Health-related quality of life in 4-to-6-year-old children with type 1 diabetes mellitus estimated by children and their mothers

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Abstract
Administration of pediatric Health Related Quality of Life (HRQoL) inventories frequently assesses both the child and parent perspectives in young children with type 1 diabetes mellitus (T1DM), but parent-proxy and child self-reports may differ, and little is known on these discrepancies. The aim is to evaluate HRQoL estimated by young children with T1DM and by their mothers, potential discrepancies in the children-maternal estimates and the factors influencing these discrepancies. Thirty-five 4-to-6-year-old children (19 boys) with T1DM admitted to the Pediatric Endocrinology Department were approached with the self-report KINDL questionnaire for children aged 4–6 years (Kiddy-KINDL for children). Their mothers were approached with the parental version (Kiddy-KINDL for parents). Both versions enable measuring child HRQoL in physical, emotional wellbeing, self-esteem, family, friends, everyday functioning, and the disease dimensions, as well as KINDL total on a 0–100 scale. Statistically significant differences were found between children’s and maternal estimates on the KINDL total and “Disease” scales, in that the maternal proxy-reports produced lower values. A statistically significant difference between self- and proxy-reports was found for the KINDL “Emotional wellbeing” scale values, and the maternal proxy-reports yielded higher estimates compared with children’s self-reports. These associations remained significant after adjustment for major potential confounders. Maternal education, maternal marital status, insulin regimen, and achievement of glycemic control modified the effect of child-maternal discrepancies.

Conclusion: Attempts should be made to improve parental understanding of child problems related to his/her disease with due account to individual family social and demographic characteristics.

What is Known:
• HRQoL in children with T1DM has been advocated as an important complementary outcome to clinical and laboratory markers.
• Self-and parental proxy-reports on HRQoL may differ, but little is known on these discrepancies and on the factors influencing them in young children with T1DM.

What is New:
• Mothers tend to underestimate general and disease-related components of HRQoL but likely to overestimate psychological wellbeing of their ill young children with T1DM.
• Maternal education, marital status, insulin regimen, and achievement of glycemic control modify estimations of HRQoL and child-maternal discrepancies.

Keywords Children · Diabetes mellitus · Health related quality of life · KINDL

Abbreviations
ANOVA Analysis of variance
BMI Body mass index
HbA1c Glycated hemoglobin
HRQoL Health-related quality of life
T1DM Type 1 diabetes mellitus

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Diabetes mellitus is a group of metabolic diseases characterized by chronic hyperglycemia resulting from defects in insulin secretion, insulin action, or both. Type 1 diabetes mellitus (T1DM) continues to be the main type of diabetes encountered in children and adolescents. Over 85% of all diabetes cases in individuals aged < 20 years worldwide are T1DM [1]. T1DM is the most common childhood chronic illness [2], and, in some reports, its incidence is increasing most rapidly among children under 5 years of age [3]. Being diagnosed with T1DM permanently changes the life of children. Treatment has some requirements such as frequent insulin injections, daily blood glucose monitoring, diet plan, and regular physical activity. Also, acute and chronic complications related to diabetes may occur. Normative dependence upon parental caretaking in early childhood translates into considerable parental responsibility for the completion of daily diabetes management tasks [4]. All of these factors can adversely affect the health-related quality of life (HRQoL) in children and adolescents with T1DM [5–9].

Health-related quality of life (HRQoL) is an important construct assessing the impact of a medical condition [10]. It is a multidimensional concept including wellbeing in terms of patient’s physical, emotional, mental, and social behaviors and is defined as the way the effects of a disease and/or its treatment are perceived by the patient [11]. It is agreed that enhancing quality of life and wellbeing is as important as metabolic control and prevention of secondary morbidity [12]. There is consensus on the importance of psychosocial support of children and adolescents to reach treatment recommendations, and HRQoL has been advocated as an important complementary outcome to clinical and laboratory markers [13, 14]. Furthermore, a satisfactory HRQoL score is shown to be a marker of people’s capacity and competence to manage their diabetes treatment and achieve treatment goals. Early detection of poor HRQoL scores is considered important, as it is hypothesized that these children are at risk of psychological maladjustment, reduced compliance and adherence to treatment, and poor metabolic control [15].

Wellbeing can be described in different forms by individuals, and the disease process may also be experienced differently. When evaluating quality of life, it should be considered that there are objective and subjective areas of HRQoL. Two people in the same situation objectively may have different perceptions of their HRQoL subjectively. Some researchers suggest that subjective assessment is more valuable because it reflects self-perception about the situation of individuals. Therefore, the evaluation of HRQoL perceived by parents as well as by the children is important to understand the children’s HRQoL correctly. Parents of young children may experience anxiety and distress related to coping with the particular diabetes responsibilities for this age group [16]. The pediatric transactional theory posits reciprocal interactions between children’s health and behavior and parents’ characteristics and experiences [17]. Therefore, administration of pediatric HRQoL inventories frequently assesses both the child and parent perspectives (proxy) of the child’s HRQoL. However, parent-proxy and child self-reports often differ [18, 19]. Little is known about what discrepant scores mean and what they may indicate in relation to child health outcomes, especially in young children. This “proxy problem” has been debated in the HRQoL literature [18; 20], but no conclusion has been reached on how to handle the apparent discrepancies. Further, analysis of a variety of demographic and disease-related factors such as age, gender, and education varied in terms of how they associated with the direction of discrepancies across the multiple studies, when reported. The direction and magnitude of HRQoL discrepancies remain unclear, and there are no data on the potential importance of discrepant scores within families in relation to clinical outcomes.

To address the issue and bridge the gap, this study aimed at evaluation of HRQoL estimated by young children with T1DM and their mothers and at evaluation of potential discrepancies in the children-maternal estimates and the factors influencing these discrepancies.

Patients and methods

Participants

Participants were recruited from children aged 4 to 6 years with diagnosed T1DM who were under supervision at the Pediatric Endocrinology Department of the V.A.Almazov National Medical Research Centre, St. Petersburg, and who were scheduled for admission to the Department in 2020 for routine check-up and therapy correction. Inclusion criteria were that all children had been diagnosed with T1DM at least 1 year previously to their planned admission, and they represented urban environment. Children with coexisting inborn malformations, chronic diseases other than T1DM, and psychiatric diseases and those with coexisting acute disease during the study were excluded. Ninety-four children met the inclusion criteria. For ethical considerations, we limited the sample size to reasonable number of participants in order to reach a balance between the study’s value and the burdens accepted by its participants. Required sample size computation was performed using G*Power 3.1.5 statistical software [21]. The study was targeted at determining a medium standardized effect size for the differences between two means (Cohen’s d = 0.5), based on the two-tailed test aimed at finding difference between two dependent means (matched pairs), with the α error equal to 0.05, and the power (1–β) equal to 0.8. This yielded the desired sample size equal to 34 children. Ninety-four eligible candidates
for selection were numbered in sequence in advance, and 40 children were selected from the list as a simple random sample without replacement using WinPepi 3.26 program [22]. On admission, these 40 selected children and their parents were approached with a request to participate in the scientific study. Of the 40 selected, 35 (88%) agreed to take part in the study, and they constituted a study group. Patients from the families refusing participation did not differ from the participants in terms of age, gender, socioeconomic status, duration of diabetes, and glycemic control.

**Measures**

**Clinical and socio-demographic characteristics**

Clinical findings were extracted from the patients’ medical records. The mothers were asked to complete the questionnaires addressing child, maternal, demographic major characteristics. To ensure reliability and to minimize recall bias, attention was paid to the thorough analysis of pre-existing medical records. Demographic variables included child sex, age, birth weight, weight and body mass index (BMI) at study, and number of siblings. Maternal marital status was classified as married, cohabiting, divorces, widowed, or single. Information on parental employment was collected. The maternal educational level was defined as the highest level of completed education and categorized as basic education (elementary school), secondary education (secondary school, college), higher education (institute, academy or university), or incomplete higher education. Clinical variables included duration of diabetes at the time of study, insulin regimen, and glycemic control. Insulin regimen was categorized as insulin pump or basal-bolus administration.

**Laboratory findings**

Blood glucose and HbA1c levels were measured at admission as a part of the routine laboratory investigation in T1DM children. HbA1c was analyzed using Bio-Rad D-10 analyzer. According to current clinical practice consensus guidelines, HbA1c values below 7% were defined as good metabolic control. This target value has been chosen with the aim of avoiding long-term microvascular and macrovascular complications of diabetes while also avoiding severe hypoglycemia and the adverse central nervous system changes associated with both hypoglycemia and hyperglycemia [23].

**Questionnaire for evaluation of HRQoL**

The KINDL questionnaires were used to evaluate HRQoL [24]. The KINDL questionnaire satisfies the demand for taking into account progress during child development and the principle of patient-generated data collection by providing different versions of the questionnaire for different age groups and both a self-report version and a proxy version. The common practice of modifying a measure originally designed for adults to make it suitable for children was avoided in this inventory. The original German KINDL questionnaire was duly validated and translated into several languages, including Russian. The psychometric testing of the KINDL indicate adequate to good reliability and convergent and discriminant validity of this inventory [24], in particular among pre-school children [25, 26], and the KINDL was proved efficient in the studies on HRQoL in children with T1DM [12]. Russian versions of the self-report KINDL questionnaire for children aged 4–6 years (Kiddy-KINDL for children) and for parents of children aged 3–6 years (Kiddy-KINDL for parents) were retrieved (https://www.kindl.org/english/language-versions/russian/) and used in this study.

The self-report version of the Kiddy-KINDL questionnaire consists of twelve Likert-scaled items, two for each of six dimensions: physical wellbeing, emotional wellbeing, self-esteem, family, friends, and everyday functioning (school or nursery school/ kindergarten). The response categories of the Kiddy-KINDL cover 3 levels (1 = never, 2 = sometimes, 3 = very often), and the children are to be questioned in a face-to-face interview. The sub-scales of these six dimensions can be combined to produce a total score.

The parents’ version of the Kiddy-KINDL consists of 24 items, four for each of the six above indicated dimensions, and the response categories cover 5 levels (1 = never, 2 = rarely, 3 = sometimes, 4 = often, 5 = all the time). In order to make up for the potentially lower information content of the self-reported responses by young children, the parents’ version of the Kiddy-KINDL contains a further 22 items which can be treated as a sub-scale in their own right.

An additional sub-scale entitled “Disease” aimed at completion in case of prolonged illness or hospitalization, as in this present study, is included both into self-report and parents’ forms and consists of six items that measure the child’s quality of life with respect to his or her illness. Similar to other sub-scales, the response categories cover 3 levels in the self-report version, while they cover 5 levels in the parent’s version.

The scores achieved on the individual KINDL sub-scales and the KINDL total score represent a quantification of the subject’s health-related quality of life from the respondent’s point of view. Higher sum score values are indicative of better HRQoL.

For each item, the children were asked to mark the response that came closest to their own personal experiences. Mothers of the children were asked to complete the KINDL questionnaire as proxies, i.e., to judge the children’s quality of life from their own point of view.
The KINDL questionnaires were analyzed by adding the item responses marked on each sub-scale. Certain KINDL items are worded in such a way that a higher item score implies a poorer health-related quality of life. Reversing the values of these items was necessary in order to ensure that higher scores correspond to a higher HRQoL for all the KINDL items and sub-scales [27].

To ensure compatibility between self-report and parents’ report, scores were summarized and transformed to a 0–100 scale using the following suggested formula [27]:

\[
[(\text{Sub-scale score} - \text{lowest possible score})/\text{possible range of raw score}] \times 100.
\]

### Statistical analysis

Descriptive and analytical statistics were used in data analysis. The internal consistency of both tools (self-report and parent’s KINDL versions) was assessed by Cronbach’s $\alpha$ and McDonald’s $\omega$ coefficients. To avoid dependence on normality of distributions, the Wilcoxon non-parametric matched-pairs signed-rank tests were run to measure differences in the KINDL sub-scales and the KINDL total score according to self- and proxy-reports. The rank-biserial correlation ($r_B$) was considered as a measure of an effect size, coefficients equal to 0.1, 0.3, and 0.5 referring to small, medium, and large effect sizes, respectively [28]. To account for multiple comparisons of the KINDL sub-scales, false discovery rate was computed for each comparison ($q$ value or adjusted $p$) using the Benjamini–Hochberg correction [29]. The mixed-design ANOVA (repeated-measures with a between-factor ANOVA) was further used to test for significant differences between paired measures adjusted for possible confounding effect(s) of several clinical and demographic characteristics and to explore possible interactions, in that the two paired measurements (either self- or proxy-report) were entered into the model as the repeated measurements, while the confounding/modifying factor in consideration was included into the model as either a between-subject factor (for a category variable) or as a covariate (for a continuous variable) [30]. Partial $\eta^2$ was used a measure of the effect size with the values equal to 0.01, 0.06, and 0.14 referring to small, medium, and large effect sizes, respectively [28]. The models were checked for potential violation of homogeneity of variance assumption using Levene’s test. When an effect of a between-subject factor was found significant, post hoc pairwise comparisons with Tukey correction were performed.

The value for significance was set at $p < 0.05$. Statistical analyses were performed using the JAMOVI statistical software version 1.6 [31].

### Results

Table 1 is the summary of the major clinical and demographic characteristics of the patients and families. The study comprised 16 girls and 19 boys with disease duration ranging between 1 and 5 years (mean 2.1). Twenty-two patients received basal-bolus insulin, while 13 were on insulin pump. The HbA1c level in the patients ranged between 5.5 and 11.2% (mean 7.4); in 19 children, it was below 7%, which was considered as an achieved glycemic control. Children and their mothers did not have any problems in understanding the language, the meaning of the questions in the KINDL questionnaires, and in selecting appropriate answers. The values of internal consistencies of the

| Overall ($N = 35$) | Sex | Girls | 16 (46%) | Boys | 19 (54%) |
|-------------------|-----|-------|---------|------|---------|
| Age, years       | 5.0 (1.0) |
| Birth weight, g   | 3118 (321) |
| Weight at study, kg | 21 (4.5) |
| Height at study, cm | 115 (10.3) |
| BMI, kg/m²        | 16.6 (2.9) |
| BMI z-score*      | 1.3 (0.7) |
| T1DM duration, years | 2.0 (1.5) |
| HbA1c, %         | 6.9 (2.5) |
| Glycemic control (HbA1c < 7%) | Yes | 19 (54%) | No | 16 (46%) |
| Insulin regimen   | Basis-bolus | 22 (63%) | Insulin pump | 13 (37%) |
| Siblings          | 0 | 20 (57%) | 1 | 8 (23%) | 2 | 5 (14%) | 3 | 2 (6%) |
| Marital status    | Married | 22 (63%) | Divorced | 7 (20%) | Widow | 1 (3%) | Cohabiting | 5 (14%) |
| Maternal education| Higher | 21 (60%) | Incomplete higher | 11 (31%) | Secondary | 3 (9%) |
| Maternal age at study, years | 27.0 (5.0) |

*Values are based on WHO standards (birth to 60 months) and WHO reference 2007 (61 months to 19 years)
self-report KINDL and parent’s KINDL questionnaires and their specific sub-scales were acceptable-to-good ranging between 0.695 and 0.953.

Table 2 shows the distributions of self- and proxy-reports on total KINDL questionnaire and different KINDL scales. Overall, a statistically significant difference was found in total quality of life estimates between a child and his/her mother, in that the maternal proxy-reports produced lower values. Likewise, compared with children, the mothers reported statistically significantly lower values on the KINDL Disease scale. Large effect sizes were found for these differences. A statistically significant difference between self- and proxy-reports was found for the KINDL “Emotional wellbeing” scale values with large effect size; however, unlike the KINDL total and the KINDL Disease scores, the maternal proxy-reports on the KINDL Emotional wellbeing score yielded higher estimates compared with children’s self-reports. A statistically significant difference between self- and proxy-reports was also found for the KINDL “Physical functioning” scale values, but this difference failed to retain its significance after correction for multiple comparisons.

The mixed-design ANOVA was further performed to test for significant differences between paired measures on total KINDL scores, the Disease and the Emotional wellbeing scores adjusted for possible confounding or modifying effect(s) of several clinical and demographic characteristics, including child sex, T1DM duration, siblings in a family, maternal education, maternal marital status, insulin regimen, and glycemic control. The two paired measurements (self- and proxy-reports) were entered into the model as the repeated measurements, while the confounding/modifying factor in consideration was included into the model as a between-subject factor (if it was a categorical) or as a covariate (if it was a continuous).

No statistically significant effects on the differences between the paired total KINDL measurements and the paired KINDL Disease measurements were found for child sex, T1DM duration, siblings in the family, and maternal age at study. Statistically significant within-subject effects related to the source of information (either self- or proxy-report) remained significant after adjustments for the above between-subject variables. A statistically significant effect on the total KINDL measurements was found for the maternal marital status as a between-subjects variable but was not found for the paired KINDL Disease measurements (Supplementary Table 3). The two paired total KINDL measurements (either self- or proxy-reports) retained its significance as a source of within-subject variance after marital status was included into the model. There was also a statistically significant interaction between the sources of total KINDL reports and maternal marital status (Fig. 2A). Following up this interaction indicates that the children and their mothers provided higher estimates on the total KINDL score when the mothers were married. However, the contrast between the self- and the proxy-estimates was more prominent when the mothers were married. Post hoc pairwise comparisons found that the values of the KINDL total scores were the highest in the self-reports of the children of the married mothers, and these values were statistically significantly higher than the values in the children whose mothers were not married (mean difference = 19.02, t(45.9) = 3.21, \( p_{\text{tukey}} = 0.012 \)). Statistically significant between-subjects effect was also found for maternal education (either higher or not) (Supplementary Table 3). The source of information on the KINDL total score (either self- or proxy-report) as a within-subjects effect retained its significance after adjustment for maternal education. There was also a statistically significant interaction between the source of report and maternal education (Fig. 2B). Following up this interaction indicates that the maternal estimates of the KINDL total score were always lower than those of their children, more so for those in maternal higher education. In the cases of maternal higher education, both children and their mothers reported lower values on the KINDL total score. Post hoc pairwise comparisons found that the children of the mothers

| KINDL scale       | Self-report | Proxy-report | \( P \) (Wilcoxon \( W \)) | \( q \)  | Effect size (rank biserial correlation) |
|-------------------|-------------|--------------|----------------------------|--------|----------------------------------------|
| The KINDL total score | 69.4 (27.8) | 42.1 (20.8)  | <.001                      | 0.007  | .886                                   |
| Physical functioning | 75.0 (37.5) | 56.3 (12.5)  | .031                       | 0.055  | .424                                   |
| Emotional wellbeing | 75.0 (62.5) | 87.5 (31.3)  | .007                       | 0.025  | .540                                   |
| Self-esteem       | 69.0 (25.0) | 62.2 (12.3)  | .072                       | 0.102  | .349                                   |
| Family            | 75.0 (50.0) | 81.3 (15.6)  | .112                       | 0.132  | .333                                   |
| Friends           | 78.0 (37.5) | 81.3 (21.9)  | .558                       | 0.564  | .126                                   |
| Everyday functioning | 75.0 (50.0) | 62.5 (18.8)  | .788                       | 0.696  | .058                                   |
| Disease           | 53.2 (37.5) | 41.7 (33.3)  | .019                       | 0.045  | .454                                   |

Statistically significant differences marked in bold
with higher education had statistically significantly lower values on the KINDL total score than the children whose mothers had educational level other than higher (mean difference = −19.84, t(46.1) = −3.36, \( p_{\text{Tukey}} = 0.008 \)). A statistically significant between-subjects effect was found for the insulin regimen (insulin pump vs. routine base-bolus administration) (Supplementary Table 3). The source of information on the KINDL total score (either self- or proxy-report) as a within-subjects effect retained its significance after adjustment for the insulin regimen. There was no statistically significant interaction between the paired differences in child-maternal reports and the insulin regimen. Therefore, the contrasts between the self- and the proxy-reports were similar under any insulin regimen, but the children receiving insulin via insulin pump and their mothers reported higher values on the KINDL total score than the pairs when the children were on a routine insulin administration. Post hoc pairwise comparisons found that the children on insulin pump had statistically significantly higher values on the KINDL total score than the children on a routine insulin regimen (Mean difference = 18.12, t(47.0) = 3.01, \( p_{\text{Tukey}} = 0.021 \)). Similarly, glycemic control was found to have a significant between-subjects effect, and paired within-subject differences retained significance after adjustment for glycemic control (Supplementary Table 3). The level of HbA1c was found to have a statistically significant effect when it was included into the model as a continuous covariate rather than a dichotomized “glycemic control” variable (\( p < 0.001 \)). Statistically significant interaction was found between glycemic control and the source of information. In the cases of accomplished glycemic control, both children and their mothers reported higher values on the KINDL total score. Post hoc pairwise comparisons found that the children who have achieved glycemic control reported statistically significantly higher estimates on the KINDL total score than the children in whom this control has not been accomplished (mean difference = 26.74, t(49.2) = 5.13, \( p_{\text{Tukey}} < 0.001 \)); likewise, the mothers of those children who have reached glycemic control reported significantly higher estimates than the mothers of the children without glycemic control (mean difference = 15.16, t(49.2) = 2.91, \( p_{\text{Tukey}} = 0.027 \) (Figs. 1D and 2).

Statistically significant between-subjects effect was found for the maternal education (either higher or not), and the source of information on the KINDL Disease score (either self- or proxy-report) as a within-subjects effect retained its significance after adjustment for maternal education (Supplementary Table 3). No statistically significant interaction between the source of information and maternal education was found. Both the children of the mothers with higher education and their mothers reported lower values on the KINDL Disease score than those pairs where the mothers did not have higher education (Fig. 2A). A statistically significant between-subjects effect was also found for the insulin regimen (insulin pump vs. routine administration). The source of information on the KINDL Disease score (either self- or proxy-report) as a within-subjects effect retained its significance after adjustment for the insulin regimen. There was a statistically significant interaction between the paired differences in child-maternal reports on the Disease score and the insulin regimen (Supplementary Table 3). Following up this interaction indicates that the estimates were higher
both in the children who were on a pump regimen and in their mothers, and the contrast between the estimates in these pairs was insignificant. The estimates in children on a routine insulin regimen and in their mothers were lower, and in these pairs, the contrast between the self- and the proxy-estimates was more remarkable and significant. The mothers of children on a routine insulin regimen provided statistically significantly lower estimates than the mothers whose children were on a pump, mean difference = 31.2, \( t(45.7) = 3.651, p_{\text{Tukey}} = 0.004 \) (Fig. 2B). Glycemic control was found to have a significant between-subjects effect, and paired within-subject differences retained their significance after adjustment for glycemic control. The level of HbA1c was found to have a statistically significant effect when it was included into the model as a continuous covariate rather than a dichotomized glycemic control variable \((p < 0.001)\). There was no statistically significant interaction between glycemic control and the source of information on the KINDL Disease score. In the cases of accomplished glycemic control, both children and their mothers reported higher values on the KINDL Disease scores than in the cases where glycemic control was not achieved. Maternal estimates were lower than children’s self-reports both in cases of accomplished and non-accomplished glycemic control (Fig. 2C).

No statistically significant between-subject effects were found when the paired KINDL Emotional wellbeing measurements were considered, and the following variables were explored for the between-subjects effect: child sex, T1DM duration, siblings in a family, and maternal age at study. As well, no statistically significant effect was found for the maternal marital status and insulin regimen. Statistically significant between-subjects effect was found for maternal education (either higher or not) (Supplementary Table 3). The source of information on the KINDL Emotional wellbeing score (either self- or proxy-report) as a within-subjects effect retained its significance after adjustment for maternal education. There was also a statistically significant interaction between the source of report and maternal education (Fig. 3A). Following up this interaction indicates that the maternal estimates of the KINDL Emotional wellbeing score were always higher than those of their children, more so for those in maternal higher education. In the cases of maternal higher education, both children and their mothers reported lower values on the KINDL Emotional wellbeing score. Post hoc pairwise comparisons found that the children of the mothers with higher education had statistically significantly lower values on the KINDL Emotional wellbeing score than the children whose mothers had educational level other than higher \((\text{mean difference} = -32.92, t(57.8) = -4.402, p_{\text{Tukey}} < 0.001)\). Glycemic control was also found to have a significant between-subjects effect, and paired within-subject differences retained their significance after adjustment for glycemic control. The level of HbA1c was found to have a statistically significant effect when it was included into the model as a continuous covariate rather than a dichotomized glycemic control variable \((p < 0.001)\). There was no statistically significant interaction between glycemic control and the source of information on the KINDL Emotional wellbeing score. In the cases of accomplished glycemic control, both children and their mothers reported higher values on the KINDL Emotional wellbeing scores than in the cases where
glycemic control was not achieved. Maternal estimates were higher than children’s self-reports both in cases of accomplished and non-accomplished glycemic control (Fig. 3B).

**Discussion**

This study aimed at comparing self- and maternal proxy-reports on different aspects of HRQoL in young children with T1DM.

One finding from this work was that, compared with their children, the mothers provided lower estimates on the KINDL total and the KINDL Disease scales. This finding is generally in accord with a fact that parents of children without chronic illness typically rate their child’s HRQoL better than the children themselves, while, in contrast, parents of children with a variety of chronic health conditions, including diabetes, typically rate their child’s HRQoL as worse than children themselves [18, 20, 32, 33]. However, across studies, the findings tend to be mixed, with discrepancies commonly reported in both directions [19, 34–37]. Evidently, parents may generally underestimate HRQoL of their ill children due to their concerns about child health, especially considering general health characteristics and disease-specific issues of the quality of life.

An important issue is psychological adjustment in children with T1DM, and the findings were that children with diabetes appear to have a greater incidence of emotional disturbances and psychological distress. Family cohesion, supportive behaviors, and collaborative solving of psychological problems are among the major constituents of care for ill children [5]. In this regard, comparisons between self- and proxy ratings of emotional wellbeing in children with T1DM are of special interest. The findings from this study were that maternal ratings of the HRQoL related to child emotional wellbeing were higher than those in children’s self-reports. Previous findings were that concordance among parents and children was greater for physical functioning than for emotional and school functioning [38]. Discrepancies in parents’ and healthy children’s reports of child emotion regulation were previously reported [39]. Clearly, there is a disconnection between parents’ and children’ perceptions of emotional state in the young age group. Given that parents are doing much of the diabetes care at this stage, many may undervalue the effect diabetes has on their young child’s emotional state. Parents of young children with diabetes may have difficulty in distinguishing diabetes-related feelings, so it may be difficult to accurately assess their child’s emotional well-being [40]. In the context of clinical care, it is important to consider these discrepancies in HRQoL reports as much as possible, since maternal reporting higher scores can indicate a worrisome lack of correspondence between the mother and the child. If providers note large discrepancies, it may be beneficial to dedicate more time to discussing HRQoL issues with families. However, more research is needed to determine whether HRQoL interventions for families with larger versus smaller discrepancies are impactful [41].

An attempt was made to study possible confounding/modifying effects of several clinical and demographic factors on child-maternal discrepancies in HRQoL estimates. Although some studies were indicative that boys with diabetes tended to report better quality of life [42], no statistically significant effect was found for child sex in this study. Better HRQoL estimates were previously reported in youth with longer diabetes duration and in those from a better socioeconomic background [19, 43, 44]. Other risk factors for unsatisfactory quality of life in diabetic children and adolescents were previously reported to be depression, one-parent family, and diabetes-related family conflict [45]. This study failed to find statistically significant effect of diabetes duration on HRQoL; at least in part, this might be due to relatively
short and similar diabetes duration in young children who entered this study. Important findings from this study were that child-maternal discrepancies in HRQoL estimates were modified by such factors as maternal marital status, maternal education, insulin regimen, and glycemic control. Generally, higher values of self- and maternal proxy reports were found in those cases where the mothers were married, had education other than higher, when a child received insulin via insulin pump, and when glycemic control was reached. Lower estimates of the child HRQoL in higher-educated mothers and their children might be due to more critical attitudes toward the child state in these families. Insulin pump therapy is the means of intensive treatment and one of the most technologically advanced methods of achieving near-normal blood glucose levels with numerous benefits ascribed to this regimen, including more flexibility in the timing of patients’ meals, exercise, and physical activity [46]. To date, however, few studies have addressed how the adjustment to pump therapy affects HRQoL of children and adolescents. Some findings were indicative that HRQoL was unrelated to regimen prescription in children [47], while the others found insulin pump therapy to be a significant predictor of the HRQoL [48]. Our own findings are the arguments that insulin pump is related to better estimates of HRQoL, both by young children themselves and by their mothers. It is in accord with the reports on the fact that intensive treatment of diabetes does not worsen quality of life and in some cases can even improve it [12, 45]. Use of the insulin pump does not appear to adversely affect quality of life [12, 47, 49]; instead, it may be associated with improved quality of life, as shown in this present study. In addition, it was found that use of continuous glucose monitoring did not seem to adversely affect quality of life [50].

The relationship between HRQoL and metabolic control, measured by HbA1c, is conflicting [12, 45]. Consistent with our findings, some authors demonstrated that better HRQoL was associated with better glycemic control [51, 52]; poorer HRQoL was associated with higher HbA1c level and greater depressive symptoms in the pediatric T1DM [49, 53]. Diabetes symptoms are associated with general health-related quality of life, which is partially mediated by diabetes management [54].

Contrasts between children’s and maternal reports on the KINDL total estimates in this study were more sharp when the mothers were married, had higher education, and when glycemic control in children was reached. Unlike in this study, some authors found that among adolescents with T1DM, larger discrepancies between self- and parental proxy reports on the generic and diabetes-specific HRQoL modules were associated with poorer glycemic control with higher HbA1c values, in that the youth reported higher scores than their parents [19]. Given adolescents are more likely to have worsening treatment adherence and poor glycemic control [41], it is possible that parental perceptions of adolescent HRQoL may take adherence or glycemic control into consideration, thus rating HRQoL lower when HbA1c is poor; however, teens’ own perceptions of HRQoL may not be related to HbA1c. Differences between parent and youth perspectives on HRQoL that encompass broader family issues, such as family conflict or poor communication in the home, may take place, as both have been associated with HRQoL and HbA1c [55, 56]. By contrast, in young children who may be more compliant with medical and parental recommendations than adolescents and who are more attached to their parents and are closely supervised by them, the reaction of a child and his/her mother to a failure in reaching glycemic control may be similarly frustrating; however, in the event of an achievement of glycemic control, not all maternal expectations about her child progress may be met, while a child may be happy with a success, thus leading to his/her higher estimates of the HRQoL and more discrepancies between self- and proxy reports. Similarly, larger discrepancies between children’s and maternal estimates on the total HRQoL scores in the cases when the mothers were married may be due to higher expectations about child health in these families.

Several limitations of this study should be acknowledged. First, the study comprised limited number of 4-to-6-year-old children from the urban setting; thus, the findings from this study cannot be easily expanded across other children categories. Another limitation may come from certain inadequacies between the child and parent KINDL questionnaire forms: except the KINDL Disease scale where both the child and the parental forms each include 6 questions, the remainder scales were based on 2 questions in the child and on 6 questions in the parental form. It is by this that, when using child self-reports, calculation of the KINDL Total score was considered as more reliable estimate [27]. Meanwhile, acceptable-to-good internal consistencies found in all self- and proxy-report subscales argued in favor of meaningfulness of the values obtained, and the score transformation to a 0–100 scale enabled direct comparisons between children’s and maternal reports. Another subject of concern may be that although the questions from the KINDL questionnaire were addressing situations related to the last week, the answers of the children might have been somehow influenced by what had happened shortly before testing and by the feelings of the child during the test, i.e., the problem of the test reliability. However, previous studies were indicative of appropriate test–retest stability of the KINDL questionnaires in 4–17-year-old children from different settings using different linguistic versions of the questionnaire, in all the subscales and the total score, even when re-testing has been performed 7 to 21 days after the initial test [57–60]. Even admitting that current life events might somehow have influenced responses, such non-differentiated response bias could
not invalidate the results of comparisons performed in this study. We must also take into account that numerous exogenous factors remaining beyond the scope of this study might have influenced HRQoL estimates produced by children and their parents. In particular, the study was conducted under the circumstances of the COVID-19 pandemic. Meanwhile, the pandemic did not interrupt the delivery of ambulatory and hospital medical care to children with diagnosed T1DM. The children and their parents who entered this study were local residents, and thus faced principally similar burdens related to the pandemic.

With all the limitations in mind and with full awareness on the fact that further studies in the field are desirable, the findings from this study support prior recommendations to use the child report in conjunction with the parent-proxy report when making determinations on child HRQoL [19, 40, 56]. When child and parent-proxy scores are discrepant, they should be considered individually. Attempts should be made to improve parental understanding of child problems related to his/her disease with special emphasis on child emotional wellbeing with due account to individual family social and demographic characteristics. Clinical assessment and discussion of HRQoL should be considered as a way to comprehensive management of diabetes spanning both physical and mental health.

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Igor A. Kelmanova: conceptualization and design of the study, analysis of data, draft of the initial manuscript.

Availability of data and material Not applicable.

Code availability Not applicable.

Declarations

Ethics approval All procedures performed in study were in accordance with the ethical standards of the institutional and/or national research committee and with the 1975 Helsinki declaration and its later amendments or comparable ethical standards. The study was approved by the Institutional ethics committee.

Consent to participate Informed consent for the participation in a scientific study was obtained from all mothers involved, and confidentiality was guaranteed.

Consent for publication Not applicable

Competing interests The authors declare no competing interests.

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