

Psychological well-being of parents of very young children with type 1 diabetes - baseline assessment

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Conflict of interest statement

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest

Author contribution statement

RH and NB coordinated the study. RH, SH, EFR, TK, CLA, CdB, FC, BR, JKM, MEW and MT designed the study. MT, JMA, KN, MF, JY, EM, DS, MF, US, AGT, DA, SS, SH, EFR, TK, CLA, CdB, FC, and BRM screened and

enrolled participants and arranged informed consent from the participants. MT, JMA, KN, MF, JY, EM, DS, MF, US, AGT, DA, JKM, HK, SS, SH, EFR,

TK, CLA, CdB, FC, and BRM provided patient care and took samples. JS managed randomisation. NC, JS, CdB and IPtC did or supported data analyses, including the statistical analyses. RH designed and implemented the glucose controller. CdB and IPtC wrote the manuscript. All authors critically reviewed the report.

CdB and IPtC had full access to all of the data in the study and take responsibility for the integrity of the data and the accuracy of the data analysis.

Keywords

Hypoglycemia fear, Parental well-being, very young children, type 1 diabetes, Child Behavior

Abstract

Word count: 235

Background: Type 1 diabetes in young children is a heavy parental burden. As part of pilot phase of the KIDSAP01 study, we conducted a baseline assessment in parents to study the association between hypoglycemia fear, parental wellbeing and child behavior.

Methods: All parents were invited to fill in baseline questionnaires: hypoglycemia fear survey (HFS), WHO-5, Epworth Sleepiness Scale and Strength and Difficulties Questionnaire (SDQ).

Results: 24 children (median age: 5-year, range 1-7 years, 63% male, mean diabetes duration: 3 ± 1.7 years) participated. 23/24 parents filled out the questionnaires. We found a higher score for the hypoglycemia fear behavior 33.9 ± 5.6 compared to hypoglycemia worry 34.6 ± 12.2 . Median WHO-5 score was 16 (8 - 22) with poor well-being in two parents. Median daytime sleepiness score was high in five parents (>10). For six children a high total behavioral difficulty score (>16) was reported. Pro social behavior score was lower than normal in six children (<6). Parental well-being was negatively associated with HSF total (r = -0.50, p < .05) and subscale scores (r = -0.44, p < .05 for HSF-Worry and HSF-Behavior), child behavior (r = -0.45, p = .05) and positively with child age (r = 0.58, p < .01). HSF, parental well-being nor daytime sleepiness are associated with the HbA1c. Conclusion: Regular screening of parental well-being, hypoglycemia fear and child behavior should be part of routine care to target early intervention.

Contribution to the field

Type 1 diabetes in young children is a huge parental burden. As part of a pilot phase of a hybrid closed loop study (Kidsap02), we conducted a baseline assessment with questionnaires in parents to study the association between hypoglycemia fear (HSF), parental wellbeing (WHO-5) and child behavior (Strength & Difficulties Questionnaire). In this assessment parental well-being was negatively associated with HSF total (r = -0.50, p < .05) and subscale scores (r = -0.44, p < .05 for HSF-Worry and HSF-Behavior), and child behavior (r = -0.45, p = .05). Based on this outcome, regular screening of parental well-being, hypoglycemia fear and child behavior should be part of routine care. An assessment with questionnaire may be useful to evaluate impact of new therapeutic approach.

Funding statement

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Ethics statements

Studies involving animal subjects

Generated Statement: No animal studies are presented in this manuscript.

Studies involving human subjects

Generated Statement: The studies involving human participants were reviewed and approved by Cambridge East Research Ethics Committee in the UK, the Ethics Committee of the University of Innsbruck in Austria, the Ethics Committee of the University of Vienna in Austria, the Ethics Committee of the University of Graz in Austria, the Ethics Committee of the Medical Faculty of the University of Leipzig in Germany, the Comité National d'Ethique de Recherche in Luxembourg. Written informed consent to participate in this study was provided by the participants' legal guardian/next of kin.

Inclusion of identifiable human data

Generated Statement: No potentially identifiable human images or data is presented in this study.



Data availability statement

Generated Statement: The datasets presented in this article are not readily available because Questionnaire data. Requests to access the datasets should be directed to bcschim@pt.lu.



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Parental well-being of children with diabetes

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KEY WORDS: Hypoglycemia fear, parental well-being, very young children, type 1 diabetes, child behavior

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ABSTRACT

Background: Type 1 diabetes in young children is a heavy parental burden. As part of pilot phase of the KIDSAP01 study, we conducted a baseline assessment in parents to study the association between hypoglycemia fear, parental well-being and child behavior.

Methods: All parents were invited to fill in baseline questionnaires: hypoglycemia fear survey (HFS), WHO-5, Epworth Sleepiness Scale and Strength and Difficulties Questionnaire (SDQ).

Results: 24 children (median age: 5-year, range 1-7 years, 63% male, mean diabetes duration: 3 ± 1.7 years) participated. 23/24 parents filled out the questionnaires. We found a higher score for the hypoglycemia fear behavior 33.9 ± 5.6 compared to hypoglycemia worry 34.6 ± 12.2 . Median WHO-5 score was 16 (8 - 22) with poor well-being in two parents. Median daytime sleepiness score was high in five parents (>10). For six children a high total behavioral difficulty score (>16) was reported. Pro social behavior score was lower than normal in six children (<6). Parental well-being was negatively associated with

<u>HFS</u> total (r = -0.50, p < .05) and subscale scores (r = -0.44, p < .05 for <u>HFS</u>-Worry and <u>HFS</u>-Behavior), child behavior (r = -0.45, p = .05) and positively with child age <u>and diabetes duration</u> (r = 0.58, p < .01, r = 0.6, p < .01). <u>HFS</u>, parental well-being nor daytime sleepiness are associated with the HbA1c.

Conclusion: Regular screening of parental well-being, hypoglycemia fear and child behavior should be part of routine care to target early intervention.

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Psychological well-being of parents of very young children with type 1 diabetes

- baseline assessment

Introduction

International recommendations propose a HbA1c of <7.0 % (<53 mmol/mol) in persons with type 1 diabetes (T1D) to prevent acute and late complications¹. Achieving near normal glucose values requires a continuous adjustment of insulin treatment around food intake, daily activities and hormonal changes. In preschool children these choices are determined by the parent/caregiver and pose continuous challenges and stress².

Especially the fear of hypoglycemia has been associated with increased stress in parents of younger children^{3–5}. Fear of hypoglycemia can lead to frequent nocturnal blood glucose measurements and subsequent sleep deprivation⁶, which in turn may affect parents' well-being as well as the (perceived) behavior of their child^{7,8}. New technology such as insulin or sensor-augmented pumps should improve metabolic control. Better control may reduce (perceived) child behavioral problems⁹ and parental anxiety around high and low blood glucose excursions^{7,10}. Regardless of treatment regimen, parents/caregivers of young children with T1D often report their anxiety caused by glucose fluctuations during the night^{11,12}. This suggests that these technologies, although providing improved metabolic outcome, may not yet alleviate parental stress levels.

Next steps in technology advancement include the use of the artificial pancreas, where different algorithms are used to steer insulin administration¹³. An initial evaluation of the use of the Cambridge algorithm in toddlers demonstrated the feasibility of hybrid closed-loop insulin delivery in young children with diabetes¹⁴, whereby parents reported reduced diabetes management burden and improved

sleep quality¹⁵. The current paper presents data from this baseline assessment with a specific focus on parental well-being in relation to fear of hypoglycemia.

Methods

The KIDSAP01 study (NCT03101865) aims to evaluate a hybrid closed-loop insulin administration system, based on the Cambridge algorithm, under free-living conditions in very young children with T1D. A feasibility study was conducted to pretest the setup of the large-scale trial and to determine the specific needs of young children with T1D and their parents. In regards to the latter, parents of children participating in study completed questionnaires on hypoglycemia fear, parental well-being, daytime sleepiness and child behavior at baseline.

Participants

The patient population has been described in detail previously 14,15 . To summarize, 24 children (median age 5 years, range 1-7 years, 63% male) with T1D from four countries (UK, Luxembourg, Germany and Austria) and seven clinics were included. Mean diabetes duration was 3.0 ± 1.7 years (range 0.5-6.4 years). All children were treated for at least three months with insulin pumps.

The majority of children (n=18) used the Medtronic 640G pumps, 4 had a

Medtronic 554 pump, 1 used an Animas Vibe and 1 a Roche Insight. The

Medtronic Guardian REAL time sensor was used by 15 children, the ABBOTT

LIBRE, was used by 4 children and 3 children did not ware a sensor at baseline of this study.

HbA1c levels at baseline were used as an indicator of metabolic control (mean HbA1c, $7.4 \pm 0.7 \%$ (57 ± 8 mmol/mol) and range 5.9-8.7% (41 - 72 mmol/mol). In the three months prior to the study, no severe hypoglycemia nor diabetic

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ketoacidosis had been observed. Twenty-three of the 24 parents filled out the questionnaire at the start of the study.

Measures

The Hypoglycemia Fear Survey (HFS) for Parents¹⁶ is a 26-item survey comprised of a HFS-worry and a HFS-behavior subscale. All items are scored on a 5-point Likert scale (1 = "never"; 5 = "always"). Higher scores reflect higher hypoglycemia fear.

The WHO-5 questionnaire has been extensively used to assess well-being in persons with diabetes and their family¹⁷. It includes five items, with item scores ranging from 0 to 5. Scores under 13 are indicative of poor well-being and a score of below 7 warrants further testing for depression¹⁸.

The Epworth Sleepiness scale evaluates sleepiness and risk to doze off during the day¹⁹. It includes eight questions, scored between 0 and 3, whereby total scores > 9 indicate excessive sleepiness.

The strength and difficulties questionnaire is a screening tool to identify psychosocial problems in (pre-)school children^{20–22}. It includes 25 items comprising four subscales measuring difficulties (emotional, conduct or peer problems and hyperactivity), and a pro-social subscale rating the strengths. The total score is based on the first 4 subscales, whereby a total score > 15 is indicative of a high risk for psychosocial problems²³.

Statistical analyses

First descriptive data will be reviewed to identify the percentage of parents scoring within (ab)normal ranges on the standardized measures. In addition, correlational analyses will be conducted to identify relationships between indicators of parental well-being, and child metabolic control and behavior.

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Based on previous research we expected positive relationships between

parental well-being, and metabolic control and child behavior. Given the

exploratory nature of the pilot study, no a-priori power analyses were conducted.

Results

Descriptive results are provided in Table 1. Results suggest that parents' fear of hypoglycemia is signified by their behavior rather than their worries. Most pronounced fear-related behaviors reflected parents making sure their child had access to fast-acting carbohydrates and getting up in the middle of the night to check the child's blood sugar levels. High mean worry scores were reported for low glucose levels whilst child was sleeping and for parent not being around when the child has low blood sugar. Poor well-being was reported by 2/23 (9%) parents, whereas 5/23 (22%) parents scored low on the question 'I woke up feeling fresh and rested'. This question also generated the lowest mean score for the sample as a whole (*M*=2.86, *SD*=1.46). Excessive daytime sleepiness (scores >10) was reported by 5/23 (22%) parents. Although parent reported child behavioral difficulties were on average within the normal range, scores for three children were in the high range (scores 17-19) and for another three in the very high range (i.e., scores ≥ 20). These scores reflect elevated scores on especially the subscales conduct and emotion. Pro-social behavior was very low for two children.

Parental well-being scores were negatively associated with fear of hypoglycemia and child behavior but positively associated with child age and diabetes duration (see Table 2). Parental well-being scores were negatively associated with fear of hypoglycemia total (r = -0.50, p < .05) and subscale scores (r = -0.44, p < .05 for HFS-Worry and HFS-Behavior) and child behavior (r = -0.45, p = .05) but positively associated with child age and diabetes duration (r = 0.58, p < .01, r = -0.58) as the property of the property o

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 $\underline{0.60, p < .01)}$. All other between-measures correlations were not significant (see table 2).

Discussion

Parents of very young children with T1D reported on their worries and behavior in relation to hypoglycemia before commencing a closed-loop intervention. The hypoglycemic fear ratings in the current sample were higher than previously reported for a sample of parents of children with T1D aged 6 and above⁵. HFS-worry subscale scores were however comparable to those reported by parents of children younger than 7 years¹². Parents indicated they especially worry about hypoglycemia during the night and will regularly get up to check the blood glucose values in order to prevent hypoglycemia or reduce fear of hypoglycemia. This confirms previous reports, suggesting that nights are particularly difficult for parents of young children with T1D¹².

No relationship was found between daytime sleepiness and other constructs nor with child age or HbA1c. This lack of association suggests that excessive daytime sleepiness in this group of parents is not directly related to the management of their child's diabetes but could rather be caused by the poorer quality of sleep of the child, general sleep deprivation or irregular sleep patterns²⁴.

Poor well-being was observed in 2/23 (9%) parents __This percentage is lower than previously reported in parents of youth with diabetes, and may indicate that diabetes and its management affects parental well-being differently depending on the developmental stage of the child²⁵. The negative association between nocturnal fear of hypoglycemia and perceived well-being suggest that fear of hypoglycemia may be one of the key contributors to the diabetes burden.

Parental well-being was positively associated with diabetes duration and child

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age, whereby child age and diabetes duration were also highly associated. It may be that caring for a very young child is generally challenging for parents and especially so when the child is diagnosed with T1D. As children grow older, parents may adjust to both parenting and diabetes management, even when this does not reduce hypoglycemia fear. Further research is needed to investigate to what extent this relates to general parenting issues or is affected by the specific challenges associated with caring for a very young child with T1D.

Twenty-six percent of the parents (6/23) reported child behavioral difficulties in the high to very high range, signified by 13% (3/23) scoring very high on subscale emotional difficulties and 17% (4/23) on conduct problems. These findings support previous research, indicating that parents often report higher levels of behavioral difficulties for children with T1D, most notably a higher incidence of anxiety and depression and externalizing problems^{7,8,26}.

A negative association was found between parental well-being and child behavior, but not HbA1c levels. Well-being of parents is considered relevant for the long-term outcome in children with diabetes and reduced parental well-being is associated with an increase in child psychosocial problems and worse glycemic control^{25,27,28}. Although our results confirm the association between parental well-being and perceived child behavior, no association with either construct and HbA1c levels was found. This may be related to the child's age, as the glycemic control in very young children may be mainly determined by the management behavior of the parent/caregiver and less affected by child behavior.

Fear of hypoglycemia was not related to HbA1c levels. This suggests that parental burden is mainly affected by parental anxiety concerning the management of their child's diabetes rather than the actual glycemic control. To

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this extent, it is imperative that intervention programs are offered to support parents in a timely manner and, in effect, improve long-term outcomes of glycemic control, child behavior and parental well-being^{8,26}.

Some limitation should be considered. Results should be interpreted with caution given the small sample size. Although post-hoc power analyses (using G*Power) indicate that the sample size of N=23 provided at least .74 power to detect effect sizes ≥ .44., medium and smaller effects may not have been detected. Furthermore, our data are all based on parent's reflective reporting. In future studies, observational data or diaries could be used to obtain other indications of parents' and children's actual behavior. In addition, the current results are based on cross-sectional data, allowing for the investigations of association. Longitudinal studies are however necessary to investigate causal relationships between the constructs.

Our study also has some specific strength. Using a multicenter and multinational design, our data can confirm that the burden of diabetes in the very young children is universal and needs to be identified and addressed within diabetes care programs to improve long-term diabetes outcomes and prevent psychological /behavioral complications. One option may be the use of hybrid closed-loop systems with age appropriate algorithms, reducing nocturnal risk of hypoglycemia and glucose variability. Such intervention has been found to reduce diabetes burden 15,29. In addition, family-based interventions, aimed to support parents in managing their child's diabetes have had positive effects on glycemic control as well as parent-child relationships 26.

Author Contributions

RH and NB coordinated the study. RH, SH, EFR, TK, CLA, CdB, FC, BR, JKM,

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MEW and MT designed the study. MT, JMA, KN, MF, JY, EM, DS, MF, US, AGT, DA, SS, SH, EFR, TK, CLA, CdB, FC, and BRM screened and enrolled participants and arranged informed consent from the participants. MT, JMA, KN, MF, JY, EM, DS, MF, US, AGT, DA, JKM, HK, SS, SH, EFR, TK, CLA, CdB, FC, and BRM provided patient care and took samples. JS managed randomisation. NC, JS, CdB and IPtC did or supported data analyses, including the statistical analyses. RH designed and implemented the glucose controller. CdB and IPtC wrote the manuscript. All authors critically reviewed the report.

CdB and IPtC had full access to all of the data in the study and take responsibility for the integrity of the data and the accuracy of the data analysis.

Declarations of interest

MT reports having received speaker honoraria from Minimed Medtronic and Novo Nordisk.

MFr has received speaker honoraria from Minimed Medtronic and has served on advisory boards for Eli Lilly.

JKM is a member in the advisory board of Becton-Dickinson, Boehringer Ingelheim, Eli Lilly, Medtronic and Sanofi, and has received speaker honoraria from ABBOTT, Diabetes Care, Astra Zeneca, Eli Lilly, Nintamed, Novo Nordisk, Roche Diabetes Care, Sanofi, Servier and Takeda.

MEW has received license fees from Becton Dickinson and has served as a consultant to Beckton Dickinson.

SEH declares speaker honoraria from Eli Lilly and Sanofi.

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EFR reports having received speaker honoraria from Minimed Medtronic and Eli Lilly, serving on advisory boards for Eli Lilly.

TMK has received speaker honoraria from Minimed Medtronic, Roche and Eli

Lilly and is member of an advisory board for ABBOTT Diabetes Care.

CdB has received speaker honoraria from Minimed Medtronic, and is member of their European Psychology Advisory Board.

FC does attend Advisory Boards and obtain speaking fees for <u>ABBOTTt</u>, Medtronic, Lilly, and NovoNordisk.

BRM reports having received speaker honoraria from Minimed Medtronic, Eli Lilly, Roche, Menarini and Novo Nordisk, serving on advisory boards for Eli Lilly. RH reports having received speaker honoraria from Eli Lilly, Dexcom and Novo Nordisk, receiving license fees from Medtronic, and being director at CamDiab. IPtC, GM, KD, CB, JMA, KN, JY, EM, DS, MFi, US, AGT, DA, HK, SS, NB, JS, NC, CK, and CLA declare no competing financial interests exist.

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Role of funding source

No sponsor had any role in the study design, data collection, data analysis, data interpretation, or writing of the report.

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Table 1. Descriptive statistics (Mean, Standard Deviation, Median, Range) for scales used in the study

	Mean (SD)	Median	Range
Baseline			
Strengths and Difficulties Questionnaire ^a			
Emotional Problems Subscale Total			7
	2.35 (2.12)	2.00	0 - 7
Score			
Conduct Problems Subscale Total Score	3.26 (2.49)	3.00	0 -10
Hyperactivity Subscale Total Score	4.48 (2.15)	4.00	0 – 8
Peer Problems Subscale Total Score	1.35 (1.56)	1.00	0 – 5
Prosocial Subscale Total Score	7.13 (2.22)	8.00	1 – 10
Total Difficulties Score	11.44 (6.65)	11.00	2 – 25
Hypoglycemia Fear Survey Total Score ^b	68.52 (16.16)	70.00	44 - 105
Behavior Subscale Total Score	33.91 (5.58)	35.00	22 – 44

Worry Subscale Total Score	34.61 (12.18)	32.00	20 – 62
WHO-5 Total Score ^C	16.78 (4.08)	16	8 - 22
Epworth Sleepiness Scale Total Score ^d	6.83 (5.38)	6.00	0 - 24

^a Subscales are on scale 0-10. Total difficulties score is on scale 0-40. For all subscales except the prosocial subscale and total scale, a higher score denotes more difficulties and fewer strengths. For the prosocial subscale, a higher score denotes more strengths and fewer difficulties. For this score, a higher score denotes more difficulties and fewer strengths.

^b Total score is on scale 26-130. Behavior subscale is on scale 11-55, and worry subscale is on scale 15-75. Higher score denotes more fear. Mean score for items on the HFSP-B subscale was 3.08 (SD=0.51). Mean score for items on the HFSP-W subscale was 2.31 (SD=0.81).

^d Scale 0-24. Higher score denotes more sleepiness.

Table 2: Spearman rho Correlation between Parental Well-being, HbA1C,
Child Age, Diabetes duration, Hypoglycemia Fear, Child Behavior and
Sleepiness (N=23)

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Correlation between Parental well-being,
hypoglycemia fear, child behavior and
sleepiness (N=23)[1]

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HbA1C	1.00	.41	<u>.26</u>	<u>.21</u>	<u>38</u>	<u>24</u>	<u>27</u>	.05	Carine De Beaufort 18/7/21 11:46
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^C Scale 0-25. Higher score denotes better well-being.

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*p<.05 **p<.01 ***p<.001

WHO5= Well-being measure; HFS-B = Hypoglycemia fear behavior; HFS-W =

Hypoglycemia fear worry; HFS-TOT = Hypoglycemia fear total; ESS =

Epworth sleepiness scale; SDQ-TOT = SDQ Total Score

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