Towards person-centred quality care for children with life-limiting and life-threatening illness: Self-reported symptoms, concerns and priority outcomes from a multi-country qualitative study

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Abstract

Background: Paediatric life-limiting and life-threatening conditions (life-limiting conditions) place significant strain on children, families and health systems. Given high service use among this population, it is essential that care addresses their main symptoms and concerns.

Aim: This study aimed to identify the symptoms, concerns and other outcomes that matter to children with life-limiting conditions and their families in sub-Saharan Africa.

Setting and participants: Cross-sectional qualitative study in Kenya, Namibia, South Africa and Uganda. Children/caregivers of children aged 0–17 years with life-limiting conditions were purposively sampled by age, sex and diagnosis. Children aged 7 and above self-reported; caregiver proxies reported for children below 7 and those aged 7 and above unable to self-report.

Results: A total of 120 interviews were conducted with children with life-limiting conditions (n = 61; age range: 7–17 years), and where self-report was not possible, caregivers (n = 59) of children (age range: 0–17 years) were included. Conditions included advanced HIV (22%), cancer (19%), heart disease (16%), endocrine, blood and immune disorders (13%), neurological conditions (12%), sickle cell anaemia (10%) and renal disease (8%). Outcomes identified included physical concerns – pain and symptom distress; psycho-social concerns – family and social relationships, ability to engage with age-appropriate activities (e.g. play, school attendance); existential concerns – worry about death, and loss of ambitions; health care quality – child- and adolescent-friendly services. Priority psycho-social concerns and health service factors varied by age.

Conclusion: This study bridges an important knowledge gap regarding symptoms, concerns and outcomes that matter to children living with life-limiting conditions and their families and informs service development and evaluation.

Keywords

Symptoms, concerns, outcomes, children, palliative care, paediatrics

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Introduction

The recent Lancet Commission highlighted the individual, family and health system burden of life-limiting and life-threatening conditions and the ‘abyss’ of pain and palliative care access in low- and middle-income countries (LMICs). In 2017, an estimated 21 million children globally needed palliative care, with over 5.3 million facing health-related suffering; 99.9% of these live in resource-limited settings. Despite the commitment to scale up access to paediatric palliative care as a universal human right, service development and access remain poor. While substantial evidence exists that palliative care reduces costs and improves outcomes in adults, the evidence for effectiveness in children is limited partly due to the lack of valid child-centred outcome measures. To generate evidence of improved treatment, outcomes and care satisfaction, it is essential to first determine what matters to children with life-limiting and life-threatening conditions and their caregivers.

To date, paediatric health care outcome measures lack palliative care relevant content validity. Specific challenges of outcome measurement in this population include difficulties in developing tools for children with life-limiting and life-threatening conditions, including variable cognitive development, impacts of advanced disease and communication impairment. Recent evidence on outcomes specific to this population found it largely drawn from studies of oncology patients in high-income countries using proxy (family and professional) views and expert opinion. More so, the evidence does not consider social and developmental outcomes for children with life-limiting and life-threatening conditions (LLCs). A recent systematic review to identify and compare definitions of indicators used to assess the impact of specialised paediatric palliative care, reported limited participation of children in the interviews because of advanced disease. Another review of instruments used to assess the impact of paediatric palliative care interventions equally noted the limited involvement of children in the reported data, and the paucity of data on psychometric properties used to assess the impact of paediatric palliative care interventions.

Consequently, an evidence-based, child- and family-centred conceptual framework of priority symptoms and concerns is needed to inform service development, adaptation and development of outcome measures, as well as ensuring child participation in care and treatment decisions to align care to their priorities. With evidence on what matters to children with life-limiting and life-threatening conditions, we can move towards the development of appropriate paediatric palliative care services, as well as developing or adapting person-centred paediatric palliative care outcome measures.

Using developmental and age appropriate research methods, children can and should be engaged in research that shapes their care.

What is already known about the topic?
- A high number of children with life-limiting and life-threatening conditions need palliative care.
- Many of these children face health-related suffering with the majority living in resource-limited countries.
- Evidence of symptoms, concerns and other priority outcomes for children facing life-limiting and life-threatening conditions is predominantly from cancer populations, largely based on proxy reporting, and from resource-rich settings.

What this paper adds?
- This study provides evidence on symptoms, concerns and outcomes that matter to children with life-limiting and life-threatening conditions.
- Demonstrates thematic variations in symptoms, and concerns and by developmental age, that can shape support throughout an illness trajectory.
- Provides an evidence base for cross-cutting themes on symptoms, and concerns and across diverse life-limiting and life-threatening conditions.

Implications for practice, theory or policy
- With evidence on what matters to children with life-limiting and life-threatening conditions, we can move towards the development of appropriate paediatric palliative care services, as well as developing or adapting person-centred paediatric palliative care outcome measures.
- Using developmental and age appropriate research methods, children can and should be engaged in research that shapes their care.

Methods

Study design and methodology

This interpretive, qualitative study adheres to COnsolidated criteria for REporting Qualitative research (COREQ) reporting guidelines. E.N., M.A., L.L. and D.H. (senior researchers and palliative care professionals, female (n = 3), and male (n = 1)) conducted in-depth interviews with children or their proxy caregivers (when children were unable to self-report, because of advanced
of pre-verbal or non-verbal children (0–5 years) and car-
egivers of older non-verbal children (6–17 years old).

Knowledge.28

Ration and interpretation of gaps in theoretical and clinical
evidence from South Africa.29 Appraisal of our initial
data collection (2016) identified a failure to achieve a het-
erogeneous purposive sample, with under-recruitment of
languages and give informed consent.

Exclusion criteria
A child or caregiver too distressed to participate was
excluded.

Sampling
We purposively sampled participants by age, sex, diagno-
sis and place of care to maximise variation in the sample. These criteria were based on palliative care needs assess-
ment evidence from South Africa.29 Appraisal of our initial
data collection (2016) identified a failure to achieve a het-
erogeneous purposive sample, with under-recruitment of
non-malignant conditions, children below the age of 5
and those receiving care in an inpatient setting. Therefore,
the further recruitment was undertaken in 2017 to ensure the
purposive sampling criteria were fulfilled. Recruitment
ceased when thematic data saturation regarding the study
aim was achieved.30

Data collection tools
We developed semi-structured topic guides, based on a
review of the literature and consultation with experts. The
topic guide addressed the experience of illness in the con-
text of the child’s developmental stage and living situa-
tion.20 We developed two proxy topic guides for caregivers
of pre-verbal or non-verbal children (0–5 years) and car-
egivers of older non-verbal children (6–17 years old).

Three topic guides were developed for children who were
able to self-report aged 7–9, 10–14 and 15–17. The guides
covered the following aspects: symptoms and concerns in
living with life-limiting and life-threatening conditions and
how these affected patients’ well-being; overall illness
treatment experiences and its associated challenges. Concerns about how the breaking of ‘bad news’ (diagno-
sis) affected patients and families; the meaning of the ill-
ness to the patients; coping with concerns and problems;
and current priorities in life were discussed with children/
caregivers of children aged 7 years and above. The topic
guides were forward and backward translated into local
languages: Luganda, Lusoga and Swahili (Uganda); Luo
and Swahili (Kenya); Oshiwambo (Namibia); and Zulu,
Sesotho and Pedi (South Africa). Each guide was pilot
tested with two caregivers/children in three countries,
and minor revisions were made.

Recruitment and data collection
At each site, the study team screened and recruited chil-
dren and caregivers. Study participants were given an
opportunity to ask questions and seek clarification. For
each interview, we obtained informed consent from car-
egivers. Self-reporting children assented in addition to the
consent from their caregivers. After informed consent
and/or assent, interviews were conducted in private
rooms, children were interviewed without their caregiv-
ers. Each interviewer completed a 2-day training course
delivered by the principal investigator (E.N.) and two child
psychologists. Interviews began with a brief period of play
and storytelling (appropriate to developmental age) to
build rapport.31 Children were encouraged to talk freely,
share their views, experiences and issues that matter to
them.31 For interviews with younger children of up to
10 years, a play interlude of 15 min was given. A social
worker was available at each study site to support partici-
pants if they became distressed. All participants were
informed about this and about their freedom to choose
who should support them and that they could decline to
answer questions or stop the interview at any time. No
repeat interviews were conducted.

Ethics
Formal ethical approval was granted in all participating
countries: Kenya (refs: KEMRI/RES/7/3/1 and ERC/B/
VOL1374), Namibia (ref: 17/3/3/EN), South Africa (ref:
HPCA 03/10) and Uganda (ref: UNCS 2S2366) and King’s
College London, Research Ethics Office (LRS-15/16-3524).
Written informed consent was obtained from caregivers
for their children to participate in the study. In addition,
written assent was obtained from self-reporting children
aged 7–17 years following approval to participate in the
study from their caregivers. All children received an
age-appropriate gift worth US$7 given at the end to
acknowledge participation but was not mentioned in any study materials and refreshments during the interviews.

Data processing and analysis

The study team digitally audio-recorded interviews, alongside field note taken that were transcribed verbatim, pseudonymised and imported into QSR NVivo V12. Interviews conducted in local languages were translated into English and reviewed by local bilingual study partners for accuracy. We analysed data using the six linked steps of inductive thematic analysis (see Table 1) to identify themes symptoms, concerns and other outcomes. Our study is underpinned by the Bronfenbrenner’s bio-ecological systems theory, adapted for palliative care. It highlights the micro, meso, exo, macro and chrono systems of the socio-ecological layers. With increasing developmental age, the child’s level of interaction at socio-ecological layer level expands. The chronosystem is concerned with the concept of changing needs by developmental age, disease stage and function, as well as interconnectedness across layers (see Supplementary Material 1). We compared theme variation by developmental age: 0–2 (pre-verbal), 3–5 (more verbal), 6–9 (pre-adolescents), 10–14 (middle adolescents) and 15–17 (adolescents).

Rigour

To ensure data credibility, the research team provided the breadth of necessary experience: paediatric palliative care, nursing, child health and psychology, qualitative methods, psychometrics and medical anthropology. To enhance transferability, rich description of the research process, setting and participants is included. To ensure dependability and confirmability, dual-coding and careful study record-keeping were undertaken.

Results

Sample characteristics

We conducted 120 interviews with 61 children (95% of those approached) and 59 family caregivers (100% of those approached). Child participants ranged from 1 day to 17 years old; 52% were female, and 81% had a non-malignant condition (Table 2). Child interviews lasted a median of 37 min (range: 35–50 min), with older children participating in longer interviews. The report excludes the play interlude. Caregiver interviews lasted a median of 55 min (range: 50–60 min). The majority of proxy reporting caregivers were mothers (n = 38, 64.4%).
Table 2. Socio-demographic characteristics of participating children (N = 120).

<table>
<thead>
<tr>
<th>Variable</th>
<th>n</th>
<th>Child (n = 61)</th>
<th>Proxy (n = 59)</th>
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<td></td>
<td></td>
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<td>13</td>
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<tr>
<td>Age group 3–5</td>
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<td>13</td>
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<td>19</td>
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<tr>
<td>Age group 13–17</td>
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<tr>
<td>Urban</td>
<td>17</td>
<td>07</td>
<td>10</td>
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<tr>
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<td>29</td>
</tr>
<tr>
<td>Male</td>
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<td>07</td>
<td>07</td>
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<tr>
<td><strong>Diagnosis</strong></td>
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<td>HIV</td>
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<td>18</td>
<td>09</td>
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<td>09</td>
<td>03</td>
<td>06</td>
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<tr>
<td>Heart disease</td>
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<td>10</td>
<td>09</td>
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<tr>
<td>Sickle cell</td>
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<td>08</td>
<td>04</td>
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<td>Other endocrine, blood and immune disorders</td>
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<td>07</td>
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<td>Congenital and acquired neurological abnormalities</td>
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<td>00</td>
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<td>14</td>
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<td>Phase of losing normality</td>
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<td>17</td>
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<td>Declining</td>
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<td>17</td>
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<tr>
<td>End of life</td>
<td>16</td>
<td>00</td>
<td>16</td>
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<td><strong>Place of care</strong></td>
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<tr>
<td>Outpatient</td>
<td>56</td>
<td>31</td>
<td>25</td>
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<td><strong>Functional performance as measured by ECOG score</strong></td>
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<tr>
<td>0</td>
<td>25</td>
<td>16</td>
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<td>4</td>
<td>08</td>
<td>00</td>
<td>08</td>
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<td><strong>Caregiver’s relationship with child</strong></td>
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<td>39</td>
<td>38</td>
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<tr>
<td>Grand mother</td>
<td>23</td>
<td>12</td>
<td>11</td>
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<tr>
<td>Father</td>
<td>07</td>
<td>02</td>
<td>05</td>
</tr>
<tr>
<td>Grand father</td>
<td>04</td>
<td>02</td>
<td>02</td>
</tr>
<tr>
<td>Sibling</td>
<td>04</td>
<td>02</td>
<td>02</td>
</tr>
<tr>
<td>Auntie</td>
<td>04</td>
<td>04</td>
<td>00</td>
</tr>
<tr>
<td>Hospice caregiver</td>
<td>01</td>
<td>00</td>
<td>01</td>
</tr>
</tbody>
</table>

N/A: not applicable; HIV: human immunodeficiency virus; ECOG: Eastern Cooperative Oncology Group.

*a*14 missing value data not captured.

*b*14 missing value ECOG not assessed for children aged 0–2.
Symptoms and concerns

Four main thematic concerns emerged from the data: physical concerns, existential/spiritual concerns, psychological issues and concerns regarding health care quality (Figure 1). These are described below with illustrative quotes (see Supplementary Material 2 online for additional quotes to support the themes) alongside the following attributes: type of respondent, age of child and country setting code.

**Physical concerns.** Commonly described symptoms included pain; fatigue; difficulty walking or moving; skin problems (rash, change in skin texture); gastro-intestinal problems (nausea, vomiting, diarrhoea); respiratory problems (breathlessness, cough, fast breathing) (quotes 1–2); and central nervous system symptoms (seizures and convulsions):

When my body swells, it causes me severe pain, my legs are heavy, and I cannot move, and my stomach is also too heavy. Somehow, I cannot sleep well, and my wellbeing is negatively affected. I just want the water to be removed and I get some relief. Also, the skin darkens, I get funny patches on the skin and the outer layer is very tight. (Child 44; age-group: 10–14 years; K.E.; quote 1)

The child starts with swollen feet, vomiting, tired, cannot breathe well, and stomach pain. (Caregiver 01; child’s age-group: 6–9 years; N.A.; quote 2)

These symptoms, particularly pain, were associated with distress. Children reported experiencing an excruciating pain not easily explained, using terms like ‘throb-bing’, ‘constant’ ‘painful insect bite’ (quote 3). Children as young as 10 expressed the desire to die to escape the pain (quote 3), while those with sickle cell disease described a life of constant uncertainty and fear of the unpredictable pain episodes (quote 4). Symptom distress was also expressed in relation to the impact on well-being, function and activities of daily living:

When the pain starts. I just feel like dying, I just admire death, I yearn for an injection to kill me off because the pain is so severe. I just yearn for death to free me from pain, it is not even equitable [comparable] to the most painful insect bite, its gnawing you just want to die and be free from the pain. (Child 41; age-group: 15–17 years; U.G.; quote 3)

I hate it because the pain comes abruptly, it can find you in the middle of town and you collapse. sometimes the pain starts from the back. It is throbbing and very severe. Sometimes it is the joints that pain or ankles, at this point you lose the capacity to walk or stand, you can crawl may be, but you need support. Sometimes the chest gets so painful other times it is the arms, It’s all about pain. (Child 29; child’s age-group: 10–14 years; K.E.; quote 4)

Caregivers frequently described using the presence of concerns, such as not feeding (especially in infants and non-verbal children), as a sign for an underlying serious problem (quote 5):

The jaws are terribly weak and so he has not been sucking and as a result he has malnutrition. (Caregiver 18; child’s age-group: 0–2 years; U.G.; quote 5)

Children expressed the importance of being able to maintain physical function in activities, such as self-care and play (quote 6). Caregivers expressed similar concerns, with dependence on others often regarded as the most difficult challenge for their children (quote 7):

The swelling of the body, especially the eyes and body, it comes with severe pain, when I stop the medication, the pain crops back so I must be on medication all the time and I just hate. I cannot lift a jerry can, I cannot wash or even run. (Child 48; age-group: 10–14 years; U.G.; quote 6)

The hardest thing about my child being sick is his dependency to other people, this is so because of mental retardation. (Caregiver 50; child’s age-group: 15–17 years; U.G.; quote 7)

**Existential/spiritual concerns.** Existential/spiritual concerns were evident across all age groups. Children were worried about death, transitioning to an afterlife and letting go of their loved ones. The children were also worried about losing their future dreams and missing out on schooling, while other expressed hopelessness and a reluctance to continue with their education, take their medication or think of their future life (quote 8):

You are concerned, you will lose your education. Sometimes you lose morale for education and feel like drop out. When I think of starting up a business, I also lose morale because I am constantly thinking about the death. So, life gets crippled you cannot move forward. (Child 10; age-group: 10–14 years; U.G.; quote 8)

Caregivers equally voiced concerns of lost dreams, regarding issues that matter to their children (quote 9):

He is not aspiring to something big in life because he is mentally challenged. (Caregiver 59; child’s age-group: 15–17 years; N.A.; quote 9)

Children reported being concerned by unanswered existential questions, including ‘Why me?’, questioning why it had to be them to suffer adversity when their siblings were healthy. All this compounded the challenges children faced around unanswered questions (quote 10):
<table>
<thead>
<tr>
<th>Physical concerns</th>
<th>Reported concerns</th>
<th>Existential / Spiritual</th>
<th>Psycho-social</th>
<th>Psycho-social</th>
<th>Health care quality</th>
</tr>
</thead>
<tbody>
<tr>
<td>Physical symptoms</td>
<td>Physical concerns</td>
<td>Gastro intestinal</td>
<td>Changes in the way food tastes</td>
<td>Diarrhea</td>
<td>Lack of appetite</td>
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<tr>
<td>Emotional</td>
<td>Emotional needs</td>
<td>Positive affect</td>
<td>Happiness</td>
<td>Feeling of contentment</td>
<td>Negative affect</td>
</tr>
<tr>
<td>Outlook on death</td>
<td>Emotional needs</td>
<td>Worry about death</td>
<td>Do not want die</td>
<td>Existential loss</td>
<td>Loss of future goals and ambitions</td>
</tr>
</tbody>
</table>

Figure 1. A framework of self-reported symptoms, concerns and outcomes for children living with life-limiting and life-threatening illnesses.
I always wonder why God brought this painful disease to me. At my church, we are told God loves children but now imagine what I am going through. I always ask myself why me, and I have never found the answer, so I have given up on asking. I just keep it to myself. (Child 42; age-group: 6–9 years; K.E.; quote 10)

Children who felt connected to a supernatural being expressed a more positive attitude towards suffering, accepting their illness as God’s will safe in the knowledge that God would support them through their illness (quote 11):

What is most important to me is God, my mother and my father. God is very important in my life because we pray to him so that we can live in this world, he created us. He also has all the authority to decide if one lives or dies. (Child 3; age-group: 10–14 years; N.A.; quote 11)

In some instances, caregivers noted that children expressed a belief that they had undergone spiritual growth because of their suffering. Their life was perceived as a gift, and their suffering was embraced as part of their life (quote 12):

She just wants to cure completely and resume living a normal life. She asks me when she will get well and go for boxing. She also tells me ‘Now I appreciate that walking is a big gift from God, I can never take it for granted again’. (Caregiver 13; child’s age-group: 6–9 years; U.G.; quote 12)

According to caregivers’ reports, children as young as 5 years were concerned about dying with unfulfilled dreams (quotes 13). More so, uncertain prognosis was associated with meaninglessness for children:

She [the child] is worried about dying young. She has not even completed her education, so she is worried. She hates going to the theatre she always tells me that there are two possible outcomes, get better or die in the process but she always notes that God is in control. (Caregiver; child’s age-group: 3–5 years; U.G.; quote 13)

Some caregivers associated severe crying of neonates with suffering due to family rejection (children born with life-limiting and life-threatening conditions are sometimes rejected by their families and are not given names, as a result they lack family tree belonging (quote 14):

Then he [the child] also very annoyed because he [the child] does not have a name yet, when the father heard about this problem, he switched off the phone he thinks this is a curse to the family. So, my son is crying, he wants to belong to family and this not falling into place, so I have a very serious problem. (Caregiver 12; child’s age-group: 0–2 years; K.E.; quote 14)

Caregivers of pre-verbal children recognised calmness as an important indicator of the spiritual well-being of their children (quotes 15):

The situation in which he is now shows that he is at peace he no-longer cries at all. (Caregiver 33; child’s age-group: 3–5 years; U.G.; quote 15)

**Psycho-social concerns.** Children described in depth the psycho-social aspects of the illness experience. They described negative emotional responses, including sadness, anger, mood swings and suicidal ideation (quote 16):

I have had to accept my situation. I am angry. I feel like it’s a punishment and I need to get over it. I keep asking Why me? I was also depressed and cried a lot. I tried to commit suicide; I thought there was no point as my CD4 was low. (Child 06; age-group: 15–17 years; S.A.; quote 16)

In addition, caregivers and children described the importance of fun, happiness and living in the moment. They also wished to engage in activities that gave them fun and meaning (quote 17–18):

She also tells me to take her back to school, she keeps referring to other children who are attending school and asks me to take her back to school. She is bed bound and is so sad because she cannot play. I thank God that she is able to laugh and ask for what she wants. (Caregiver 23; child’s age group: 6–9 years; K.E.; quote 17)

This year I was going in P5 but due this sickness i wasn’t able to go to school I am now here. So, I miss all my friends at school, and I may never see some of them again. I also lose out on being in school, I want to be a mechanic. I am sick but I am happy because I play with my colleagues here, we don’t fight. When I play well with my colleagues, when the health workers play with us, it makes me very happy. (Child 20; age group: 10–14 years; U.G.; quote 18)

There were also treatment-/medicine-related concerns such as pill burden, taste of medication and medication fatigue, all of which contributed to poor treatment adherence (quotes 19):

I really want the experts to come up with a cure, I have lived with this disease for long, but I am still dreaming about the cure (laughs)). If those doctors can reduce on the number of tablets I take in a day, it’s too much you cannot live like this forever. (Child 28; age-group: 10–14 years; U.G.; quote 19)

The caregivers also described concerns related to treatment poor treatment adherence, which was associated with worry about child survival. Other concerns included behaviour that put children in conflict with family members (quote 20):
I am worried that [name] is not taking medications, I am afraid she will die. She is also cheeky and causing family problems and conflict. (Caregiver 18; child’s age group: 15–17 years; S.A.; quote 20)

Caregivers also highlighted behavioural concerns exhibited by the children such as negative emotions, particularly suicidal ideation, and loss of interest in previously enjoyed activities (quote 21):

Anger, she does not understand why she has HIV. She is also depressed and tried committing suicide this year, by hanging herself. (Caregiver 53; child’s age-group: 15–17 years; S.A.; quote 21)

Caregivers also noted that cognitive or developmental concerns related to the onset of the life-limiting and life-threatening conditions were cited (quote 22):

I think her reasoning is declining, she [is] not as intelligent as she used to be, she cannot add simple number. For me I am very worried, could be doctors are not telling us the truth about what happened to her. She is very quiet I do not know why; no single sound is coming out. (Caregiver 21; child’s age-group: 3–5 years; K.E.; quote 22)

For adolescents, much of the focus was on body image and wasting (quote 23):

My biggest concern is the fact my body swells, and I just look so bad. It affects my image, my legs and face the most beautiful parts of my body so why they swell and lose the beauty I feel so bad. (Child 25; child’s age-group: 15–17 years; U.G.; quote 23)

Children and caregivers attached high importance to normalcy, being able to achieve age-appropriate developmental milestones, including such simple outcomes as crying or feeding, and being treated like unaffected peers (quotes 24–25):

He cannot kick his legs around like normal children, he cannot even move his body at all, he breastfeeds well and passes urine like a normal child but there are those problems. (Caregiver 12; child’s age-group: 0–2 years; K.E.; quote 24)

I want to have life, I want to live up to the age of 25, get married and have a child, just like other children. That is the most important thing in my life. (Child 21; age-group: 15–17 years; U.G.; quote 25)

School-going children were concerned about missing school, and the negative impact of interruptions impacted on the quality of social interactions, relationships and life goals (quote 26). Children wanted time to interact with peers, feel understood, listened to, respected and appreciated:

This year I was meant to join primary 5, but due [to] this sickness I couldn’t go to school I am now here. So, I miss all my friends at school, and I may never see some of them again. I also lose out on being in school. When I play well with my colleagues, when the health workers play with us, it makes me very happy. (Child 39; child’s age-group: 6–9 years; K.E.; quote 26)

More so, according to caregivers of the non-verbal children, spending less time with significant others impacted on the quality of social relationships (quote 27):

He recognises me we are enjoying our small world. As soon as I sit on the bed he stops crying. He is calm and quiet. (Caregiver 06; child’s age-group: 0–2 years; U.G.; quote 27)

Regarding family relationships, children highlighted the importance of having a positive family environment as part of their social support structure. They valued love, physical presence and quality time with their significant others (quote 28):

Despite all this, what I like about it all is that I am very special to my mother am very close to her all so that is a good thing on my side. (Child 2; child’s age-group: 6–9 years; K.E.; quote 28)

Genetic illnesses such as sickle cell commonly strained family relations and at times led to separations. This happened on two levels; children were treated like outcasts as male partners posited that their family trees could not be associated with such illnesses. Some children also questioned their lineage as these illnesses seemed to affect them alone and their siblings were healthy (quote 29):

The men always blame the women. Yet this is a genetic problem which could be on the man’s side. So, my mother was thrown out of the home and we are struggling. Men believe all the bad things are brought in by the women. This should stop. I want to know why I have this disease and why is it me of all the children. It is only me with this disease, we are five children and I am the last born. Why is it me only, why isn’t it at least two or three people in the family? I want an answer. (Child 16; child’s age-group: 15–17 years; N.A.; quote 29)

Children also worried about protecting their families from any bad experiences. For example, one child explained why she would withhold news about her worrying symptoms because she was concerned that it would cause distress to her family (quote 30):

When I fail to breathe, I just keep quiet. I do not want to worry other people; some people are already ill so if you share such bad news they can collapse into coma and die. So, for me I just rely on God he decides if you must die or survive. (Child 34; age group: 10–14 years; K.E.; quote 30)
Caregivers of pre-verbal children reported cases of psycho-social distress linked to lack of a ‘name’ which connected the children to a family tree and gave them a sense of belonging (quotes 31):

Then he also very annoyed because he does not have a name yet, when the father heard about this problem, he switched off the phone he thinks this is a curse to the family. So, my son is crying, he wants to belong to family and this not falling into place, so I have a very serious problem. (Caregiver 12; child’s age-group: 0–2 years; K.E.; quote 31)

Some children reported being given names associated with the suffering of living with life-limiting and life-threatening conditions; for some of them, this was a source of value and strength (quote 32):

When am ill I always ask my mother, why is it me who got this disease and she tells me she does not know? It is just that. So, I think it was God’s decision, God decided that of all people in my home it should be me to live this way. That is why I was given this name [name] just to say it is God who decided. I am like this because it is God’s choice. In fact, the full name is [name translated as [name] what God decides]. I had another name [name] but I prefer to use the latter, am what I am because it was God’s choice. In fact, I was given this name when my parents first learnt of my diagnosis. (Child 57; age group: 15–17 years; quote 32)

Caregivers were concerned about their children having to deal with stigma and bullying due to looking different from others, such as when facing severe weight loss, or peers knowing about their diagnosis (quote 33):

My brothers’ physical appearance, is he going to stay small forever, will he ever grow. He is bullied and teased at school, because he is small for his age. (Caregiver 31; child’s age-group: 15–17 years; S.A.; quote 33)

Children also noted negative stereotypes about children living with life-limiting and life-threatening conditions. This negatively impacted on their self-esteem and personal identity. Negative body image and fears of stigma, discrimination and rejection also impacted on adolescents wishing to initiate sexual relationships (quote 34):

There are several people who laugh at [me], I got a woman she had a child and died. So, my family members started saying you see, a ‘sickle’ should not engage in sex, they should not have children or even have relationships. Even now I have a girlfriend, but I am very cautious, am hesitant to go and have sex with her, I still live with my father. I am very worried I could die during sex, so I just try to avoid it. It is shameful if you die during this act. (Child 9; child’s age-group: 15–17 years; K.E.; quote 34)

Finally, caregivers described the catastrophic financial expenditures associated with care and struggles to balance the need to work and earn money, and to care for terminally ill children and their siblings. Some adolescents had taken on the onus of meeting their own transport-related costs and increasingly found it difficult to meet such demands (quotes 35–36). Participants also described the need for practical support to meet the physical need of food:

Financial problems. I live far away from the hospital and living off grants. Transport is expensive. I need more financial support. (Caregiver 04; child’s age-group: 6–9 years; N.A.; quote 35)

As I told you earlier on that at home, we are not financially okay so sometimes I would fail to get money for transport to take me to [name of health facility] and get my treatment so I would miss. But even if I explained to them, they could not listen much as I told them the truth. They took me to the counsellor and the counsellor told me that they were going to write me off their program. (Child 26; 17 years; U.G.; quote 36)

Caregivers of children with mobility problems mentioned need for support with wheelchairs to ease the mobility of their children (quote 38):

He is not able to turn himself, if you leave him in a certain position, you will find him in that position no matter how far you went so the condition affect his body movement, he always need someone to turn him, bath and feed. I was sent to a hospital-based self-help project to buy a wheel chair using my own money. (Caregiver 03; child’s age-group: 10–14 years; N.A.; quote 38)

Health care quality. Children felt it critical to have a positive and friendly working relationship with health service providers due to their need for chronic care until death. Those who reported friendly and supportive staff attitudes felt happy to remain in care and adhere to clinical treatment; those reporting negative staff interactions wanted to disengage (quote 39):

That day they abused her [mother to the child] and she came back home upset, so when I went back for the next visit, I asked them why they abused my mother since that time they became ruder to me. And they told me that for us we have never abused your mother, however the health workers try to empower parents show more responsibility to their children. But I felt so bad, when they abused my mother. Even tomorrow is my treatment day and I must go there but I have fears of what will happen to me, I don’t know what they are going to tell me this time. (Child 44; child’s age-group: 15–17 years; K.E.; quote 39)

Caregivers of hospitalised verbal children described the difficulties of sharing wards with adult patients (participants reported this situation as a problem in Uganda’s
Even now I have a girlfriend, but I am very cautious, am hesitant to go and have sex with her, I still live with my father. I am very worried I could die during sex, so I just try to avoid it. It is shameful if you die during this act. No, there is no privacy around here, so I do not have an opportunity. There are younger children, so you cannot ask such questions. (Child 09 age group: 15–17 years - UG [quote 45])

They intimidate us, you are a young girl and the doctor is much older and superior laughter, they also show us that they are very busy. So somehow there is no opportunity for us to ask any questions. We have a television here I wish they could bring us some educative tapes and we learn on our own. (Child 36; child’s age-group: 15–17 years; U.G.; quote 44)

Theme variation by developmental age. We also conducted a stratified analysis by developmental age, and the findings by theme variation are presented in Table 3. The themes on pain and other symptoms, physical function, indicators of spiritual well-being, connection to creator, positive and negative emotions, quality of social relationships, stigma and discrimination were overarching across all developmental age groups. Normalcy, outlook on self, spiritual growth, existential loss and worry, severe pain, were mentioned by children or caregivers of children (aged 6 years and above). Severe pain triggering search for escape were dominant themes for self-reporting older children (7 years and above).

Discussion

This study describes the domains of health-related suffering faced by children living with life-limiting and life-threatening conditions and their families. Emergent themes included symptoms and concerns in children living with life-limiting and life-threatening conditions and their families in the domains of physical, existential/spiritual, psycho-social concerns and health care quality. While this supports the WHO statement that difficulties affecting adults with life-limiting and life-threatening conditions are relevant for children,37 this study describes the additional and unique concerns related to activities of daily living (such as play, life goals and values, going to school, ‘name’ happiness, biological growth and development that matter to children and caregivers).

Children described the psycho-social impact of life-limiting and life-threatening conditions specifically in relation to their ability to engage in their usual activities, such as playing, having fun, attending school and achieving their dreams when they become adults.40 They described concerns about normalcy and maintaining their ‘child identity’, and the shame, anger and stigma of not being able to do what their peers can do, which in turn led to discrimination and reluctance to access services. The theme of ‘name’ was a unique finding. In the African culture, personal names may have meanings, affect one’s personality and their ability to thrive in society.41 A such, the lack of a ‘name’ is associated with lack of cultural identity, a critical ingredient for
Table 3. Display of selected data by domain and subtheme based showing theme variation by developmental age.

<table>
<thead>
<tr>
<th>Domain</th>
<th>Subtheme</th>
<th>Key issue, by age group</th>
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<tbody>
<tr>
<td></td>
<td></td>
<td>0–2 years: pre-verbal</td>
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<tr>
<td></td>
<td></td>
<td>3–5 years: more verbal</td>
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<tr>
<td></td>
<td></td>
<td>6–9 years: mid-childhood</td>
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<td></td>
<td></td>
<td>10–14 years: pre-adolescent</td>
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<tr>
<td></td>
<td></td>
<td>15–17 years: adolescent</td>
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<tr>
<td>Physical concerns</td>
<td>Pain</td>
<td>Disease- or treatment-related pain</td>
</tr>
<tr>
<td></td>
<td>Other symptoms</td>
<td>Feeding/crying indicates well-being to carers</td>
</tr>
<tr>
<td></td>
<td>Symptom distress</td>
<td>Severe symptom = impending death (caregiver)</td>
</tr>
<tr>
<td></td>
<td>Physical function</td>
<td>Want child to be ‘normal’</td>
</tr>
<tr>
<td></td>
<td>Physical needs</td>
<td>Physical presence of caregivers</td>
</tr>
<tr>
<td></td>
<td>Spiritual, existential concerns</td>
<td>Calmness associated with being in good spirits</td>
</tr>
<tr>
<td></td>
<td>Beliefs, practices, faith</td>
<td>Show religiosity (e.g. prayer) and belief in a supernatural being (e.g. ‘God’)</td>
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<tr>
<td></td>
<td>Outlook on death</td>
<td>Existential worry and death concerns</td>
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<tr>
<td></td>
<td>Existential loss</td>
<td>Concern over loss</td>
</tr>
<tr>
<td></td>
<td>Spiritual resources</td>
<td>Participate in spiritual rituals (e.g. prayer)</td>
</tr>
<tr>
<td></td>
<td>Connection to Creator and others</td>
<td>Present personal concerns to Creator</td>
</tr>
<tr>
<td></td>
<td>Spiritual growth</td>
<td>Search for meaning of adversity, purpose of fragile life</td>
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<tr>
<td></td>
<td></td>
<td>Trust and hope in significant others (e.g. caregivers if &lt; 10 years)</td>
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<tr>
<td></td>
<td></td>
<td>Need for love and self-worth (from interacting with caregivers)</td>
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<td></td>
<td></td>
<td>Appreciate life as a special gift; want suffering to ‘pay’ for family’s good health</td>
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<tr>
<th>Domain</th>
<th>Subtheme</th>
<th>Key issue, by age group</th>
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</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>0–2 years: pre-verbal</td>
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<tr>
<td>Psycho-social concerns</td>
<td>Emotional</td>
<td>Positive emotions</td>
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<tr>
<td></td>
<td></td>
<td>Smiling, etc. seen as signs of psychological well-being (in &lt;10 s)</td>
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<tr>
<td></td>
<td>Behavioural</td>
<td>Adherence to treatment therapy issues</td>
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<tr>
<td></td>
<td></td>
<td>Antisocial behaviour more common</td>
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<tr>
<td></td>
<td>Cognitive</td>
<td>Children and caregivers concerned about effect of illness on learning</td>
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<tr>
<td></td>
<td></td>
<td>Concern by carers about effect of illness on speech/communication development</td>
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<tr>
<td></td>
<td>View of self</td>
<td>Want quality interaction with significant others</td>
</tr>
<tr>
<td></td>
<td>Quality of relationships</td>
<td>Physical presence of carer is critical</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Want play with peers, socialise with significant others</td>
</tr>
<tr>
<td></td>
<td>Initiate and maintain</td>
<td>Keen to attend school for social reasons</td>
</tr>
<tr>
<td></td>
<td>relationships</td>
<td>Goal is to become responsible adult</td>
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<td></td>
<td>Stigma and discrimination</td>
<td>Want to initiate and maintain relationships</td>
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<tr>
<td></td>
<td></td>
<td>Some express sexual and reproductive health concerns</td>
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<td></td>
<td>Perspective of others</td>
<td>Sexual/reproductive health concerns dominant</td>
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<tr>
<td></td>
<td>Life values</td>
<td></td>
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<tr>
<td></td>
<td></td>
<td>Isolation from peers to avoid stigma, discrimination</td>
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<tr>
<td></td>
<td></td>
<td>Caregivers protect child from peers by isolation</td>
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<tr>
<td></td>
<td></td>
<td>Child isolates self to avoid stigma and discrimination</td>
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<tr>
<td></td>
<td></td>
<td>Protect their loved ones from suffering/distress associated with their situation</td>
</tr>
<tr>
<td></td>
<td>Name</td>
<td>Have equal opportunities like ‘normal children’</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Achieve life goals</td>
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<tr>
<td></td>
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<td>Need for a name for family tree and community belonging</td>
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<tr>
<th>Domain</th>
<th>Subtheme</th>
<th>Key issue, by age group</th>
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<tr>
<td></td>
<td></td>
<td>0–2 years: pre-verbal</td>
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<tr>
<td>Health care quality</td>
<td>Staff attitude</td>
<td>Existential longing to be home or in appropriate environment</td>
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<td></td>
<td>Environment of care</td>
<td>Location of care affects well-being (e.g. if share ward with adult patients)</td>
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<tr>
<td></td>
<td>Information, communication</td>
<td>Want child/adolescent-friendly services</td>
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<td></td>
<td></td>
<td>Concern over care transitions</td>
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<tr>
<td></td>
<td></td>
<td>Need information about disease, treatment, positive living</td>
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<tr>
<td></td>
<td></td>
<td>Want place to ask private questions</td>
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<td></td>
<td></td>
<td>Involvement in decision-making</td>
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<td></td>
<td>Disclosure</td>
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Cross cutting themes are bolded. No variations in themes were noted.
ability to thrive and hence worth addressing in paediatric palliative care.

In addition, the children also described distress related to future unfulfilled dreams and concerns about death and the afterlife. The prominence of the psycho-social and spiritual concerns described by even very young children was surprising but highlights the need to address these concerns to improve child well-being, self-efficacy, resilience, coping and retention in care.18,42–44

The experience of symptoms also differed by age for children and their caregivers. For many children, pain was associated with fear, but for young children pain was associated with fear of impending death, while for adolescents it mediated desires for death to escape distress and suffering.45 For some children, anticipated treatment and procedure-related pain impacted most heavily on their well-being.46 These findings illustrate the multidimensionality of pain and its impact on the whole person (i.e. ‘total pain’).47

Children and caregivers expressed the importance of child-friendly services, drawing a link between issues such as emotional distress associated with exposure to high mortality rates in inpatient wards, unapproachable staff and their negative impact on their well-being.48,49 These findings suggest that to improve care, health care quality and the additional existential/spiritual and psycho-social concerns should be considered.

Finally, children may not always be able to self-report, so proxy reporting is important to consider. There was considerable overlap in the concerns expressed by children and caregivers; however, while caregivers could report on visible and functional concerns, self-report is required for emotional and abstract constructs.50

Strengths and limitations

This study contributes to the evidence gap related to the problems and concerns of children with life-limiting and life-threatening conditions19 through a large data set, based on a comprehensive purposive sampling frame and includes self-report by children (and caregivers where that was not possible); across diverse age and diagnostic groups; and in resource-limited settings, increasing the transferability of the findings. We also undertook stratified analysis by developmental age and generated important evidence on differences in concerns. Our study has limitations: due to recruitment difficulties and variation in access to palliative care services,4 the number of participants from each country varied, with the greatest proportion of participants from Uganda (43%) and the least from South Africa (12%). Moreover, we interviewed children about complex constructs, which they might find challenging to elaborate upon comprehensively given their limited vocabulary. Resultantly, some details could have been missed due to inadequacies in language. However, consistent with the existing developed country–based evidence, we found multi-dimensional concerns which have been posited to be important in patients living with life-limiting and life-threatening conditions, reinforcing the validity of our findings.19,51

What this study adds

This study provides evidence on symptoms and concerns that matter to children with life-limiting and life-threatening conditions and their families within the core domains of physical, spiritual/existential, psycho-social and quality of care. We, therefore, recommend a multi-professional approach to care with attention to quality of services, as well as the multi-dimensional child and family concerns.

Our study also demonstrates the feasibility of self-report in children and the use of age-appropriate or colloquial language, which should be a key consideration for research involving children. These novel data address gaps in the literature by collecting primary self-report data in low- and middle-income countries.

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Author contribution

E.N., K.B., F.E.M.M., J.D., R.A.P., M.A. and R.H. contributed to study concept and design, analysis and interpretation of the data, drafting of the manuscript and critical revision of the manuscript for important intellectual content. I.J.H., M.A., L.L., E.B.K.L. and F.N.M.-P. contributed to analysis and interpretation of the data, drafting of the manuscript and critical revision of the manuscript for important intellectual content. Z.A., D.H. and S.G. contributed to study supervision, drafting of manuscript, administrative, technical and material support. All authors approved the final manuscript as submitted and agree to be accountable for all aspects of the work.
Availability of data

The transcripts are not publicly available for confidentiality reasons, but we share anonymised verbatim quotes to support the themes generated as primary data.

Declaration of conflicting interests

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Supplemental material

Supplemental material for this article is available online.

References