CLINICAL RESEARCH

Quality of life in children participating in a non-selective INR self-monitoring VKA-education programme

Qualité de vie des enfants sous AVK participant à un programme d’éducation thérapeutique non sélectif à l’auto-mesure de l’INR

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KEYWORDS
Quality of life; INR

Summary
Background. — The quality of life (QoL) of children receiving vitamin K antagonist (VKA) treatment has been scarcely studied.
Aim. — To assess QoL of children, and its evolution, throughout our non-selective international normalized ratio (INR) self-monitoring education programme.

Abbreviations: CHD, congenital heart disease; M3C, French National Reference Center for Complex Congenital Heart Diseases; QoL, quality of life; VKA, vitamin K antagonist.

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Methods. — Children and parents completed QoL questionnaires (Qualin, PedsQL) during education sessions. Scores were compared with those from controls.

Results. — A total of 111 children (mean ± standard deviation age 8.7 ± 5.4 years) were included over a 3-year period. Indications for VKA treatment were congenital heart diseases (valve replacement [42.3%], total cavopulmonary connection [29.7%], myocardopathy [11.7%], coronary aneurysm [7.2%], venous/intracardiac thrombosis [4.5%], pulmonary artery hypertension [1.8%], arrhythmia [0.9%] and extra-cardiac disease [1.8%]). Eighty children, 105 mothers and 74 fathers completed the QoL questionnaires. QoL was good among children aged 1 – 4 years and moderately impaired in those aged between 5 and 18 years. There was no significant relationship between self-reported QoL and patient’s sex, type of VKA, number of group sessions attended, disease duration or time of diagnosis (prenatal or postnatal). QoL scores were significantly lower among children with congenital heart diseases compared with other diseases. There were few differences in QoL between children under transient VKA treatment and those treated for life. Parental proxy QoL scoring correlated well with but was significantly lower than child self-assessments. QoL reported by mothers increased through the education programme, independently of any improvement of the health condition.

Conclusions. — This QoL study provides original data from a large cohort of children and their parents participating in a formalized INR self-monitoring education programme for VKA treatment.

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Background

More consideration is now given to patient-related outcomes [1], especially in assessing the quality of life (QoL) of children with congenital heart diseases (CHD) [2]. QoL has been defined as the "overall life satisfaction" [3], but most clinical trials require a more operational definition and use health-related quality of life questionnaires [4].

Survival of children with more complex CHD have led to new burdens related to the need for long-term anticoagulation with vitamin K antagonist (VKA) treatment in a variety of conditions. The use of selective international normalized ratio (INR) self-monitoring using a self-measurement device at home has proven safe in children receiving VKA therapy [5–7]. Whereas most patients claim that they are comfortable with self-measurement devices, QoL has been scarcely evaluated in this population. QoL studies in adults have delivered contradictory findings [8–14]. Data from paediatric studies with small cohorts suggest QoL improvement during INR-self-monitoring programmes [15,16], but without any data from the child or their parents.

Our non-selective child-focused education programme appeared to be useful in maintaining efficacy, safety and compliance to anticoagulation and its monitoring [17]. Here, we aimed to assess the QoL of children included in our education programme in comparison with a control population. We also assessed QoL evolution throughout the education programme and compared the child’s with the mother’s and father’s proxy evaluations.

Methods

This prospective cohort study was carried out in Necker M3C national reference centre, Paris, France, with a 3-year recruitment and observation period. Children aged 0 to 18 years who were receiving VKA treatment for any reason, prescribed for >3 months and using an INR self-monitoring device were eligible. All patients and/or parents participating in our INR self-monitoring education programme were invited to complete a generic QoL questionnaire during each education session.

INR self-monitoring and VKA education programme

As previously reported [17], our non-selective educational programme starts with an initial session divided into five parts: VKA theoretical training, self-measurement device practical training (CoaguChek XS), group session (game, discussion), administration of QoL questionnaires, and evaluation. Six months later, a reinforcement (i.e. support) session is organized. Participation in the support session is not mandatory, but is based on the family, child or physician’s demand. The global clinical status of the child (stable, worse, better) was established by a paediatric cardiologist at the last session.

Health-related QoL questionnaires

Two generic paediatric QoL questionnaires were used: the Qualin questionnaire for infants <2 years of age [18] and the PedsQL questionnaire for children aged between 2 and 18 years [19,20]. Both parents could participate and completed their proxy questionnaires separately. We used the following versions of the PedsQL, depending on the patient’s age: proxy reports for children aged 2 to 4 years, and self and proxy reports for children aged 5–7, 8–12, and 13–18 years. PedsQL self-reports were completed by the children under trained nurse supervising. All families were administered a questionnaire after the first session of the education programme, 6 months later, and, when applicable, if VKA was stopped before 6 months.

The Qualin questionnaire is a 34-item generic French QoL instrument designed for infants (aged 0–3 years) and was completed by parents or caregivers. Each answer scores from −2 (totally false) to +2 (quite true). An item with a score >1 corresponds to favourable QoL. The psychometric properties are satisfactory (acceptability 90%, Cronbach’s alpha coefficient >0.75, correlations between two raters >0.50, good construct validity) [18].

The PedsQL generic QoL questionnaire has four multidimensional scales: physical functioning (8 items), emotional functioning (5 items), social functioning (5 items), and school functioning (5 items). The three summary scores are total scale score (23 items), physical health summary score (8 items), and psychosocial health summary score (15 items). Each item uses a 5-point Likert scale from 0 (never) to 4 (almost always). Items are reversed scored and linearly transformed to a 0–100 scale, higher scores indicating a better QoL. Psychometric properties showed reliability, validity and responsiveness to clinical change over time [19]. The translation and cultural adaptation into French was performed by MAPI Research Institute (www.mapi-trust.org) following the international guidelines [21]. The psychometric properties of the French version of the PedsQL appeared to be acceptable [22].

Control population

The normal values from the literature were used for healthy children aged <8 years [18,19,23]. For children aged from 8 to 18 years, we used the data from our own recently reported comparative QoL study: 180 families of children aged 8 to 18 years were recruited in randomly selected schools and completed self and proxy QoL questionnaires [2].

Formal aspects

This study complies with the declaration of Helsinki. It was approved by our regional Ethics Committee (Ile-de-France III) and is registered on ClinicalTrials.gov (number NCT01202916). The Ministry of Education Regional Authority authorized schools’ participation. Informed consent was obtained from all parents.

Statistical analysis

Patient characteristics are presented using mean and standard deviation (SD) for continuous variables and as frequencies and proportions for qualitative variables. Total QoL scores and subscore dimensions were calculated for each questionnaire. QoL scores were compared with the
parametric Student’s t-test when the distribution was Gaussian, and with the Mann–Whitney test otherwise, for the following variables: sex, pathology, time of diagnosis (prenatal or postnatal), type of VKA, duration of VKA prescription (life-long or transient), and type and number of therapeutic education sessions attended. Correlations between time since diagnosis and QoL scores were studied using Pearson’s or Spearman’s coefficient. Comparisons and correlations were performed independently for Qualin and PedsQL, for initial and final evaluations and for children, mothers and fathers. QoL scores were compared between self-reports, maternal reports and paternal reports, and between baseline and final evaluations, using paired Student’s or Wilcoxon tests depending on the distribution. The Qualin and PedsQL internal reliability were evaluated with Cronbach α coefficient.

The two-sided significance level was 0.05. SAS version 9 (SAS Institute, Cary, NC) was used.

Results

Patient characteristics

A total of 111 children participated in our INR self-monitoring programme. No family refused to be enrolled in the QoL study. Data on the indications for VKA, types of VKA and education programme participation are detailed in Table 1. For the 111 children, 426 QoL questionnaires (25 Qualin, 401 PedsQL) were completed by 259 people (80 children, 105 mothers, 74 fathers) (Table 2).

Qualin QoL scores

Two parents of infants aged < 1 year participated in the education programme and completed the Qualin questionnaire. The QoL global score was low (−0.1 ± 0.1) (vs a normal value of 0.9).

Eleven mothers and 10 fathers of children aged > 1 and < 2 years participated in the education programme and completed the Qualin questionnaire. Parents’ proxy QoL global scores were close to normal values: 1.1 ± 0.2 for mothers and 1.2 ± 0.3 for fathers (vs a normal value of 0.9) (Fig. 1A and Supplementary Table 1).

The Qualin internal reliability was satisfactory, with a Cronbach coefficient of 0.89 for mothers, 0.93 for fathers and 0.91 globally.

PedsQL QoL scores

In all three subgroups of age (5–7, 8–12 and 13–18 years), children’s self-reported QoL scores were significantly lower than controls globally and in all dimensions except school and physical dimensions in the group aged 5–7 years and school and emotion dimensions in the group aged 13–18 years (Fig. 1B–D and Supplementary Table 2).

In the group aged 2–4 years, mother-reported proxy scores were significantly lower than controls globally and in all dimensions except social, whereas father-reported proxy scores were not significantly different from controls (Fig. 2A). In the group aged 5–7 years, mother-reported scores were significantly lower than control scores in all dimensions whereas father-reported scores were significantly lower than controls in the school and psychosocial dimensions only (Fig. 2B). In the groups aged 8–12 and 13–18 years, parent-reported QoL scores were significantly lower than controls globally and in all dimensions except emotion for children aged 13–18 years (Fig. 2C and D and Supplementary Table 3).

The PedsQL internal reliability was satisfactory, with a Cronbach coefficient of 0.91. When the questionnaires completed by children, mothers and fathers were separated, internal reliability remained satisfactory, with Cronbach coefficients of 0.88, 0.92 and 0.92, respectively.

Factors associated with PedsQL scores

PedsQL self-reported scores were not affected by sex, type of VKA, type of education sessions attended, duration of VKA prescription (life-long or transient treatment) or time of diagnosis (prenatal or postnatal), at both baseline and final
evaluations. QoL evaluation by mothers and fathers was not impacted by sex or type and number of education sessions (data not shown).

At the final evaluation, mother-reported QoL scores were higher for children treated with warfarin versus fluindione in several dimensions: school (79.9 ± 15.4 vs 67.1 ± 24.2; P = 0.02), physical (80.0 ± 17.5 vs 65.1 ± 24.3; P = 0.03), social (80.4 ± 15.4 vs 69.0 ± 18.2; P = 0.04), and total score (77.1 ± 14.4 vs 66.7 ± 19.5; P = 0.03). At the final evaluation, mother-reported QoL scores were lower for children with prenatal versus post-natal diagnosis in the emotion (53.8 ± 15.3 vs 71.6 ± 20.9; P = 0.005) and psychological (64.5 ± 12.1 vs 74.9 ± 17.0; P = 0.04) dimensions. At baseline, the father-reported physical score was higher for children with long-life treatment than for those with transient treatment (71.4 ± 22.8 vs 55.6 ± 24.2; P = 0.02).

Disease duration was negatively correlated with several father-reported scores at the final evaluation: school (r = −0.57; P < 0.01), emotion (r = −0.49; P < 0.01), physical (r = −0.60; P < 0.001), psychological (r = −0.57; P < 0.001), social (r = −0.55; P < 0.001) and total score (r = −0.62; P < 0.001). Disease duration was negatively correlated with...
the mother-reported physical score at the final evaluation ($r = -0.23; P = 0.05$).

QoL was impaired in children with CHD (heart valve diseases and total cavopulmonary connections, $n = 80$) compared to children with other diseases (Table 3). Almost all dimensions were affected in self, maternal and paternal proxy reports at baseline and in the final evaluations. The QoL of children receiving life-long VKA after valve replacement ($n = 47$) was not significantly different from that of other children for all dimensions and at each step of the programme.

**Evolution of PedsQL scores throughout the education programme**

Mothers’ reported PedsQL scores increased significantly throughout the programme globally and in physical, school and psychosocial dimensions (Table 4). Self-reported and father-reported PedsQL scores did not vary significantly between initial and final sessions. Only a tendency toward increased scores was observed for self-reports in the school dimension and for paternal reports in school and physical dimensions.

No statistical link was found between the change in PedsQL scores throughout the programme and the evolution of the global clinical status of the child (stable, worse, better) evaluated by the paediatric cardiologist ($P > 0.05$ in all dimensions).

**Comparison between child, maternal and paternal reports**

No difference between QoL scores was found when comparing pairs of maternal and paternal reports with the Qualin and PedsQL questionnaires (Table 5) administered at the same time, and for their first administration. Maternal reported PedsQL global scores were significantly lower than child self-reports, with all dimensions affected (Supplementary Table 4). Paternal reported PedsQL global scores were also lower than child self-reports, with only the physical dimension significantly affected (Supplementary Table 4).


Table 3  PedQL scores in children with CHD versus other diseases according to mother-, father- and self-reports at baseline and final evaluations.

<table>
<thead>
<tr>
<th>Dimension</th>
<th>Baseline</th>
<th>Final</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>CHD</td>
<td>Other diseases&lt;sup&gt;a&lt;/sup&gt;</td>
<td>CHD</td>
</tr>
<tr>
<td></td>
<td></td>
<td>P</td>
<td></td>
</tr>
<tr>
<td>School</td>
<td>64.5 ± 19.3</td>
<td>71.0 ± 25.9 0.12</td>
<td>65.3 ± 20.0</td>
</tr>
<tr>
<td>Emotion</td>
<td>61.5 ± 22.6</td>
<td>70.8 ± 23.0 0.15</td>
<td>63.1 ± 23.6</td>
</tr>
<tr>
<td>Physical</td>
<td>64.9 ± 23.2</td>
<td>69.3 ± 26.7 0.46</td>
<td>65.7 ± 23.8</td>
</tr>
<tr>
<td>Psychological</td>
<td>64.5 ± 17.4</td>
<td>74.0 ± 21.0 0.03</td>
<td>65.9 ± 17.7</td>
</tr>
<tr>
<td>Social</td>
<td>67.3 ± 21.1</td>
<td>79.7 ± 20.0 0.03</td>
<td>68.2 ± 20.1</td>
</tr>
<tr>
<td>Total</td>
<td>64.7 ± 17.8</td>
<td>71.8 ± 21.7 0.13</td>
<td>66.1 ± 18.6</td>
</tr>
</tbody>
</table>

Values are means ± SD. CHD: congenital heart disease.
<sup>a</sup> Other diseases: myocardiopathy, coronary aneurysm, venous or intracardiac thrombosis, pulmonary arterial hypertension, arrhythmia, extracardiac disease.

Discussion

This prospective cohort study assessed the QoL of a cohort of 111 children who participated with their families in a non-selective INR self-monitoring education programme for VKA treatment. QoL was good among children aged 1–4 years and was moderately impaired between 5 and 18 years of age. This non-selective education programme is dedicated to all children who require VKA therapy in our institution [17]. As opposed to previous studies, we did not select patients or families by their native language, type of VKA or adherence rate to INR testing [15].

Interestingly, the QoL in the youngest children was quite good. The “disability paradox” [24] might explain good QoL levels in adults with serious chronic diseases, as well as among teenagers or young adults with CHD [2,25]. A recent

Table 4  PedQL scores: comparison between baseline and final evaluations for self- and parent-reports.

<table>
<thead>
<tr>
<th>Dimension</th>
<th>Baseline&lt;sup&gt;a&lt;/sup&gt;</th>
<th>Final&lt;sup&gt;a&lt;/sup&gt;</th>
<th>P</th>
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<tbody>
<tr>
<td></td>
<td>n = 59</td>
<td>n = 59</td>
<td></td>
</tr>
<tr>
<td>School</td>
<td>70.34 ± 19.7</td>
<td>74.47 ± 17.6 0.09</td>
<td></td>
</tr>
<tr>
<td>Emotion</td>
<td>72.07 ± 18.7</td>
<td>72.20 ± 20.4 0.87</td>
<td></td>
</tr>
<tr>
<td>Physical</td>
<td>71.68 ± 19.0</td>
<td>74.15 ± 19.7 0.13</td>
<td></td>
</tr>
<tr>
<td>Psychosocial</td>
<td>71.84 ± 15.5</td>
<td>73.54 ± 16.0 0.36</td>
<td></td>
</tr>
<tr>
<td>Social</td>
<td>73.45 ± 19.8</td>
<td>74.07 ± 18.7 0.78</td>
<td></td>
</tr>
<tr>
<td>Total score</td>
<td>71.82 ± 15.4</td>
<td>73.79 ± 15.0 0.27</td>
<td></td>
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</table>

<table>
<thead>
<tr>
<th>Dimension</th>
<th>Mothers’ reports&lt;sup&gt;a&lt;/sup&gt;</th>
<th>Fathers’ reports&lt;sup&gt;a&lt;/sup&gt;</th>
<th>P</th>
</tr>
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<tbody>
<tr>
<td></td>
<td>n = 9</td>
<td>n = 9</td>
<td></td>
</tr>
<tr>
<td>School</td>
<td>66.20 ± 19.8</td>
<td>65.80 ± 20.3 0.83</td>
<td></td>
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<tr>
<td>Emotion</td>
<td>61.44 ± 23.4</td>
<td>64.70 ± 22.9 0.21</td>
<td></td>
</tr>
<tr>
<td>Physical</td>
<td>64.26 ± 23.6</td>
<td>64.98 ± 24.7 0.92</td>
<td></td>
</tr>
<tr>
<td>Social</td>
<td>66.66 ± 19.3</td>
<td>66.92 ± 17.6 0.40</td>
<td></td>
</tr>
<tr>
<td>Total score</td>
<td>65.02 ± 19.4</td>
<td>66.11 ± 18.5 0.48</td>
<td></td>
</tr>
</tbody>
</table>

<sup>a</sup> Completed at the same time and at first administration for each pair. Values are means ± SD.
A large-scale international study found that overall QoL in 4000 adults with CHD from 15 countries was generally good; indeed, Apers et al. suggested that variation in QoL was related to patient characteristics but not country-specific characteristics [26]. In our study, we can assume that the optimization of healthcare system might have directly benefited these young children and their parents in terms of QoL. That includes management of pain [27], reduction in duration of hospital stay with outpatient management even in critical conditions [28], availability of parent and child rooms at the hospital, full social security cover, childcare allowance, etc. The creation of a formalized VKA education programme is in line with this global care of young patients and their families. Similarly, recent studies suggested a better QoL for children entering a self-monitoring VKA programme [15,16].

QoL among children older than 5 years, as reported by the children themselves or by their parents, was significantly lower than that of controls. However, long-term VKA treatment, as opposed to temporary treatment, did not impact the children’s QoL in our study. Besides, QoL did not seem to be affected by the duration of the disease. Indeed, several mechanisms interfere with the severity and the duration of the chronic disease and therefore may act together or in opposition to determine QoL. For instance, in our cohort, the subgroup of children with CHD had lower QoL scores compared with children receiving VKA for another reason. These children were born with a congenital disease, underwent open-heart surgery, probably experienced some degree of physical limitation, and while growing up became progressively aware of their health condition. Therefore, added to long-term VKA therapy, the burden of the CHD certainly influenced their QoL. Because of this multifactorial effect on QoL status, we purposely chose generic QoL questionnaires in our VKA education programme, as opposed to studies using specific instruments dedicated to QoL in anticoagulation [29].

In our study, the QoL scores evaluated by the children and their fathers remained stable throughout the education programme whereas QoL scores evaluated by mothers increased during the programme, independently of any improvement in the cardiac condition. Mothers represent the largest sample of completed QoL questionnaires. In a recent study, we found that the actual physical impairment (peak oxygen uptake) of children with a CHD was better correlated with parent-reported QoL than with self-reported QoL [30]. Similarly, in the study from Jones et al., parents reported a significant improvement in PedsQL scores (mean increase: 11.1) of 35 children commencing home INR self-testing, whereas the children themselves did not [16]. In our study, no significant differences were observed between both parents when participating together in the same education session. The fathers who participated in the education sessions might have been particularly involved in their child’s health condition and were therefore more accurate (vs fathers who did not participate) in their QoL reports. Therefore, parents’ reports are essential in paediatric QoL studies and are usually required in paediatric clinical trials.

At the end of the programme, QoL as reported by mothers was significantly better for children under warfarin than for those under fluindione. Warfarin has the longest half-life of 36 to 42 hours and is usually recommended in paediatrics, supposedly for a better INR stabilization [31,32]. This QoL impairment in the fluindione group might reflect the impact of numerous INR adjustments and the burden of frequent blood monitoring on children.

Study limitations

Children with laboratory INR monitoring were not included, which represents a selection bias; indeed, all our patients participated in the INR self-monitoring programme and we no longer have such patients in our centre. Patients, families, specialist nurses and physicians request INR self-monitoring, therefore we could not randomize our VKA population into two groups (self-monitoring versus laboratory) for ethical reasons. We cannot draw any conclusion from the QoL impairment in the youngest group of our study (<1 year of age), as a small number of patients of children of that age participated in the education programme. QoL questionnaires and scorings were heterogeneous; as a result, some children switched from an instrument (Qualin) to another (PedsQL) between baseline and final sessions. Our control population was heterogeneous, was not contemporary for the infants [18], and was derived from a foreign culture for children aged 2 to 7 years [19]. We plan to achieve the constitution of a large contemporary control paediatric population for further QoL studies. Families participating in formalized education programs might represent a selected “privileged” population. This might explain why fathers’ and mothers’ QoL scores were so close. However, our education programme is non-selective, suggesting limited bias [17].

Conclusions

This prospective study provided original data about the QoL of a cohort of 111 children and their parents who were participating in a formalized INR self-monitoring education programme for VKA treatment. QoL scores were good among children <4 years of age, and were moderately impaired for children aged 5 to 18 years. Child-reported QoL was higher than the proxy values reported by their parents. Paternal and maternal proxy reports were similar, globally and for each dimension. QoL among the subgroup of children with CHD under VKA was the most impaired, and probably requires specific education programmes in addition to an INR self-monitoring programme. QoL scores increased during the education programme according to maternal reports only and independently of any improvement in the health condition.

Acknowledgments

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Disclosure of interest

The authors declare that they have no competing interest.

Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at http://dx.doi.org/10.1016/j.acvd.2017.05.013.

References
