Dynamics on the field: A focused ethnographic study of pediatric pain management at four Ghanaian hospitals

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Abstract

Background

As part of efforts to develop and implement a short course educational program on pediatric pain management, the current study sought to understand the culture and contextual factors that influence children’s pain management in order to improve the practice in pediatric care settings.

Methods

Guided by Bourdieu’s theory of practice, a focused ethnographic study was conducted from October, 2018 to February, 2019. The study was contextualized at four Ghanaian hospitals among purposefully sampled nurses, physicians, hospitalized children and their families. During the 20-week study period, three ethnographers spent 144 hours conducting participant-observation sessions. Formal and informal interviews were held with participants in addition to review of hospital records.

Results

Analysis of the field data resulted in four themes. “Children’s pain expression and response of caregivers” described the disposition (habitus) of both children and caregivers to act in particular ways due to children’s incomplete health status (bodily capital) which caused them pain and also resulted in discomforting procedures. “Pharmacological pain management practices and attitudes” elucidated the use of analgesics as the mainstay disposition (habitus) in children’s pain management due to high level of respect (symbolic capital) given to such interventions on the pediatric units (field). “Managing pain without drugs” illustrated healthcare providers and family caregivers’ disposition (habitus) of using diverse nonpharmacological methods in managing children’s pain. “Communication and interaction between pain actors” depicted how children’s access to care givers (social capital) can serve as a powerful tool in influencing pediatric pain assessment and
management disposition (*habitus*) on the pediatric units (*field*).

**Conclusions**

The *habitus* of pediatric pain actors toward pain assessment and management practices are influenced by various forms of capital (*social, cultural, symbolic, bodily and economic*) operating at different levels on the pediatric care *field*. Quality improvement programs that seek to enhance pediatric pain management should use the insights obtained in this study to guide the development, implementation and evaluation stages.

**Background**

Even though the knowledge and methodologies to relieve pain exist [1, 2], children still experience needless pain during hospitalization [3, 4]. The effects of unrelieved pain among children cannot be overemphasized. Inadequately treated pain can lead to both short-term and long-term biopsychosocial consequences on affected children. These include: activation of the stress response [5], impaired functional ability [6], delayed recovery, prolonged hospitalization, increased cost of healthcare [7], absence from school [8], post-traumatic stress [9], isolation [10], chronic pain and negative impact on children’s quality of life [11]. Unrelieved children’s pain also affects the social, psychological and financial aspects of the family and society [12].

Considering the negative consequences of unrelieved pain, it is not surprising that suboptimal pain management has been considered an international health tragedy [13] and the freedom from pain regarded as a basic human right [14]. Furthermore, the International Association for the Study of Pain [15] declared the right of all people to have access to nondiscriminatory pain management, the right to pain acknowledgment and information on its assessment and management, as well as the right to have access to appropriately trained healthcare professionals. In order to prevent the numerous associated effects of undertreated pain, the American Academy of Pediatrics also
advocates for the expansion of pediatric pain assessment and management knowledge among healthcare providers [16].

As part of efforts to improve pediatric pain management, several guidelines have been developed to guide the assessment and management of children’s pain [17–19]. In spite of these best practice guidelines, pain management largely remains under-prioritized in children [20, 21]. The under-prioritization of children’s pain management may be the result of failure to translate best practice guidelines into practice, demonstrating the importance of context in implementation science [22]. Earlier studies have demonstrated the important value of cultural context in shaping evidence-based practices by caregivers in the assessment and management of pain in pediatric care settings [23, 24].

Culture refers to the behaviour, attitudes, values, systems of meaning and skills shared by a group of people [25]. It guides beliefs and attitudes pertaining to meaning of illness, healthcare seeking behaviours, degree of receptivity to healthcare interventions, and healthcare practices [26, 27]. The sociocultural context is critical in enhancing our understanding of pediatric pain management, especially in low-middle income countries where culture has been identified as a barrier to optimal children’s pain management [28, 29]. Although efforts have been channeled toward addressing the knowledge, attitudes, self-reported practices, impact of pain care [30], experiences and perceptions of stakeholders involved in children’ pain management [31]; there is limited evidence on the context and cultural factors that underpin the assessment and management of pediatric pain in clinical practice.

Earlier studies have enhanced our understanding on the influence of context in shaping the complexity involved in pain management in clinical settings [24, 32]. It appears from the reviewed literature that, the culture and context of pediatric pain management has not been explored from Sub-Saharan African perspective. The effectiveness of healthcare
practice is also highly dependent on the resources available to the practitioner as well as
the environment and culture that forms the framework for interaction with the patient
[33]. As part of efforts to develop and implement a short course educational program on
pediatric pain management, the current study sought to understand the culture and
contextual factors that influence children’s pain management in order to improve the
practice in these settings.

Methods

Study design

Guided by Bourdieu’s theory of practice [34], a focused ethnographic study was conducted
over the course of five months from October, 2018 to February, 2019. This approach was
chosen as the researchers intended to understand the processes involved in the
assessment and management of children’s pain within the pediatric care settings of four
Ghanaian hospitals through the use of three foundational concepts (field, capital and
habitus). According to this theory, field refers to the social space and structures within
which individuals practice. Capital signifies resources or power over a field and the
individuals operating within it; this power may take the form of social (peers, networks),
cultural (education, socio-demographic), economic (salary, finances), bodily (health
status) or symbolic (reputation, respect, status). Habitus represents dispositions or
inclinations which cause individuals to behave in particular ways over time and is
demonstrated through an ongoing and emerging relationships between the individual
(agency) and the collective (structure); it is underpinned by personal experiences,
backgrounds, professions and circumstances.

The overall goal of this focused ethnographic study was to gain an understanding of both
the emic and etic perspectives. Emic perspective describes how members or insiders of a
particular group perceive and understand their world whereas the etic perspective
characterizes an outsiders’ understanding of an observed culture [35]. The present study formed part of a larger study that sought to explore the educational needs on pain management in children.

Setting

The study was contextualized at four hospitals in the Ashanti region of Ghana. For the purposes of ensuring anonymity and confidentiality, the four hospitals have been labelled as A, B, C and D. These hospitals were purposively chosen as they all had specially designed in-patient children’s care settings and were located in diverse geographical locations (urban, peri-urban, rural). Hospital A was a specialist private children’s hospital with a bed capacity of 20. Hospital B was also a specialist children’s hospital owned by the government of Ghana; it had 26 beds in the facility. Hospital C was a quasi-governmental hospital with a 22-bed capacity pediatric unit. Hospital D was a mission-based hospital with 19 beds in the pediatric unit. All four hospitals admit children under 13 years and oversee to those with medical or minor surgical conditions.

The pediatric care settings were colourfully painted and had drawings of flowers, fruits, cartoons, rainbows, balloons among others. The units were divided into various sections to cater for children based on their age or disease condition. There were specially designated nurses’ station where newly admitted children and their families were received and examined. The units also had an emergency or resuscitation area where they triaged and cared for children in need of such services. The in-patient beds had a side locker for keeping patients’ belongings and a bedside chair for the parent or guardian. As part of the hospital protocols, hospitalized children are housed with at least one parent or guardian at all times in the pediatric units. However, the hospital had limited accommodation facilities for these parents or guardians. All the pediatric care settings had notice boards on which the ward protocols and duty rosters were displayed.
During the period of the study, the weekly admission rates within the children’s unit ranged from six to 15. The pediatric care settings are staffed by 10-20 nurses and three to six physicians who work on a shift basis. The number of nurses per shift ranges from two to six and that of physicians varies from one to two. Nursing work within the children’s unit was daily operationalized on a three-tier shift system: morning, afternoon or night duties. This system ensured the provision of a 24-hour continuous nursing care to hospitalized children and their families. On a daily basis, physicians visited the wards individually or as a team to review the conditions of the admitted children. Physicians are also consulted outside their working hours as and when deemed necessary.

**Participants**

Participants for the current study comprised of nurses, physicians, hospitalized children and their families. Nurses and physicians were purposefully sampled if they were working in the pediatric units of the included hospitals. Hospitalized children were selected if they had pain complains as one of the symptoms of their present medical condition or were undergoing an invasive or skin-breaking procedure. Families of such children were also purposively sampled to participate in the study.

**Data collection procedures**

Data collection for the study began following administrative and ethical approvals from the respective hospitals and ethics committee. The researchers approached the nurse managers and the nurse-in-charges of the children’s units and briefed them about the purpose and procedures involved in the study. The nurse in-charges then introduced the researchers to eligible nurses, physicians, hospitalized children and their families. The researchers then briefed the eligible participants on the scope of the research and gave them the opportunity to ask questions. Answers and clarifications were given to participants before the study began and during the entire research period. Reflexivity was
ensured by keeping a detailed journal during data collection and analysis, where personal reflections and perceptions were written so that both the emic and ethic experiences did not bias the participants’ accounts. Prior to each fieldwork, the ethnographers (AKA, JKD & CKA) documented their mood and expectations before entering the unit to serve as a check when identifying the research themes at a latter period. Field data for the current study comprised of observations, interviews (formal and informal) and review of documents (audit).

**Observations**

Observational data collected within the first three sessions (morning, afternoon, evening) in the children’s unit of each hospital were not counted as the ethnographers (AKA, JKD & CKA) wanted some familiarization with the participants in order to reduce the Hawthorne’s effect [36]. At the beginning of each day’s fieldwork, the researchers took part in friendly conversations and other non-pain-related activities with the nurses, physicians, children and their families. The researchers actively kept a moderate level of participation by balancing participation with observation [37]. This approach enhanced co-operation and a positive working relationship with the participants throughout the data collection period. Two of the ethnographers were female nurses and the third was a male nurse, all of whom did not work in any of the included hospitals. They entered the children’s units, wearing their nursing uniforms and blended in as natural participants [38]. The ethnographers participated in daily nursing activities as shadow nurses whilst keenly observing and documenting activities related to children’s pain assessment and management. The ethnographers were present at each of the hospitals for two to three days per week and, on average for four hours per each observation. During the 20-week study period, the ethnographers spent 144 hours conducting participant-observation sessions with 36 hours spent in each hospital.
Observations were guided by a checklist to keep the researchers’ focus on children’s pain assessment and management. Specifically, observations were focused on the ward environment, number of participants present during each observation (nurses, physicians, number of children on admission, families of hospitalized children), pain assessment, pharmacological and nonpharmacological pain management activities, and documentation of pain assessment and management interventions, as well as the feelings of all pain actors present in the unit. In addition, data on the availability of pain tool, its location and interactions among the participants (nurses, physicians, children and families) were also observed.

**Formal interviews**

Twenty-eight (28) nurses and 12 physicians working in the children’s unit of the included hospitals were purposively chosen to participate in the formal interviews. Consideration was given to age, gender, working years in the health profession and in the children’s unit. Formal interviews were held each with hospitalized children who were above five years (20) and their corresponding family member who was resident with them in the hospital (20). Apart from the child’s ability to communicate effectively, the selection process considered the gender, age, medical condition or procedure performed. Efforts were also made to include diverse categories of family care-givers such as mothers, fathers, grandmothers, aunties, uncles among others. The children, family caregivers, nurses and physicians from the four different hospitals were purposefully selected to achieve maximum variation to enhance our understanding on the socio-cultural context of pediatric pain management in Ghana.

Individual or group interviews were conducted on scheduled dates with the participants using English or Asante Twi (a popular indigenous dialect spoken in Ghana) language. The interviews were recorded with participants’ permission and lasted from 10 to 40 minutes.
per each session. With the aid of a semi-structured guide, four authors (AKA, JKD, EO and CKA) facilitated the interview sessions at private, quiet rooms within the hospital premises. At least two out of the four authors were present during each interview to allow for notes taking and the smooth running of the sessions. Notwithstanding, the first author (AKA) was present at all interviews either as a facilitator or notes taker during the sessions. All interviews were completed in one hospital before moving to the next. The nurses, physicians and family care givers responded to questions which included: their views on the prevalence of children’s pain, methods they used in assessing and managing children’s pain, their communication and preparedness towards their role in children’s pain assessment and management. The hospitalized children were also asked similarly about the extent of pain they experience as a result of their medical conditions or procedures performed on them. They were also asked about the role the other participants (nurses, physicians, family care givers) played in the assessment and management of their pain and their wishes regarding pain care. Recorded interviews were transcribed verbatim by two of the researchers (EO and CKA) after which they were analyzed by all the researchers involved in the study before the next data collection session. The interviews that were conducted using the Asante Twi language were translated into English and back translated into the original language with the assistance of a language translator.

**Informal interviews**

Informal interviews were casual conversations, in which the ethnographers clarified observations with participants. These were noted in shorthand by the researchers in their field notes and later transcribed in full for analysis. Forty (40) nurses, 12 physicians, and 72 children-family dyads were informally interviewed over the study period.

**Document audits**
Over the course of five months, a total of 108 patient folders were reviewed for pain assessment and management reports. Patient folders of those children who had pain complains during hospitalization and had been on admission in the unit for at least 24 hours were selected. Thirty-six (36) nursing reports and physician notes on hospitalized children’s conditions were thoroughly examined over the same time period. Documents displayed on the notice boards of the wards and the pediatric care settings were also reviewed.

**Data analysis**

Using Bourdieu’s theory of practice [34] as the theoretical lens, an iterative process of data collection and inductive analysis followed the data analysis method developed by Leininger [38] (refer to Figure 1). Recorded field notes, transcribed interviews and document review reports were transported into NVivo 12 Plus software for data management. Identification of themes was carried out by reading through field notes, transcripts and document review reports of each study site multiple times. Coding was conducted by the ethnographers and verified with the assistance of two experienced qualitative researchers. Concepts and themes were continuously clarified, revised and updated during data collection and analysis period. Documented personal reflections were consulted in the data analysis process to ensure that the researchers’ experiences did not bias participants’ accounts. Data saturation was achieved within and across study sites as no new information or themes emerged from the field data. Information gathered during the fieldwork were merged during thematic analysis to create a holistic sense of the culture and context of pain assessment and management in the children’s units of hospitals in Ghana.

**Trustworthiness**

The trustworthiness of this focused ethnographic study was ensured by adhering to Guba
and Lincoln's [39] principles of credibility, conformability, dependability and transferability. Credibility and conformability were achieved through triangulation and member-checking. Triangulation was enhanced using multiple study settings (four hospitals with different ownership and geographical locations), data sources (observations, interviews, document reviews), data collectors and analysts. Member-checking was operationalized through the sharing of data interpretations and conclusions with participants for clarifications, corrections and additional information as deemed necessary.

In making sure that the information gathered was consistent and dependable, two researchers (AKA & JKD) conducted a pilot study with the study instrument (checklist) before it was used in the four selected hospitals. Comparison of data collection and thematic analysis was done to ensure consistency of the identified themes. Detailed field notes of the fieldwork also facilitated auditability of the study’s findings. Although qualitative research does not aim at generalization, the settings and procedures involved in the study have been extensively described to enhance transferability of the findings into similar settings.

Results

Participant characteristics

The characteristics of the participants have been presented in Table 1.

Themes

Children’s pain expression and response of caregivers

One of the recurring themes of the current study was “children’s pain expression and response of caregivers”. Pain was a common complaint among hospitalized children who could verbally communicate. Different levels of pain (mild, moderate and severe) were exhibited by the children at the time of their admission and intermittently during
hospitalization. The pain experienced was caused by their medical conditions (such as fractures, sickle cell disease) and or skin breaking procedures (such as intramuscular injections, intravenous cannulation and medication, lumbar puncture, surgical operation, wound dressing among others). Children’s incomplete bodily capital thus served as a source of discomfort and subsequently subjected them to such discomforting procedures. As a result of the pain experienced during skin breaking procedures, the children cried loudly and clung unto their family caregivers for relief. Family caregivers thus, served as a form of social capital for the children in distressing times. Nevertheless, there was no escape from pain as healthcare providers regarded the procedures to be of important value and went ahead with them even if it meant applying physical restraints to these children. The high level of respect (symbolic capital to these procedures influenced healthcare providers’ habitus of restraining these helpless children. Some family caregivers also felt helpless and even cried as they witnessed their children go through the pain associated with skin breaking procedures and some clinical conditions. In some cases, family members could not withstand the traumatic experience their children were going through and excused themselves during such procedures, leaving the children solely in the care of the healthcare providers. A mother of a 6-month old baby expressed it this way “Madam I cannot watch, it makes me sad to watch my child in pain like that”. The children were also observed to exhibit the non-cooperative behaviours that were demonstrated during the skin breaking procedures in subsequent non-painful procedures such as checking of their body temperature, auscultation of their heart and breath sounds among others. According to the children, they kept memories of these painful events which sometimes prevented them from co-operating with healthcare providers during subsequent procedures. The recollection of their previous personal experiences thus influences their noncooperative habitus.
Healthcare providers admitted to difficulties in assessing pain among children who could not verbally communicate as a result of their developmental stage or medical condition. According to the healthcare providers, this could be attributed to their lack of *cultural capital* in this area of pain care. They believed crying was the language of infants and toddlers and did not regard this as a cause for concern among these vulnerable population. Their *habitus* of not doing much in such situations on the *field* resulted from the low *symbolic capital* attributed to children’s cry. They further intimated their lack of access to *cultural capital* as they experienced difficulties in distinguishing crying due to pain from other causes. On the other hand, some healthcare providers believed children were more likely to be in pain if the assessment findings of the four traditional vital signs (temperature, pulse, respiration and blood pressure) were higher than the normal range of values. Healthcare providers heavily relied on family caregivers as a source of *social capital* in determining pain and its causes among the children. Whilst some family caregivers reported of using changes in their children’s activity, behaviour and verbalizations to detect pain, others admitted to being clueless in obtaining a definite sign or symptom of their children’s pain. A father of an 18-month child had this to say, “Hmmm Madam he’s been moaning and turning aaarrrrhh... but I can’t tell if he’s feeling any pain though he cries intermittently too”. Hence, *social capital* had a dualistic role in the *field*: serving as a catalyst for pediatric pain assessment on one hand or not favourable on the other hand. Family caregivers and healthcare providers believed children’s expression of the extent of their pain was dependent on the child’s age which served as a source of *cultural capital* and their lack of access to *bodily capital* due to their medical conditions. A nurse puts it in this way:

“Oh the pain, it usually depends on the condition of the child, what the child will bring in... so maybe if a child has trauma, maybe a child is involved in an accident, compared to a
child who has a mild fever or malaria, the pain would be different. So it depends on the
condition that, the child would bring.”

Healthcare providers’ individual habitus of not prioritizing pain was influenced by the
collective habitus operating in the field as pain was not actively assessed compared with
the other four traditional vital signs (temperature, pulse, respiration rate and blood
pressure). Additionally, there were no specially designed documentation sheets for pain,
nor was pain assessment pages or sections integrated into the patient’s folders, unlike the
four traditional vital signs. This could be attributed to the symbolic capital given to this
aspect of pain care and the institution’s lack of access to economic capital in providing
these resources. Thus, pain documentations if any only occurred in the physicians’ and
nurses’ notes. An example of such report was:

“Patient complained of severe headache, 375mg of intravenous Paracetamol administered.
Vital signs checked and recorded. Patient tepid sponged to reduce the recorded
temperature of 39.6°C to 37.8°C after 40 minutes. Child is made comfortable in bed and
mother reassured”

Observational data from the field work also revealed the presence of two pain assessment
tools (Faces Pain Scale and FLACC-Faces, Leg, Activity, Crying, Consolability Scale) in only
one of the four hospitals. The few pain assessment tools were also not used by the
healthcare providers in assessing children’s pain. The hospitals’ lack of access to
economic capital in purchasing these assessment tools influenced healthcare providers’
habitus of not systematically assessing children’s pain and its subsequent lack of
communication during ward rounds and handing over activities. Some healthcare
providers were not familiar with pain assessment tools due to their own deficiency of
cultural capital and the institutions’ lack of economic capital; both of which underpinned
their habitus of assessing children’s pain in a suboptimal manner on the field. Some of the
healthcare providers also attributed their *habitus* of not using standardized tools in guiding their pain assessment of children to heavy workload and shortage of staff in the demanding pediatric care *field* due to deficiencies in both *cultural capital* and the institutions’ deficiency in accessing *economic capital*.

**Pharmacological pain management practices and attitudes**

Another recurring theme from the coded field data was “pharmacological pain management practices and attitudes”. Pharmacological interventions served as the mainstay *habitus* in children’s pain management due to high *symbolic capital* given to such interventions. The common pharmacological agents used in managing children with mild to moderate pain were Paracetamol, Diclofenac and Ibuprofen (Non-Steroidal Anti-Inflammatory Drugs – NSAIDs) which were administered via enteral, parenteral and anal routes. These drugs were alternated in some circumstances due to their mechanism of action and to reduce their side effects. According to the healthcare providers, children who were admitted on account of burns were given Ibuprofen syrup or Paracetamol suppository thirty minutes before wound dressing to reduce pain. This *habitus* of healthcare providers on the *field* was influenced by their desire to reduce children’s pain and discomfort during such procedures.

Physicians’ *habitus* of seldomly prescribing opioids for children in pain even when it was the best medication in a particular situation was influenced by different forms of *capital* operating on the *field*. Some of these include: the *symbolic capital* associated with these class of analgesics, limited access to *cultural capital* on opioids, influence of other healthcare providers who served as a source of *social capital*, the institutions’ lack of *economic capital* in providing these medications and some families inability to afford these relatively expensive pain medications due to their deficiency in *economic capital*.

Nurses’ *habitus* of infrequent administration of prescribed opioids for children were also
underpinned by the above-mentioned forms of capital on the pediatric care field. Thus, opioid medications (Morphine, Pethidine and Tramadol) which were to be administered when necessary ended up not being given at all even if the child’s condition demanded it. According to the nurses, they deliberately gave other analgesics even when physicians had prescribed opioids. Some nurses further confessed that they have never administered an opioid in their practice even when it was prescribed.

“I know we are to give Pethidine or morphine, but I have never administered some for any child. After surgery, they come with Pethidine to be given where necessary, but they even do well with ibuprofen and Paracetamol administered intermittently so most times we don’t give. You know, it comes with its own issues.” (Nurse 2, hospital A)

The habitus of not prescribing and providing opioids (such as Morphine) to children and their families to be taken at home was underpinned by healthcare providers’ prejudiced symbolic capital associated with these analgesics and their insufficient cultural capital on these drugs. In spite of misconceptions associated with opioids, many healthcare workers admitted that they have not witnessed children experiencing side effects of opioids in their practice during the interview sessions. A physician expressed it this way “I am not sure I have encountered any side effects from the opioids”. A review through the hospital records (doctor’s notes, nurses’ notes and report books) also confirmed this as there were no documented evidence of pain medication side effects.

**Managing pain without drugs**

Another recurring theme from the field data was “managing pain without drugs” which largely interconnects with the previous theme of “pharmacological pain management practices and attitudes”. Some of the pain experienced by the children were managed without drugs on the field. Both healthcare providers and family caregivers served as a source of social capital for the children and were observed to be working together to
manage the pain of the children in this regard. Healthcare providers and family caregivers were observed to be engaged in the *habitus* of using nonpharmacological methods such as cuddling, stroking, coaxing, consoling, positioning and breastfeeding in distracting children's attention from pain during skin breaking procedures or upon self-report of pain from the children. A typical example of this was the observation of a mother who was softly singing and talking to her baby to stop crying after a venipuncture. Some healthcare providers were also observed to be consoling, cuddling and stroking the back of babies who were crying during and after skin-breaking procedures. A classic instance is when a nurse explained to a seven-year old after a wound dressing to stop crying as he will be leaving the hospital soon. She stated: *“Oww don’t cry my darling... I will take you home today”*

The pediatric care field of the hospitals were equipped with television sets, toys, play areas, and colourful wall paintings with child-friendly designs (such as rainbow, cartoons, etc); reflecting the institutions’ ability to access *economic capital* in the provision of these resources. According to healthcare providers, these facilities were meant to distract children’s attention from painful circumstances during hospitalization. Healthcare providers also said that the environmental layout of the ward as a non-pharmacological pain management strategy had the greatest impact on newly admitted children as they tend to engage with them in the earlier stages of admission. They however, felt its effect diminished over time as children stayed in the ward for longer periods.

Conversely, the children and family caregivers expressed happiness and contentment with the availability of these therapeutic nonpharmacological facilities which diverted children’s attention during painful procedures and served as a form of pain relief. Some children were observed playing on wooden and plastic animal toys in the play area of the hospitals. Other children were also handed teddy bears and toys by healthcare providers
before skin breaking procedures to distract their minds and subsequently manage the pain associated with such procedures. Notwithstanding these, we observed shortfalls in these resources based on the number of admitted children on the field which posed as a challenge and influenced children’s habitus of using these distraction methods in managing pain. The children further appealed for the provision of additional toys on the field due to the insufficient numbers and the bad state of some playing materials; indicating the hospital’s deficiency in accessing economic capital in meeting this demand.

**Communication and interaction between pain actors**

The final recurring theme was “communication and interaction between pain actors”. Observation of the interactions that occurred among the participating physicians, nurses, children and their families revealed a cordial and respectful engagement with each other. Physicians and nurses had separate ward rounds but there were periods where both groups came together and reviewed the hospitalized children and their families. The physicians and nurses engaged the children and their families by addressing them by their names and asking how they were doing including any pain they were experiencing. Family caregivers whose children were not capable of communicating effectively gave reports on their well-being and asked for clarifications on issues which were unclear to them. Most of the time, family caregivers and nurses informed physicians about children’s pain complaints after which pain medications were prescribed for the children. The nurses also informed their colleagues to reaffirm their pain assessment findings prior to informing the physicians. All of these interactions reflect how children’s access to social capital can serve as a powerful tool in influencing pediatric pain assessment and management habitus.

Informal conversations with some hospitalized children and their family caregivers unearthed that they were satisfied and happy with how the healthcare providers engaged
with them concerning their wellbeing including that of pain. Nevertheless, some children and their family caregivers were unhappy about their short duration of communication with healthcare providers and wished to be engaged more than what they received. In addition, the dissatisfied family caregivers felt that healthcare providers did not inform them about the treatment being prescribed and administered for their children. Resultantly, some healthcare providers were perceived as good whilst others were perceived by the children and their family caregivers as unkind. This further supports the dualistic potential of social capital in either facilitating or impeding pediatric pain care communication. An interaction with a mother of a six-year old boy who was unhappy about communication with healthcare providers expressed it this way:

“Hmmmm, since I came around at 4am, no nurse has personally come to me to interact with me... Madam it’s their work and since other patients equally need their care, I don’t want to be seen as disturbing them, but I wish one of them could come and talk with me.”

On the other hand, healthcare providers (physicians and nurses) were generally satisfied with the relationship that existed between them and the hospitalized children as well as their families.

The children clung unto their family caregivers in the hospital environment and perceived them as advocates, who made decisions in their best interest. In situations where children were asked questions by the healthcare providers, they turned toward their family caregivers even before giving responses. Thus, family caregivers mainly served as a source of social capital and the mouthpiece in identifying pain and its causes among children who could not talk. On the other hand, some family caregivers felt reluctant in constantly reporting their children’s pain for fear of being tagged as “troublesome” or “medical attention seekers”. They therefore preferred to wait for routine ward rounds before relaying any changes in their children’s condition. Again, this reaffirms the
dualistic role of social capital in working for or against improved pediatric pain care. Family caregivers also assisted with the activities of daily living of the children including bathing, grooming, feeding and playing. They were constantly beside the children and offered support in care giving whenever it was required; reflecting the power of social capital on the field.

Discussion

The multi-faceted complexities and dynamic environments within which healthcare systems operate imply that the same interventions are not likely to work in the same manner in different settings [41, 42]. Thus, understanding the culture and context of pain care is critical for the successful development and implementation of a sustainable short-term educational program targeted for nurses on pediatric pain management. The current study, guided by Bourdieu’s theory of practice [34] was grounded in the assumption that the assessment and management of children’s pain is largely influenced by different forms of capital and habitus operating in a particular field. Our findings revealed that the habitus of the pediatric pain actors toward pain assessment and management practices were influenced by various forms of capital (social, cultural, symbolic, bodily and economic) operating at different levels on the pediatric care field.

Children’s incomplete health status (bodily capital) caused them pain, resulting from discomforting skin breaking procedures and this stimulated the habitus of both children and caregivers to act in particular ways. In this study, pain was a common complaint among hospitalized children as reported in previous studies [43, 44]. Consistent with earlier studies [45, 46], procedural pain was described by the participating children as the most distressing part of the hospitalization process and memories of such events influenced their habitus by discouraging them from co-operating in subsequent procedures. This situation also resulted in a distressing and discomforting habitus among
some family caregivers as reported in a previous study [47].

Although pain is regarded as of high *symbolic capital* and considered as the fifth vital sign by international bodies [48, 49] , we found that pain did not receive the same level of attention for assessment as the traditional four vital signs (temperature, respiration, pulse and blood pressure) in these Ghanaian hospitals. Healthcare providers admitted to not being able to assess pain in non-verbal children due to their lack of *cultural capital* in this regard and heavily relied on the *social capital* provided by family caregivers who were sometimes clueless about how to assess pain in their children. This also reflects the dualistic potential of social capital in either stimulating or impeding optimal pediatric pain care [50].

In spite of the exponential growth in pain assessment tools for diverse categories of children [51, 52], few pain assessment tools were available at the pediatric care *field* reflecting the institution’s lack of *economic capital* in providing these resources. The few tools were also not utilized in practice to assess children’s pain owing to healthcare providers’ deficiencies in *cultural capital*, adding to the numerous published reports of infrequent pain assessments among hospitalized children [53, 54]. Additionally, there existed limited documentation of pain assessments and evaluations in the studied hospitals, which further underscored the low *symbolic capital* given to pain in the Ghanaian setting. While these pain assessment inadequacies are not unique to the included hospitals [55, 56], they are unacceptable and influences the prolonged unnecessary suffering of vulnerable children. The leadership and management of these hospitals and healthcare in general should take advantage of their *symbolic* and *economic capital* to educate and motivate practitioners to prioritize pediatric pain assessments and provide them with the needed pain assessment tools and documentation charts in practice.
The use of NSAIDs served as the mainstay habitus in children’s pain management due to high symbolic capital given to such interventions on the field. The use of these drugs have been established as a safe treatment option for children with mild to moderate pain [1, 57]. Healthcare providers’ habitus of seldomly prescribing and administering opioid analgesics was also attributed to various forms of capital (symbolic, cultural, social and economic) controlling the pediatric care field. Misconceptions on side effects of opioid analgesics (such as drug dependence, respiratory depression and addiction) have been reported to be prevalent among health care providers in earlier studies [58, 59]. However, these fears are usually unfounded and some of the perceived side effects can be alleviated or totally eliminated by using multi-modal pain treatment approaches [60, 61]. Positive changes in the habitus of healthcare providers on opioid use must be actively pursued through regular, short-duration educational interventions. They could be educated on ways of mitigating identified risks and side-effects, minimum doses to achieve effective analgesia, regular assessment for opioid side effects or for indications to discontinue treatment, and also provided with practical and published evidences to correct their misconceptions.

Pain actors’ habitus of using diverse nonpharmacological methods to manage pediatric pain were influenced by children’s access to social capital and the availability of nondrug resources which reflected the field’s access to economic capital. Notwithstanding these, healthcare providers reported of the time-consuming nature of some non-drug interventions which prevented them from using such methods. The shortfalls observed in the provision of playing resources on the field also posed as a challenge and influenced children’s habitus of using these distraction methods in managing pain. Additional nonpharmacological interventions were thus desired by the children despite their contentment with the institution’s economic capital in providing such resources on
the field. The use of wall designs, cartoons, toys and other playing materials in the hospital has been noted as a form of active distraction during painful procedures among children [62, 63]. The reported wide range of effective nondrug pain relief methods such as ball squeezing [64], oral glucose administration [65] and musical mobiles [66] should be explored and encouraged in practice as these methods are cheaper, simple, minimally invasive and can serve as useful adjuncts to analgesics [67].

Communication and interaction between the pain actors depicted how children’s access to social capital can serve as a powerful tool in influencing pediatric pain assessment and management habitus on the field [68]. The process of optimal pediatric pain management occurs in a context which is reliant on effective communication among key stakeholders where the pain care needs of the child take center stage. Observation of the relationship and interactions that occurred among the participating physicians, nurses, children and their families revealed a cordial and respectful engagement with each other, unlike Clancy’s report of frustration and expression of anger in their treatment of children’s pain among six healthcare providers in sub-Saharan Africa [56]. This notwithstanding, some family caregivers were dissatisfied with the quality of communication existing in the field, and therefore desired for more attention. This is unfortunate, as these family caregivers who serve as a source of social capital give voice to the pain of their children, provide information for assessment and support pain management modalities [50, 69]. As such, they should be prepared for their role in the assessment and management of children’s pain so that they can be of help in such circumstances. Healthcare providers should be trained to effectively communicate with these important members in order to improve pediatric pain assessment and management. The healthcare providers in this study also complained of heavy workload in the highly demanding pediatric care field which prevented them from pain assessment and management interventions which also
impacted their communication with some family caregivers. Heavy workloads impact on the ability of healthcare providers to carry out their responsibilities such as assessments, administering treatments, and communicating with clients and family caregivers about progress of treatment [70]. This calls for an improvement in the healthcare provider-patient ratio; additional auxiliary staff can also be added to the healthcare workforce to assist with the non-technical duties so that the professional staff can concentrate on the technical duties and have more time with children and their families.

Measures which were employed to ensure trustworthiness (credibility, conformability, dependability and transferability) in the current study is considered as one of the study’s strengths. The use of Bourdieu’s theory served as a useful lens for the examination of the sociocultural context of pediatric pain assessment and management at the four Ghanaian hospitals. In spite of the above stated strengths, we reckon that the presence of the ethnographers on the field might have influenced the behaviours of the pain actors even though measures such as not recording the earlier field work were done to safe guard against this. Though, the ethnographers were reflective during the generation of themes, we cannot preclude biases inherent in this process.

Conclusion

The *habitus* of pediatric pain actors toward pain assessment and management practices are influenced by various forms of capital (*social, cultural, symbolic, bodily and economic*) operating at different levels on the pediatric care field. This intricate process is heavily influenced in the hospitalized settings by the culture of “how things are done around here”. The current study has provided useful information on the contextual and cultural factors that influence the assessment and management of children’s pain in hospitalized settings. Quality improvement programs that seek to enhance this area of practice should use the insights obtained in this study to guide the development, implementation and
evaluation stages.

It is not necessarily the establishment of the gap which is interesting, but the exploration of why the gap between beliefs and knowledge about pain management, and the performance of it, actually exists. Effective clinical leadership is therefore required if we are keen on improving pain care outcomes for vulnerable children and their families as it influences resource provision and determines the standards for acceptable and unacceptable behaviour through role modelling and expectation setting. This study has further demonstrated the importance of context in influencing healthcare practices.

Abbreviations
NSAIDs – Non-steroidal anti-inflammatory drugs

Declarations

**Ethics approval and consent to participate**

Administrative approvals were given by the respective hospitals before obtaining ethical approval from the Committee on Human Research Publications and Ethics (CHRPE), School of Medical Sciences (SMS), Kwame Nkrumah University of Science and Technology (KNUST) (Reference number: CHRPE/AP/574/18). Hypothetical names were assigned to the hospitals and the participants as a method of guaranteeing anonymity and confidentiality of the research findings.

Consent from the participants (nurses, physicians, children and families) were continuously collected orally by asking permission to observe or interview and ensuring that the participant being observed or interviewed did not object to the data collection method used. None of the participants objected to be observed or interviewed in the study. The participants who were engaged in formal interviews further gave written informed consent as evidence for their voluntary participations. Participants were
informed about the voluntary nature of their participation and their right to withdraw at any stage of the study without receiving any sanctions. The research data were accessible to only the researchers involved in the current study.

**Consent for publication**

Not applicable.

**Availability of data and materials**

The dataset supporting the conclusions of this article would be provided upon reasonable request from the corresponding author.

**Competing interests**

The authors have no competing interests to declare.

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The study was self-funded by the authors.

**Authors’ contributions**

The study was conceptualized and designed by all authors (AKA, EO, VB, JKD, CKA & AA). AKA, EO, JKD and CKA collected the data; all authors participated in the data analysis (AKA, EO, VB, JKD, CKA & AA). Manuscript was written by AKA and EO, all authors critically reviewed, revised and approved the final draft for publication (AKA, EO, VB, JKD, CKA & AA).

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Table

Table 1. Study participants

| Group                    | Observations | Formal Interviews | Informal Interviews | Document |
|--------------------------|--------------|-------------------|---------------------|----------|
| Nurses                   | 40           | 28                | 40                  | 36       |
| Physicians               | 12           | 12                | 12                  | 36       |
| Hospitalized children    | 72           | 20                | 72                  | 108      |
| Family caregivers        | 72           | 20                | 72                  |          |

Figures

Figure 1

Four phases of Leininger’s data analysis
