factor for successful management of type 1 diabetes [11]. Indeed, patient satisfaction has been linked with persistent use of devices and improved glycemic control [21, 22]. For the purposes of our analysis, HCL and AHCL users were pooled together, however there is evidence in the literature that users' acceptance is increased with AHCL as compared with earlier systems, maybe due to frequent alarms and need for calibration by fingerstick glucose to maintain the Auto-Mode with the latter [23, 24].

With regards to both glycemic outcomes and patient satisfaction, the results of our adult cohort are in line with those of the recently published pediatric study [12], therefore confirming that diabetes devices are beneficial in the whole spectrum of patients with type 1 diabetes, and HCL/AHCL systems represent nowadays the gold standard of insulin replacement treatment [25]. Nevertheless, use of technological devices is still limited in Italy, with only 40.8% and 24% of patients with type 1 diabetes being on CGM or an insulin pump, respectively, the levels of uptake being even lower among adults and in southern regions [26]. Inadequate number of professionals in the diabetes team, need for high-level training of both healthcare professionals and patients, insufficient allocation of economic resources, and heterogeneous reimbursement policies are well-known barriers for a wider spread of diabetes devices [27, 28]. We hope that our research may convince both the healthcare professionals and the payers of the irreplaceable role of technology for the management of type 1 diabetes mellitus.

Interestingly, completing a "high" level of education was independently associated with reaching $\geq 70\%$ of TIR among CGM users. This is not surprising when one recalls that self-management of type 1 diabetes requires numerical skills and simultaneous consideration of multiple variables (e.g., deviation from target glucose, glucose trend, carbohydrate intake, insulin sensitivity factor, insulin on board, etc.) before making treatment decisions [29, 30]. However, glucose-driven automated insulin delivery in PLGM and HCL/AHCL systems may compensate some patients' deficiencies [31, 32], and therefore "democratize" insulin treatment.

Increasing age and time spent for physical activity were similarly linked to optimal metabolic control. In the literature, conflicting results in terms of overall glycemic control have been reported with exercise in individuals with type 1 diabetes, with some studies demonstrating benefits [33–35] and others no improvement in HbA1c following aerobic or resistance training [36, 37]. To achieve enhanced glycemic control while avoiding hypoglycemia, a skillful balance of insulin dosing and food intake is required before, during, and after exercise [38]. Technological advances may help accomplish these tasks with less effort [39].

The independent association of diabetes duration with device satisfaction is also intriguing, in agreement with recent research showing better technology utilization in patients with long-standing disease [40].

The major strengths of our study are the large cohort of participants, the consecutive enrolment, and the great number of outcomes and characteristics that were considered. However, there are some limitations that should be acknowledged. First of all, the cross-sectional design formally prevents causal and temporal inferences between treatments modalities and their glycemic and psychosocial correlates. However, randomized clinical trials and their meta-analyses have extensively clarified that HCL/AHCL systems lead to unprecedented improvements in both aspects of diabetes management [10, 11]. In this scenario, our research shows that treatment goals are achieved also in the Italian real-world setting, therefore providing a reassuring insight to both healthcare professionals and payers. Second, participants on MDI+SMBG were few in number as compared with other groups. In this regard, it has to be considered that our research was conducted in centres with high levels of uptake of diabetes technologies and high expertise in this field, where technology naïve patients are undoubtedly a minority. Third, as laboratory-measured HbA1c levels were not available for the majority of participants, these data were not analyzed. Finally, we used a single tool for the assessment of PROMs, and some important aspects including fear of hypoglycaemia, sleep quality, and diabetes distress were not evaluated. However, the DIDS scale is a short and easy-to-administer tool that is adequate for use across all insulin delivery devices, and has shown robust psychometric properties in individuals with type 1 diabetes [13].

Conclusion

In adults with type 1 diabetes from different areas of Italy, real-life use of advanced technologies for glucose monitoring and/or insulin delivery, particularly HCL/AHCL systems, is associated with improved glucose metrics and device satisfaction. In this population, education attainment, but not family income, may impact on glycemic outcomes. While a definitive cure for type 1 diabetes is not yet achievable, it is crucial to increase the uptake of the most efficacious treatment options to everyone who can benefit from it.

Supplementary Information The online version contains supplementary material available at <https://doi.org/10.1007/s00592-024-02381-3>.

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Data availability Aggregated data might be made available upon reasonable request via email to the corresponding author.

Declarations

Informed consent All participants signed informed consent forms prior to inclusion in the study.

Competing interests The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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