Association of Paediatric Emergency Medicine

1683 PARENTAL PERCEPTIONS OF PAEDIATRIC EMERGENCY DEPARTMENTAL ATTENDANCE IN CHILDREN DURING THE COVID-19 PANDEMIC IN UK (PPECID). THE QUALITATIVE VIEW

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Background The COVID-19 pandemic has massively impacted on how healthcare is accessed. Paediatric Emergency Departments (PED) attendances have fallen dramatically in the UK during this period. Studies so far have focused on quantitative data when exploring this topic; whereas, our study uses qualitative methodology to review parental attendance motivations. Our study addresses a critical gap in understanding the factors that may influence caregivers’ PED attendance in the context of the COVID-19 pandemic within our region.

Objectives The aim of the study was to explore and understand how the COVID-19 pandemic influenced caregivers’ health-seeking behaviours. Specifically, we assessed how caregiver behaviour has changed, the perceived factors affecting presentation to the paediatric emergency department, how information around health has been received by caregivers and how the information should be communicated by health-care professionals to meet the caregivers’ needs.

Methods This qualitative sub-study was part of a mixed method study assessing parental perceptions and experience of PED. It used a purposive and convenience sampling method to recruit caregivers attending PED with varied age, gender and ethnicity. Using in-depth interviews, we explored caregivers’ experiences and perceptions of PED attendance among caregivers who presented with their children at our trust, comprising of two hospitals, during November - December 2020. Informed written consent was obtained from caregivers. Twenty semi-structured interviews were conducted. Interviews were audio recorded and transcribed verbatim. Data were analysed thematically using coding to identify emergent themes, patterns and concepts from participants’ accounts. The thematic content analysis was conducted by two study members for inter-rater reliability.

Results The findings coalesced around three main themes: caregivers’ lived experience of the pandemic, healthcare seeking perceptions and behaviour, and information sources/communication channels. When describing experiences of the pandemic, almost all participants expressed intense emotions such as fear, anxiety, depression and the need to adapt to new social norms. Decision-making around when and where to seek care for a sick child was predominantly influenced by the severity of the child’s condition, professional medical advice and perceptions of the SARS-COV-2 transmission risk posed by visiting the hospital: ‘we’ve heard … of people going into hospital for something unrelated and catching it’ ID8. Some caregivers reported how hospital staff actions and/or the hospital environment itself could mitigate and/or propagate these concerns in equal measures, such as wearing masks correctly. Caregivers also expressed worries around overburdening the NHS ‘I’ll just ring the GP see what they say … make sure they (the hospital) wouldn’t think we were wasting their time’ ID4. Despite the varied range of communication channels reportedly providing health information, which included government, health services, mainstream and/or social media, many participants cited distrust in one or several of these sources. Health professionals, schools and social networks were cited as trusted and preferred sources for disseminating information around paediatric health issues.

Conclusions Our findings show that caregivers are fearful of bringing sick children to PED and some mainstream sources of information are perceived to be untrustworthy. Public health messaging should emphasize the availability and safety of hospital facilities.

Quality Improvement and Patient Safety

1688 JUST IN CASE TRAINING

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Background The Trust introduced Just in Case (JIC) training in 2014 in the ITU areas under the global PediRES-Q research study. The initiative reinforces traditional annual resuscitation training, delivering refresher and preparatory training at the bedside to help staff become more focused and aware of essential skills and interventions that may be required for individual patients, meeting the learners needs when it arises, through collaborative learning with peers. It supports at the point where greater performance is required, promoting a confident and responsive workforce, providing a timely, child centric approach to the delivery of resuscitation skills at the bed-side where all clinical staff can be engaged and appreciate the end goal of identifying those at risk of deteriorating and prevention.

Objectives To improve the recognition and enable early intervention and management of the acutely unwell child in order to prevent deterioration into cardiorespiratory arrest and rapid response in paediatric resuscitation by providing Just in Case training to clinical staff.

Methods In response to the impact of the Covid 19 pandemic, there were opportunities to extend the JIC training, bringing additional expertise, support and reassurance to all clinical areas but especially where Covid 19 patients were identified, increased acuity of patients, staffing levels where stretched, the PEWS >9, ward teams, Clinical Site Practitioner’s (CSP) or parents had identified a high risk of deterioration or collapse. Also, to support to staff redeployed from the North Central London Paediatric Network, ward-based training regarding the Trust emergency response systems and familiarisation and use of Trust emergency equipment was delivered.

Success led to an extension of the initiative and collaboration with the CSP team and clinical staff, identifying JIC...
opportunities, including a refresher of the skills of effective bag-valve-mask ventilation, application of defibrillator pads and quality CPR. Reviewing emergency processes such as algorithms and protocols, highlighting situational awareness including bedscape preparation, role allocation and clinical decision-making is supported. Furthermore, expertise within the team encourages the staff to explore clinical conditions of patients, giving context to the disease process including support for modified approaches to resuscitation. Increased visibility in the clinical areas, has resulted in increased requests from staff for this training to develop their confidence, supporting the complex and progressive clinical needs of the child requiring a higher dependency.

Results

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<thead>
<tr>
<th></th>
<th>Jan 2019-Dec 2019</th>
<th>Jan 2020-Dec 2020</th>
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<tbody>
<tr>
<td>Total 222 calls attended by Clinical Emergency</td>
<td>147</td>
<td>125</td>
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<tr>
<td>Team</td>
<td></td>
<td></td>
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<tr>
<td>Cardiopulmonary arrests</td>
<td>13</td>
<td>10</td>
</tr>
<tr>
<td>Respiratory arrests</td>
<td>34</td>
<td>48</td>
</tr>
<tr>
<td>Unplanned admissions to Critical Care Units</td>
<td>158</td>
<td>168</td>
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<tr>
<td>Number of staff trained</td>
<td>-</td>
<td>384</td>
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</tbody>
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Conclusions Aside from the earlier escalation and interventions, resulting in a decrease in 2222 calls overall and cardiorespiratory arrests, the positive impact of this additional bedside teaching has been very well received and praised by the staff in clinical areas, especially those caring for complex, high risk patients at the point of care and in context of the specific disease process.

British Society of Paediatric Endocrinology and Diabetes

1692 CONGENITAL HYPOTHYROIDISM WITH GLAND IN SITU: EXAMINING RISK FACTORS, HEALTH, AND EDUCATION OUTCOMES USING LINKED ADMINISTRATIVE DATA

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Background Since the introduction of newborn screening 40 years ago, the birth prevalence of Congenital Hypothyroidism (CH) has doubled in the UK. This appears to be due to an increase in screen-detected congenital hypothyroidism with gland in situ (CH-GIS). It is not clear why CH-GIS prevalence has increased, nor are there recent studies examining the health and education outcomes for children with CH-GIS.

Objectives Study objectives are to 1) establish a research-ready clinical CH database of children who screened positive for CH through the North Thames Newborn Screening Programme and were referred to the Endocrinology Team at Great Ormond Street Hospital (GOSH) from 2006 onwards; 2) investigate the reasons for the increase in screen detection of CH-GIS; 3) examine health and education outcomes for children with CH-GIS compared to children who have other forms of CH and to those who do not have CH-GIS.

Methods To address objective 1, we used REDCap (Research Electronic Data Capture) to collate data on children attending GOSH who had screened positive for CH; data had been stored in a number of repositories/formats: EPIC, Excel, Word and GOSH legacy databases. To address objective 2, we will establish a cohort study of children born in the North Thames region and screened for CH through the North Thames Newborn Screening Programme between 2000 and 2020 (~2.2 million children) to investigate perinatal risk factors for CH. These children will be linked to ONS birth and death registration data and to the CH database. To address objective 3, we will use a case-control design. All children on the CH database between 2006 and 2020 will be linked to Hospital Episode Statistics (HES), the National Pupil Database (NPD), and national community dispensing data. Controls will be selected from a pseudonymised national birth cohort created from linked HES-NPD data. We started the process of applying for ethical and Confidentiality Advisory Group (CAG) approval and negotiating with data providers (NHS Digital, GOSH, Department for Education and Public Health England) in May 2020. Public involvement activities have also been undertaken with parents and children/young people, including the British Thyroid Foundation and GOSH Young Persons’ Advisory Group (YPAG).

Results The CH database contains 109 variables on 1928 children, with follow-up information on treatment and diagnoses until December 2020. We received ethical approval in November 2020, however negotiations with data providers and CAG are ongoing. As of early March 2021, it has taken 1 research associate 10 months working full-time to obtain these approvals and we expect it to take a further 6 months before the study is fully approved.

Conclusions We have demonstrated the utility of REDCap for developing a clinical database from which datasets for research can easily be extracted. The process of obtaining approvals for projects using linked administrative and health data in England continues to be complex and time consuming, leading to severe delays for data-focused research projects; this needs to be addressed in order to allow population-based research into rare diseases in children using linked clinical and administrative datasets.

Paediatric Critical Care Society

1693 RISING TO THE CHALLENGE OF COVID-19: HOW OUR PAEDIATRIC INTENSIVE CARE UNIT REFLECTED ON OUR EXPERIENCE CARING FOR ADULTS

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Background During the first wave of the COVID-19 pandemic our Paediatric Intensive Care Unit (PICU) was one of 7 in the UK to care for adults with COVID-19. This strategy allowed the hospital to expand its adult critical care capacity. Towards