World Journal of Pediatric Surgery

Learning from parental experience in a neonatal surgical unit: a qualitative service evaluation

Anna Littlejohns , ^{1,2} Emile Crouzen, ² Rebecca Mernenko, ² Fiona Metcalfe, ^{2,3} Waaka Moni-Nwinia, ² Hemma Chauhan, ² Bethan Johnson, ² Douglas McConachie, ⁴ Elizabeth Lawson, ⁴ Victoria Tricklebank, ³ John G McElwaine, ¹ Gurdeep S Sagoo, ⁵ Liz McKechnie, ³ Gary Latchford, ^{6,7} Jonathan Sutcliffe²

To cite: Littlejohns A, Crouzen E, Mernenko R, *et al.* Learning from parental experience in a neonatal surgical unit: a qualitative service evaluation. *World J Pediatr Surg* 2023;**6**:e000596. doi:10.1136/wjps-2023-000596

➤ Additional supplemental material is published online only. To view, please visit the journal online (http://dx.doi.org/10.1136/wjps-2023-000596).

Received 8 March 2023 Accepted 31 May 2023



© Author(s) (or their employer(s)) 2023. Re-use permitted under CC BY. Published by BMJ.

¹Anaesthesia and Critical Care, Leeds Teaching Hospitals NHS Trust, Leeds, UK

²Paediatric Surgery, Leeds Children's Hospital, Leeds, UK ³Leeds Centre for Newborn Care, Leeds Children's Hospital, Leeds, UK

⁴Faculty of Medicine and Health, University of Leeds, Leeds, UK ⁵Academic Unit of Health Economics, University of Leeds, Leeds, UK

⁶Institute of Health Sciences, University of Leeds Leeds, Leeds, UK

⁷St James's University Hospital Department of Clinical and Health Psychology, Leeds, UK

Correspondence to
Jonathan Sutcliffe; jonathan.
sutcliffe@nhs.net

ABSTRACT

Objectives Patient experience is directly related to health outcomes, and parental experience can be used as a proxy for this in neonatal care. This project was designed to assess parental experience of neonatal surgical care to inform future service developments and improve the care we provide.

Methods This was a qualitative study using rapid qualitative analysis. The study was carried out in a large neonatal surgical intensive care unit in the UK. Parents of infants treated by the neonatal surgical team between March 2020 and February 2021, during the COVID-19 pandemic were included. Purposive sampling was used to ensure that a representative range of parents were interviewed. A semistructured interview was created and tested in a previous phase of work. This questionnaire was used to ask parents open questions about different aspects of their infants' healthcare journey from the antenatal phase through to discharge from the neonatal unit (NUU). **Results** Rapid qualitative analysis was employed. and parental experiences were grouped into five main categories: before admission to the NNU, initial admission to NNU, information and support, COVID-19 and discharge. Within these five groups, we highlighted positive experiences to be fed back to the healthcare teams to reinforce good practice, areas that warranted improvement and suggestions for service development. Conclusions The wealth of data generated from the

Conclusions The wealth of data generated from the interviews has been summarized and shared with healthcare teams who are putting the service improvement suggestions into practice. The tool is available for services that wish to measure parental experience.

INTRODUCTION

Positive patient experience is associated with improved health outcomes¹ as well as being intrinsic to the delivery of humane care. Within the neonatal setting, parental experience can be used as a proxy for patient experience.² The Neonatal Critical Care Review emphasizes the need for enhancing the family experience.³ Nevertheless, the 'Getting It Right First Time' (GIRFT) report for pediatric general surgery and urology in

WHAT IS ALREADY KNOWN ON THIS TOPIC

Parental experience is hugely important in the neonatal surgical setting, particularly in the context of family integrated care. However, formal assessment of this is lacking.

WHAT THIS STUDY ADDS

⇒ This is the first use of our recently-developed interview tool to gain an in-depth understanding of parental experience on the neonatal unit from admission to discharge.

HOW THIS STUDY MIGHT AFFECT RESEARCH, PRACTICE OR POLICY

⇒ Using the observations of a representative range of families, key areas of good practice and areas for improvement have been identified, leading to actionable suggestions for service improvements. Workstreams have been created to implement these improvements. Since these improvements were based partly on the perspective of 'difficult to access' groups, they may be more likely to successfully improve parental experience for the whole cohort of families we look after. In turn this is likely to reduce health inequality, be more effective and reduce wasted effort. While our findings can be adapted to assess and improve neonatal surgical care in other centers, the tool may be applied to other settings to define and address the needs of patients, service providers and other key stakeholders.

England and Wales acknowledges that 'the method of collecting patient experience data is lacking for pediatric surgery', ⁴ and the National Institute of Health and Care Excellence (NICE) guideline 'Babies children and young people's experience of healthcare' notes that since particular groups may be less likely to provide feedback, their views should be actively sought.⁵

Family integrated care (FICare) is an important component of modern neonatal practice. It establishes parents as partners in care by providing education and psychosocial



support to enable them to gain confidence and become their infant's main caregiver. FICare improves health outcomes, including parental experience, ⁶⁷ and our unit introduced a model of FICare in 2017.

In March 2020, the SARS-CoV-2 (COVID-19) pandemic abruptly changed the delivery of healthcare. A large-scale review demonstrated that the restrictions significantly negatively affected the care provided for neonates and led to a poor experience for parents, the wider family and healthcare professionals. The authors highlight how bonding and developmental care practices suffered and articulate the unique characteristics of high-quality neonatal care and the extreme vulnerability of many neonatal patients. A key message was that an in-depth understanding of the unintended consequences that COVID-19 has had in a neonatal setting was needed. There was also a need to create tools and guidelines to be able to adapt to any ongoing or future changes.

Our project was designed in the early stages of the pandemic to capture how parental experience of neonatal surgical care had been affected and to inform future service developments.

METHODOLOGY Participants

Participants in the study include parents of infants treated by the neonatal surgical team during the COVID-19 pandemic, between March 2020 and February 2021 were reruited in this study.

Design

A semistructured interview was developed and tested (online supplemental table 1). Key characteristics of patients and families had been defined previously to guide purposive sampling. 10 Information about the project was advertised using posters, social media and through the neonatal unit (NNU) staff. Families interested in participating were provided with written and verbal information available in a range of languages. Each interview was conducted by two members of the project team. Audio recordings were taken to enable accurate transcription. One interviewer transcribed the interviews verbatim. The interview team comprised 10 members and included nurses, a nurse manager, a trainee advanced clinical practitioner, and trainee and consultant surgeons and neonatologists. All were trained and coached in interview techniques and qualitative analysis by a clinical psychologist.

The project was delivered without funding except for translation services, supported by our pediatric surgery department.

Patient and public involvement

Parents are used as proxies for the patients in the neonatal setting. Parents have been integral to this work from design to completion. Parents were key stakeholders involved in the initial design, creation and cognitive testing of the interview tool. A different group of parents

were the participants who were interviewed to generate our results and they have given suggestions for service improvements going forwards.

Analysis

Qualitative analysis is typically complex, time-consuming and arguably unsuitable in situations where information is sought quickly, such as during a health crisis. Thus, the project was informed by the 'rapid assessment process' and particularly 'rapid qualitative analysis', which was adapted for this study. Traditional qualitative methods involve detailed and time-consuming analysis of transcriptions which the researcher reads and re-reads repeatedly while they identify emergent themes and their relationship to each other, attempting to capture the experiences of participants. In contrast, rapid qualitative analysis is a form of 'top-down' analysis where many of the parameters are defined from the start, it is designed to answer specific questions about the service rather than produce a theoretically driven account of patient experience.

The interview questions provided the framework for the analysis. Each question was summarized using a neutral domain name, for example, 'preparation for leaving the unit'. A summary template was written listing all questions and domains, with columns for participant responses and quotations. An example of a summary template used in the analysis (online supplemental table 2). Any responses that did not fit existing domains were added to a new category. The summary template was piloted by the six members of the analysis team on one interview to test suitability. Minor changes were made to domains and an extra column detailing possible service implications was added. When consistency was established, transcripts were divided between the team for analysis, with each analyzed independently by two team members who agreed on a final summary for each participant. All summaries were combined to produce an initial matrix for all participants. The matrix was then divided between five members of the team who produced summary matrices for five aspects of the patient journey: before admission to the NNU, initial admission to the NNU, information and support, COVID-19 and discharge. A final summary of parental themes and service implications was created.

RESULTS

Twenty-four participants were recruited. Each of the characteristics deemed important by the stakeholder analysis in the phase I work were represented by at least one of the families recruited (table 1). A total of 18 interviews were carried out, 6 interviews with parents together and 12 interviews with parents separately. A mixture of virtual and face-to-face interviews were undertaken as determined by participant preference. Interviews were typically between 30 min and 90 min. Descriptive results are presented further. The summary results tables (online supplemental tables 3–7).



Participant characterist	ics: infant		Participant character	istics: parent	
	Acute	96	T di dolpant ondi dotoi	1–5	54
	Acute	90		>5	42
Presentation (%)	Elective	4	Depravation index (1–10) (%)	Unknown	4
resentation (70)	0	38	(1-10) (70)	<20	8
	1	33		20–25	8
	2	4		26–30	21
	3+	21		31–40	33
lumber of siblings (%)	Unknown	4	Mother's age (years) (%)	40+	17
difficer of sibilings (70)	Yes	71	(70)	Yes	13
	165	7 1		No	83
fultiple morbidity (%)	No	29	Single parent (%)	Unknown	4
iuitiple morbidity (%)	Yes	25	Sirigle parent (%)	Single	21
	169	20		Cohabiting	38
				Married	33
				Separated/	33
				divorced	0
Intenatal diagnosis (%)	No	75	Marital status (%)	Unknown	8
<u> </u>	1–14	8	· · · · · · · · · · · · · · · · · · ·	White British	67
	15–31	42		Other European	17
				African	4
				Asian	4
ength of stay (days) (%)	>31	50	Ethnicity (%)	Unknown	8
	Yes	79		English	76
Care at another hospital				Other European	17
%)	No	21	First language (%)	Other	8
,	Yes	25	<u> </u>	None	4%
				Some high school	13
				High school	17
				College	17
				Bachelor's degree	13
Care in another lepartment within trust			Highest education	Master's degree	8
%)	No	75	level (%)	Unknown	29
•	Ward	13	,	Yes	58
	HDU	8		No	13
lighest level of care (%)	NICU/PICU	79	Internet at home (%)	Unknown	29
	Upper GI/thoracic	12.5	. ,	Yes	0
	. ,			No	67
Specialty (%)	Lower GI	87.5	Disability (%)	Unknown	33
	24–27	33	, ,	<20	21
	28–31	4		20–39	4
	32–35	21		40–59	38
Gestation at birth			Travel time to hospital	60+	8
weeks) (%)	36+	42	(min) (%)	Unknown	33

Continued

Table 1 Continued						
Participant characterist	ics: infant		Participant characte	Participant characteristics: parent		
	24–27	4		Yes	21	
	28–31	21		No	60	
Gestation at presentation	32–35	21		Unknown	29	
(weeks) (%)	36+	54	IVF (%)			
	24–27	0		Yes	8	
	28–31	4		No	88	
	32–35	4				
Current gestation (weeks) (%)	36+	83	Multiple pregnancy			
	Unknown	8	(%)	Unknown	4	

GI, gastrointestinal; HDU, high-dependency unit; IVF, In vitro fertilization; NICU, neonatal intensive care unit; PICU, paediatric intensive care unit.

Before admission to NNU

There was sometimes confusion among the parents who received a diagnosis postnatally as to why the diagnosis had been made antenatally, even for conditions not typically identified antenatally. A sense that a diagnosis may have been missed lowered confidence in the clinical team:

'We thought if anything would have been wrong, it would have shown on the amniocentesis.'

The lack of diagnosis led to an inability to prepare for admission:

'[It was] overwhelming at first...felt hysterical at first but calmed down.'

When a diagnosis had been identified antenatally, both partners being able to access counseling and appointments together was important:

'It is really upsetting to talk about it to be honest. I don't think I have processed it myself yet.'

'I had to do all the scans on my own which I found really upsetting.'

During admission to NNU

The initial transfer to the NNU was identified as a particularly stressful event:

'The transfer from one unit to another unit was the worst part of it all. It took a long time for the transfer to happen.' 'I think a midwife should have taken me to neonatal, that would have been very helpful; they could have then said 'this is [baby]'s mum, could someone please show her around.'

A recurrent theme was the importance of personalizing care:

Young mums need a bit more support and a bit more explanation.'

'I had had a C-section on a different ward and were unable to go anywhere, while [baby] was going to theatre, I could not come to [baby].'

Staff kindness was easily recognizable, and the importance of the broader team on health outcomes was emphasized:

'To say it was a really rubbish situation it was really lovely, they were all great, understanding.'

Comments on accommodation and non-clinical areas provided actionable information for service planning for established and potential new build:

'It's not a hotel but everything is perfect still.'

'Got permit for free parking but there was still not always a parking space. Drove around a lot.'

Comments on the clinical areas highlighted the need for increased privacy and natural light:

"... was ridiculous to deprive the babies of this much light."

The financial burden was acknowledged:

'It is extremely expensive to have a baby admitted to [hospital] due to the costs around it.'

The availability of a kitchen to help decrease costs for families, lockable cupboards, free coffee and tea, reliable free Wi-Fi and a video link to access the ward when needed were all regularly raised and described as 'a life-saver'. These interventions are relatively inexpensive ways to improve experience.

Information and support

Communication was a key theme, and overall, information was felt to be clearly presented and regularly updated:

'The amount of information about the baby was enormous; thanks to the quality of the team, I understand everything.'



Diagrams were felt to be particularly useful, for example, to help visualize aspects of anatomy. Some families felt that staff were not always available, for example, at weekends. Understanding how to access the team was not always clear, especially early in the stay: what to do if the ward round did not enter their cubicle, for example.

Honesty was highly valued, including for bad news:

'I didn't feel that they were keeping any secrets. That's sometimes what you worry about, is there something they are not telling me.'

However, when there had been a loss of trust, this had important consequences:

'There was an incident when [baby] got an abscess from a cannula that was inserted. We did not know how honest they were about this. ... but it felt like they were trying to shove it under the carpet in fear of us complaining.'

Continuity of care across staff groups appeared important:

'It is almost like you have the same nurse for the whole time really that you are in hospital because they all know what is going on and it is great.'

Most knew and appreciated the offer of talking to a counsellor, but the team, particularly the nurses, were also an important source of support. Information about the experience of bereavement was striking; small things mattered and had a lasting impact:

'The stuff they did before [X] died and afterwards, it was never too much. Like we left some of his clothes here and they posted them out to us... I don't think I could have been as strong as I was if they weren't as strong as they were as well.'

Participants with limited English appreciated the efforts to find interpreters but expressed a preference for quicker solutions:

'I would be happy with Google translate sometimes.'

COVID-19 impact

Parents generally felt they and their baby were safe although desperate to get them home to their 'little bubble'. They worried they were potentially vulnerable:

'COVID was always at the back of the mind.'

Strict implementation of infection prevention guidance was seen as reassuring. There was frustration over discrepancy between testing for visitors and staff:

'[We] Don't really believe in COVID but accept restrictions.' It was just strange that the medical staff were not being swabbed.'

Parents were able to clearly articulate the impact of restrictions:

'It must be horrible for babies to just feel rubber and plastic all the time.'

'The only people who were touching her were medical staff. For 5 weeks we did not hold our baby, which made the bond we initially had disappeared. The only people touching her were medical staff for interventions.'

Discharge

Parents generally felt well prepared for discharge; they noted help with paperwork and good follow-up from the surgeon and the team. Support from clinical nurse specialists, the community team and outreach nurses was important. This included emotional support, noted particularly by one parent who sadly lost their baby:

'There is nothing I can imagine that could have helped us more.'

The distress generated when discharge was delayed was striking. Provision of 'goal-based' criteria for discharge might allow less focus on a specific time:

It is a strange feeling to have your baby at home after long invasive support, which is why still having access in the community is essential.'

With the benefit of hindsight, parents often acknowledged that they were not as well prepared psychologically as they thought, and they have become hypervigilant:

'A bit of a shock to the system, but nothing more could've been done.'

'I just have to be more careful and really watch my baby for any signs.'

Service improvements

The insights into parental experience have led to local service improvements and influenced projects already underway. Various workstreams were created to formulate meaningful responses to the parents' concerns, thereby developing our service and improving the care we provide. A summary of the resultant improvements is shown in table 2. A comprehensive list of the improvements made and further improvements planned can be found in online supplemental tables 3–7.

DISCUSSION

The value of measuring parental experience in the neonatal setting is well recognized. Despite this, NICE and GIRFT have highlighted the lack of mechanism for collecting these data and make a clear recommendation that this should be addressed.⁴⁵

The impact of COVID-19 on neonatal care is emerging in the literature. Given the importance of FICare, it is understandable how visiting restrictions have been particularly detrimental in this setting. Work from across the globe has demonstrated how restrictions have negatively impacted parental well-being, increased parental concerns about bonding and childhood development, and reduced parental confidence in caring for their infant. 9 14-16 Our study adds support to these findings and

Table 2 Examples o	Table 2 Examples of service improvements			
Aspect of patient journey	Examples of service improvements informed or instigated			
Before admission to NNU	 Monthly fetal medicine multidisciplinary team meetings and 6 monthly reviews of fetal medicine clinic. Video tour of NNU now available prior to admission. Midwives now bring mothers to the NNU on their first visit, and the family care team provides orientation to the unit. 			
During admission to NNU	 Psychology service now in place. Parents are actively encouraged to attended ward round. Their attendance is recorded and audited across the service. Badgernet video diaries used frequently to connect families with their baby when not on the unit. New screens purchased to help provide more privacy. Parking permits available for families, some designated parking spaces protected for neonatal family use. 			
Information and support	 Joint neonatal and surgical ward rounds from Monday to Friday and weekly multiprofessional team meetings to have holistic oversight of progress and ongoing care planning. Every patient has a named neonatologist and named surgeon, with this displayed by the bedside. Poster with details and picture of staff uniforms and job roles to help families understand different staff roles. Hospital chaplaincy team starting to facilitate weekly coffee morning for families. Our new NNU podcast 'Unexpected Beginnings: The Neonatal Unit'. This is hosted by veteran neonatal parents and runs through key aspects of being a parent on the NNU to provide support for other parents. 			
COVID-19 impact	 Regular communication and letters given to all parents in relation to any infection prevention control issues. Parental feedback on COVID-19 concerns disseminated to all teams involved in neonatal care. 			
Discharge	 Implementation of criteria-led discharge to help manage parental expectations and reduce delays on day of discharge. Multidisciplinary discharge meetings arranged for more complex infants, district general hospital teams invited virtually if there is a surgical neonate returned to local center. Weekly 'discharge huddle' to discuss patient flow, outstanding tasks and any family needs. Extra basic life support training sessions for staff so more staff is able to support parental training and reduce delays on discharge. 			
NNU, neonatal unit.				

offers further insight from groups that are representative of the range of families we look after, including families traditionally seen as 'difficult to access'. We have interviewed parents with a range of characteristics (eg, deprivation index, health literacy, first language, single-parent and two-parent families, ages, and distance travelled) whose infants themselves had a range of characteristics (eg, gestational age, disease complexity, and length of stay). We believe that this has made the information obtained more likely to reflect real-world experience and make the attempt to improve service delivery more likely to be effective.

It is valuable to have the perspective of families who have observed the service closely for many hours. Many of the issues raised have simple solutions, and we are collaborating with healthcare teams to develop and implement change. A number of the areas for improvement can be addressed by building on the 'culture and values' already in place. Other recommendations, such as the importance of kitchens, can be used to inform the plans for our new children's hospital, currently under

development. The insights into the direct and indirect impact of COVID-19 will inform preparation for potential future challenges. However, despite the context of the pandemic, most of the themes were not related exclusively to COVID-19. Therefore, we believe our results can give insight more broadly into the general experience of parents on the surgical NNU.

The amount of information obtained was extensive. This is hard to present academically and challenging to manage clinically. The main value of this project is taking these results forward so they can lead to service development. Several workstreams have been created, for example, the antenatal workstream, with input from relevant healthcare groups to prioritize the themes and actionable improvements.

From the beginning of this project, we understood the potential for scalability to other clinical pathways. Other aspects of care (clinical outcomes, process, and resource use) might usefully be considered by assessing cognitive diversity, and the methodology used here may be applicable. A recent study into crisis leadership in the



health service in Slovakia during the pandemic found that having a detailed understanding of the perspective of representative stakeholders allowed cognitive diversity and was a source of trust, satisfaction, and engagement in medical teams and helped inform rapid decision making.¹⁷ As in many fora outside medicine, it seems likely that capturing a variety of perspectives, and understanding the different needs and thought processes of a curated range of people will usefully add rigour to how services are designed.

We hope our interview tool and findings can be of use in other settings. While some observations may be specific to our center, others may be directly applicable elsewhere. The interview tool, however, is something that can be used widely.

Acknowledgements We thank all the parents who gave their time to this work and are very grateful to them for sharing their experiences. We also thank Bhanumathi Lakshminarayanan (consultant paediatric surgeon, Leeds Children's Hospital), Jenny King (chief research officer, Picker Institute), Suzanne Abrahams (general manager, Leeds Children's Hospital), John Ingleson (perioperative practitioner and branch chair of UNISON) and Sean O'Riordan (consultant paediatrician and COVID-19 lead for Leeds Children's Hospital) for their input, guidance and support.

Contributors All authors (AL, EC, RM, FM, WM-N, HC, BJ, DM, EL, VT, JGM, GSS, LM, GL, JS) were involved in the conceptualisation of the work. GL, LM and JS led on study design. EC, HC, BJ, AL, GL, FM, LM, RM, WM-N and JS contributed to the data collection. EC, AL, FM, LM and JS contributed to data analysis, led by GL. AL, GL, LM and JS led on the written manuscript, with contributions from all authors (EC, RM, FM, WM-N, HC, BJ, DM, EL, VT, JGM, GSS) and all authors have approved the final version of the submission. JS acts as guarantor and takes full responsibility for the work.

Funding The authors have not declared a specific grant for this research from any funding agency in the public, commercial or not-for-profit sectors.

Competing interests None declared.

Patient consent for publication Not applicable.

Ethics approval This study involves human participants, but upon review by the Trust Research and Innovation Research Governance Lead, this project was confirmed as a service development project and thus did not require Health Research Authority or NHS Research Ethics Committee approval. Approval for this project was also obtained from the Trust Caldicott Guardian. All participants gave informed consent to participate in the study before taking part.

Provenance and peer review Not commissioned; externally peer reviewed.

Data availability statement Data are available upon reasonable request.

Supplemental material This content has been supplied by the author(s). It has not been vetted by BMJ Publishing Group Limited (BMJ) and may not have been peer-reviewed. Any opinions or recommendations discussed are solely those of the author(s) and are not endorsed by BMJ. BMJ disclaims all liability and responsibility arising from any reliance placed on the content. Where the content includes any translated material, BMJ does not warrant the accuracy and reliability of the translations (including but not limited to local regulations, clinical guidelines, terminology, drug names and drug dosages), and is not responsible for any error and/or omissions arising from translation and adaptation or otherwise.

Open access This is an open access article distributed in accordance with the Creative Commons Attribution 4.0 Unported (CC BY 4.0) license, which permits others to copy, redistribute, remix, transform and build upon this work for any purpose, provided the original work is properly cited, a link to the licence is given, and indication of whether changes were made. See: https://creativecommons.org/licenses/by/4.0/.

ORCID iD

Anna Littlejohns http://orcid.org/0000-0003-1042-4096

REFERENCES

- 1 Doyle C, Lennox L, Bell D. A systematic review of evidence on the links between patient experience and clinical safety and effectiveness. *BMJ Open* 2013;3:e001570.
- 2 Burger SA, King J, Tallett A. Parents' experiences of neonatal care in England. *Patient Experience Journal* 2015;2:45–52.
- 3 NHS England and NHS Improvement. Implementing the recommendations of the neonatal critical care transformation review. 2019. Available: https://www.england.nhs.uk/publication/implementing-the-recommendations-of-the-neonatal-critical-care-transformation-review/ [accessed 29 Jun 2023].
- 4 Simon K. Paediatric general surgery and urology GIRFT Programme National Specialty Report GIRFT clinical lead for paediatric surgery. 2021. Available: https://gettingitrightfirsttime.co.uk/surgical_ specialties/paediatric-general-surgery-and-urology/ [accessed 26 Jun 2023].
- 5 Babies, children and young people's experience of Healthcare NICE guideline [Internet]. 2021. Available: www.nice.org.uk/guidance/ ng204 [Accessed 26 Jun 2023].
- 6 Church PT, Grunau RE, Mirea L, et al. Family integrated care (FICare): positive impact on behavioural outcomes at 18 months. Early Hum Dev 2020;151:105196.
- 7 O'Brien K, Robson K, Bracht M, et al. Effectiveness of family integrated care in neonatal intensive care units on infant and parent outcomes: a multicentre, multinational, cluster-randomised controlled trial. Lancet Child Adolesc Health 2018;2:245–54.
- 8 Ryan L, Plötz FB, van den Hoogen A, et al. Neonates and COVID-19: state of the art: neonatal sepsis series. *Pediatr Res* 2022;91:432–9.
- 9 Muniraman H, Ali M, Cawley P, et al. Parental perceptions of the impact of neonatal unit visitation policies during COVID-19 pandemic. BMJ Paediatr Open 2020;4:e000899.
- 10 Mernenko RK, Littlejohns A, Latchford G, et al. Developing a method to capture parental experience in a neonatal surgical centre in the context of COVID-19: a qualitative study. BMJ Paediatr Open 2022:6:e001434.
- 11 Beebe J. Rapid assessment process: an introduction. AltaMira Press, 2001. Available: https://archive.org/details/rapidassessmentp 0000beeb [accessed 5 Jul 2023].
- 12 Hamilton A. Qualitative methods in rapid turn-around health services research. 2013. Available: https://www.hsrd.research.va.gov/for_ researchers/cyber_seminars/archives/780-notes.pdf [accessed 29 Jun 2023].
- 13 Hamilton AB, Finley EP. Qualitative methods in implementation research: an introduction. Psychiatry Res 2020;283:112629.
- 14 Kostenzer J, Hoffmann J, von Rosenstiel-Pulver C, et al. Neonatal care during the COVID-19 pandemic - a global survey of parents' experiences regarding infant and family-centred developmental care. EClinicalMedicine 2021;39:101056.
- 15 Garfield H, Westgate B, Chaudhary R, et al. Parental and staff experiences of restricted parental presence on a neonatal intensive care unit during COVID-19. Acta Paediatr 2021;110:3308–14.
- 16 Kynø NM, Fugelseth D, Knudsen LMM, et al. Starting parenting in isolation a qualitative user-initiated study of parents' experiences with hospitalization in neonatal intensive care units during the COVID-19 pandemic. PLoS One 2021;16:e0258358.
- 17 Joniaková Z, Jankelová N, Blštáková J, et al. Cognitive diversity as the quality of leadership in crisis: team performance in health service during the COVID-19 pandemic. Healthcare (Basel) 2021;9:313.

Date and Time of Interview

Consent recorded? Y/N

Purpose of the interview and what we want them to do

Set out the main areas there will be questions so each get enough focus (e.g, before admission and early on, then during the stay, then communication, COVID-19, and thoughts around leaving the unit)

Thinking about when you were pregnant and when your baby was first born

- 1- What were your main worries about the thought of coming into hospital?
 - a. Were you worried about the pandemic, or did the pandemic have any other affect?
- 2- Did you know what to expect for you and your baby's care once your baby was born?
 - a. Did you think that the pandemic might have an effect?
- 3- Did you know who the team looking after your baby were?
 - a. Did you know which consultant neonatologist and which consultant surgeon was in charge of your baby's care, and how the 2 link?
- 4- Were you made to feel welcomed by the team looking after your baby?
 - a. Did you feel that they were approachable?

Thinking about the neonatal unit

- 1. What are your thoughts on the room(s) where your baby was?
- 2. What do you think about your privacy while on the ward?
 - a. Do you think the pandemic affected your privacy?
- 3. Do you have any other comments regarding practical things that made a difference to you? For example, the parents' accommodation, refreshment areas, anything to do with your ability to travel to and from the hospital to visit or parking?
 - a. Did you notice any effects on these that might have been caused by the pandemic?
- 4. Is there anything else about practicalities of being on a neonatal unit we should have asked you?

Considering the information and support you received

- 1. For your personal circumstance did you need additional help (for example an interpreter) and was this provided for you?
- 2. Did you understand your baby's diagnosis and treatment as much as you would have liked?
 - a. Were you able to talk to nursing staff and the doctors as much as you wanted?
- 3. How do you feel about the amount of information you were given?
 - a. Too much? Too little?
- 4. Were you ever given conflicting information?
- 5. Were you aware of how to access emotional and practical support for you before, during or after your baby's admission? Was there anyone who was a 'constant' person all the way through?

- 6. Is there anything you would have liked to have known that wasn't in the information given to you?
- 7. Do you think the pandemic affected the communication you received?
- 8. Were the rules about COVID-19 easy to understand?
 - a. Were the rules always followed the same way by different staff members or families? Did you notice differences between rooms? What happened when rules weren't followed?
- 9. Do you recall the rules changing?
 - a. If so, how soon did you get to know?
- 10. Did you feel confident that information about your baby's care was shared well between the members of the team, including nurses, doctors and everyone else in the team?

Thinking about COVID-19

- 1. Did the pandemic change how safe you felt for you and your baby?
- 2. Do you think that the pandemic and its regulations influence how you and anyone else important for your baby (for example your partner) were able to bond with your baby?
- 3. Do you have other children? Has having your baby in hospital affected them?
 - a. Do you think that the pandemic affected this?
- 4. Are there ways in which you think the pandemic might have affected the way that staff cared for your baby?
- 5. Are there any other sources of stress that we have not asked about?

Leaving the unit

- 1. When you leave the neonatal unit, do you know if it will be to another ward, or to go home? How prepared do you/did you feel for discharge from the neonatal unit?
- 2. Did someone make you aware of the support that is available to you for when you do get home, for example, coming to A&E, seeing your GP, support from family?
- 3. Do you have any particular concerns for your baby once they are at home, having had surgery?
- 4. Are there any additional concerns or practical problems caused by the pandemic?

Time Interview Finished

Supplementary Table 1. Interview Questions

Question	Domain			
Thinking about when you were pregnant and when your baby was first born				
What were your main worries about the	General anticipatory worries (including effect of			
thought of coming into hospital?	COVID-19)			
Were you worried about the pandemic, or				
did the pandemic have any other affect?				
Did you know what to expect for you and	Expectations of newborn care (including effect of			
your baby's care once your baby was born?	COVID-19)			
Did you think that the pandemic might have				
an effect?				
Did you know who the team looking after	Knowledge of team			
your baby were?				
Did you know which consultant				
neonatologist and which consultant surgeon				
was in charge of your baby's care, and how				
the 2 link?				
Were you made to feel welcomed by the	Feeling welcomed			
team looking after your baby?				
Did you feel that they were approachable?				
Thinking about the neonatal unit				
What are your thoughts on the room(s)	Parental view: room			
where your baby was?				
What do you think about your privacy while	Parental view: privacy (including effect of COVID-			
on the ward?	19)			
Do you think the pandemic affected your				
privacy?				
Do you have any other comments regarding	Parental view: practicalities (including effect of			
practical things that made a difference to	COVID-19)			
you? For example, the parents'				
accommodation, refreshment areas,				

	T
anything to do with your ability to travel to	
and from the hospital to visit or parking?	
Did you notice any effects on these that	
might have been caused by the pandemic?	
Is there anything else about practicalities of	
being on a neonatal unit we should have	
asked you?	
Considering the information and support	you received
For your personal circumstance did you	Presence of additional help (if needed)
need additional help (for example an	
interpreter) and was this provided for you?	
Did you understand your baby's diagnosis	Understanding /ability to talk to team
and treatment as much as you would have	
liked?	
Were you able to talk to nursing staff and	
the doctors as much as you wanted?	
How do you feel about the amount of	Information amount
information you were given?	
Too much? Too little?	
Were you ever given conflicting	Presence of conflicting information
information?	
Were you aware of how to access	Access to emotional/practical support:
emotional and practical support for you	Before – During – After; consistency
before, during or after your baby's	
admission? Was there anyone who was a	
'constant' person all the way through?	
Is there anything you would have liked to	Unmet information needs
have known that wasn't in the information	
given to you?	
Do you think the pandemic affected the	Information about COVID-19
communication you received?	
Were the rules about COVID-19 easy to	Understanding of COVID-19 rules
understand?	(differences/changes/consistency?)
Were the rules always followed the same	
way by different staff members or families?	
Did you notice differences between rooms?	
	I

World J Pediatr Surg

What happened when rules weren't	
followed?	
Do you recall the rules changing?	
If so, how soon did you get to know?	
Did you feel confident that information	Parental view: information was shared between
about your baby's care was shared well	team
between the members of the team,	
including nurses, doctors and everyone	
else in the team?	
Thinking about COVID-19	
Did the pandemic changed how safe you	Feeling safe
felt for you and your baby?	
Do you think that the pandemic and its	Impact on bonding with baby
regulations influence how you and anyone	
else important for your baby (for example	
your partner) were able to bond with your	
baby?	
Do you think that the pandemic affected	
this?	
Do you have other children? Has having	Impact on siblings
your baby in hospital affected them?	
Do you think that the pandemic affected	
this?	
Are there wave in which you think the	Parantal vious impact on stoff
Are there ways in which you think the	Parental view: impact on staff
pandemic might have affected the way that	
staff cared for your baby?	
Are there any other sources of stress that	Other sources of stress
we have not asked about?	
Leaving the unit	
When you leave the neonatal unit, do you	Preparation for leaving unit
know if it will be to another ward, or to go	
home? How prepared do you/did you feel	
for discharge from the neonatal unit?	

Did someone make you aware of the	Awareness of support available
support that is available to you for when	
you do get home, for example, coming to	
A&E, seeing your GP, support from family?	
Do you have any particular concerns for	Concerns at home
your baby once they are at home, having	
had surgery?	
Are there any additional concerns or	Other concerns
	Other concerns
practical problems caused by the	
pandemic?	

Supplementary Table 2. An example of a summary template from our analysis

Before Admission					
Knowledge of the baby's condition					
Themes	Quotes	Actions	Service improvements		
Difficulty processing antenatal diagnosis The need for clear information in this	"Was told but didn't want to believe it" "Conflicting information carried on throughout the whole antenatal period If	Feedback to staff on real world positive and negative effect of antenatal counselling	Monthly fetal medicine MDT clinic- includes neonatologist, obstetrician, surgical nurse, midwife and paediatric surgeon		
period	you are already anxious, nervous and have no idea what is going to happen, it is	Need updated parent information leaflets	Fetal medicine clinic reviewed 6 monthly		
Potential for unintended consequences of knowing there may be an abnormality	"It is really upsetting to talk about it to be honest. I don't think I have processed it		Antenatal counselling- includes support information- paediatric immediate life support, practical advice		
Importance of balanced antenatal discussion to managing expectations	myself yet."		Close links with family care team on NNU- meet parents, orientate to the ward, answer practical questions		
			Video tour of NNU now available		
The value of trust in staff by parents who need confidence in professionals	"Felt safe already, staff brilliant" "I know someone from my hospital that was transferred here with their heart the wrong way round; so I know it is a good hospital"	Feedback to staff and the potential to roll out feedback network-wide	Family care team Poster about good communication tips for staff- considering parent feedback		
Confusion when some congenital abnormalities were not detected on antenatal scans	"We thought if anything would have been wrong, it would have shown on the amniocentesis" "overwhelming at first'felt hysterical at first but calmed down"	Need to conduct work to determine what this group of families need to know in the immediate, short term and long-term Consider postnatal meeting between foetal medicine and families if further explanation needed. This could include review of antenatal scans and review of performance to serve as audit	Neonatology team can link to obstetric team to meet parents for de-brief if required		

Quotes	Actions	Service improvements
"petrified" "didn't know what was going to hannen"	Video tour to show the rooms and incubator	Video tour of unit now done and being updated. This is shared antenatally
alan t know what was going to happen	'Supersibs' and sibling support packs potentially useful	Neonatal operational delivery network (ODN) work stream to include videos for other units
		Our neonatal unit podcast "Unexpected Beginnings- The Neonatal Unit". Hosted by veteran neonatal parents going through key aspects of being a parent on the neonatal unit to provide support for other parents, including an episode on 'Juggling Siblings'
	T	In
"I had to do all the scans on my own which I found really upsetting" "It's like being separate families as being split up so often!" "It was too much information to take in on your own" "Husband felt left out and couldn't do anything"	Consider video consultations as a standard offer when a parent is unable to attend because of COVID Consider partner in terms of ongoing bonding and long- term well-being of the child as well as parents	Regular communication and letters given to all parents in relation to any IPC issues Families kept updated
"scared that [baby] would get COVID-19 and she is obviously very vulnerable" "The rules kept on changing all the time. You would ask someone questions about our appointments regarding COVID-19 regulations, and nobody really knew the	Collaborate with local, regional and national teams Regional Q&A or FAQ as an efficient mechanism of information dissemination	Parental feedback on COVID-19 concerns and communication disseminated to all teams involved in neonatal care Presented regionally and nationally also
	"didn't know what was going to happen" Quotes "I had to do all the scans on my own which I found really upsetting" "It's like being separate families as being split up so often!" "It was too much information to take in on your own" "Husband felt left out and couldn't do anything" "scared that [baby] would get COVID-19 and she is obviously very vulnerable" "The rules kept on changing all the time. You would ask someone questions about our appointments regarding COVID-19	"didn't know what was going to happen" "didn't know what was going to happen" "Supersibs' and sibling support packs potentially useful "I had to do all the scans on my own which I found really upsetting" "It's like being separate families as being split up so often!" "It was too much information to take in on your own" "Husband felt left out and couldn't do anything" "scared that [baby] would get COVID-19 and she is obviously very vulnerable" "The rules kept on changing all the time. You would ask someone questions about our appointments regarding COVID-19 regulations, and nobody really knew the

Themes	Quotes	Actions	Service improvements
Parents found transfer very stressful and found it hard not to be allowed to travel with baby	"The transfer from one unit to another unit was the worst part of it all. It took a long time for the transfer to happen, but the people were lovely"	Feedback to local transport network and involve them	Members of research team are actively involved in Neonatal Operational Delivery Network (ODN) co-production work streams to include project outcomes in all work eg-transfer of surgical neonate information resources about all neonatal units in the region -align practiced regionally to reduce the variability of care across the regional NNU Working with the neonatal ODN and local neonatal transport service to help families access information and resources about transfer and the different type on neonatal
			units Our regional neonatal transport team routinely collect feedback on care and transport experience with a quick response (QR) code on the incubators/beds. As a transfer involves care in multiple centres thi inherently collects feedback on the differences between centres. Our unit is looking into using bedside QR codes as a toc to collect timely feedback.

Supplementary Table 3. Before admission

	Admission to Hospital					
Initial phase of care						
Themes	Quotes	Actions	Service Improvements			
Appreciation for welcoming team	"to say it was a really rubbish situation it was really lovely, they were all great, understanding"	Feedback to all staff, especially nursery nurses, the very positive comments	Fetal medicine MDT workstream commenced to support service improvements- 6 monthly review meetings			
	'felt like a family'		Veteran parent volunteers back to support families			
			Metric to ensure parents spoken to in first 24 hours			
			Psychology service now in place			
			We need to reintroduce 'super sibs'- volunteer service supporting siblings on the neonatal unit			
The need more support around transitions and initial admission	"I started crying at the cot of a baby that was not mine, because I thought that was [baby]. That is how little information I had upon admission" "I think a midwife should have taken me to	Consider an 'Induction' for new families as we do for staff. Perhaps this could be a role of the surgical liaison neonatologist Consider video tours of the unit that are updated with any ward change with in	Welcome posters on NNU- practical information Family care team involvement- orientation to unit etc Named neonatologists/Surgeons			
	neonatal, that would have been very helpful; they could have then said 'this is [baby]'s mum, could someone please show her around"	future waves of the pandemic	Two neonatal consultants are now assigned as designated links to the surgical team Communicated to midwife team to bring			
			mother on 1 st visit			
			Plan to develop photo books of the unit/equipment etc			

			"Unexpected Beginnings- The Neonatal Unit" podcast episodes on 'what is a neonatal unit?' and 'meeting your baby for the first time'
During Admission			
Themes	Quotes	Actions	Service improvements
Physical and emotional challenges will exist for all new mothers following the birth; some will be particularly vulnerable Language barriers were addressed well. Interpreters ere offered and available but they were not always very effective. Some families preferred faster, easier and less official means	"The beads, Vcreate etc., white board were all very helpful. It was nice we could decorate the cot, it was all personal" "young mums need a bit more support and a bit more explanation" "I had had a C-section on a different ward and unable to go anywhere, while [baby] was going to theatre, I could not come to [baby], nobody was able to come and get me" "Language barriers are professionally dealt with" "Some interpreters difficult to understand" "I would be happy with google translate sometimes"	Tailor information and advice to the specific family. We need to be aware of specific groups and characteristics that may need tailored care. These could include; 'young' mums, family that conceived through IVF, those with language barriers, families with other health needs Feedback on interpreting services and their use. Think how best to use interpreting services; Offer translation in different ways (person/app/google) for day to day work. Need to consider information governance risks phone applications	Simple measures were very appreciated and helpful Badgernet videos regularly utilised Events to promote with staff on special occasions eg Mother's Day, Fathers Day, Easter, birthdays etc Staff utilising virtual interpreting services more often in addition to face-to-face: telephone, video, google translate

Operational issues				
Themes	Quotes	Actions	Service improvements	
Importance of clear introductions and clear channels of communication	"important to have one person you can talk to, one person you are happy with, you could go to talk to and trust" 'big unit, moved rooms a lot so didn't get to know many nurses well; Was told who surgeon was but did not meet until after surgery'	Named nurses and named consultants needed (medical and surgical) and ensure the link between the neonatal and surgical teams is clear Reminder to staff importance of introducing themselves and their role	Parents actively encouraged to attended ward round. Their attendance is recorded and audited across the service Named consultants Name boards at bed space for families to see Joint neonatal and surgical ward rounds daily Discussion with expert transport teams to develop feedback mechanisms so families can feedback on whole of healthcare journey	
Building, environment, a				
Themes	Quotes	Actions	Service improvements	
Appreciation of accommodation with the clean, spacious, private rooms nearby. This eased financial	"a life-saver" "it's not a hotel but everything is perfect still"	Acknowledge the financial strain associated for these families	Family care team and lead oversee all aspects of parental care & support	
worries	"It is extremely expensive to have a baby	Inform planners of new building that the kitchen is very important, and parents	Family integrated care well established on NNU	
Appreciation for WIFI	admitted to LGI due to the costs around it"	need their own space within this. And note the importance of wifi to the Digital	Financial support from hospital charity for families	
Travelling and parking a common source of stress as well as the indirect expense of the admission	"Got permit for free parking but there was still not always a parking space. Drove around a lot"	Leeds project mentioned by commissioners No easy parking solution. Perhaps promote public transport and be clear	Some meal and refreshment provision. Ability to order parent food out of hours for families transferred in	
		with parents what the parking situation is like, so they know what to expect	Families have access to hospital parent accommodation/ facilities-free of charge. Designated parent facilities-kitchen, sitting/dining room	

			Free parking permits available for families. Some designated parking spaces protected for neonatal family use
Issues with building structure and lack of privacy	"all parents could hear about our child and we knew about theirs" "There was no natural light. I could not believe the room would be designed like this, and this was ridiculous to deprive the babies of this much light." "The rooms are so loud, what with the building being this old, which is bad for the babies I believe."	With the planned new build, we need a mix of single rooms and open bays, private space for families and more reclining chairs Ensure appropriate access to screens/rooms/secure environment for expressing	New screens purchased to help provide more privacy in surgical newborns area. Screens have an 'under the sea' theme to match the room décor. The screens are wall mounted between each bed space There is a designated breastfeeding room and parent consultation room already in place for additional privacy

Supplementary Table 4. Admission to hospital

Information and Support on the NNU				
General				
Themes	Quotes	Actions	Service Improvements	
Information was clearly, presented, diagrams were useful	"The amount of information about the baby was enormous; Thanks to the quality of the team, I understand everything"	Encourage use of diagrams, have some standard diagrams of normal anatomy and some showing abnormalities we encounter what we can then draw on	Neonatal project work stream set up to help review and implement service improvements Photo books planned to be developed- linking	
Parents distressed when they searched online to find	"It is difficult to understand; diagrams have solved all this; It is not the difficulty of the information	Ensure good written and web resources for	in with ODN co-production work streams	
information	that was upsetting, but the content and how rushed everything inevitably was"	parents provided or signposted	Surgical outreach signpost parents to resources/websites/charities	
	"a bad move because you only ever remember the bad stuff from those websites"		All parent information leaflets reviewed for quality of information and will be available shortly for families and staff	
Parents felt the team were always honest, even if giving bad news	"as much as I didn't want to hear it, looking back I think well, I needed to hear it. As much as it's a job for them, they're doing the best possible thing which is being honest."	Feedback to staff their candour is noticed and appreciated	Continue this care	
	"I didn't feel that they were keeping any secrets. That's sometimes what you worry about, is there something they are not telling me."			
Care felt personal	"is almost like you have the same nurse for the whole time really that you are in hospital because they all know what is going on and it is great" "to have a bit of a giggle amid all the seriousness	Aim for consistency in staff over different shifts	Positive feedback to staff given to share feedback from families and to encourage continues good practice for supporting families	
	felt very human"		We need to review process for continuity of care from nursing team	
	"The family care team was absolutely amazing. Always present, every single day. 'Do you need this', 'do you need that'. It was brilliant"			
Feeling of being overwhelmed by having an unwell baby	"I basically said goodbye to him before surgery thinking the worst. I completely understand that they would paint the worst picture"	Prepare parents as best as possible with explanations, photos and videos	Development of psychosocial MDT underway - links with unit safeguarding nurses	

Supplemental material

me poor communication	"I never knew anything about neonatal medicine, so it was completely new" "We never got the amount of information about the stoma we wantedNobody came to sit down with us about that procedure before it happened"' "Communication could have been better and promises should be kept: Excluded from room during precious visiting time because of procedures with other babies; procedures delayed and parents not updated as promised" "Would like to have spoken to surgeons more, the surgeon actually doing operation we only caught by chance and we had about 30 seconds to speak to them"	We need to work on improving communication. Specific areas highlighted that we need to be aware of include: - Anomaly screen is particularly stressful – i.e., looking for other abnormalities when one has been found - Less information is available at weekends - Parents sometimes too tired to take things in – need things repeating - Hard to hear staff when wearing a mask	Better joined up MDT working to help with communication, decision making and consistent care planning including: - Joint neonatal and surgical ward rounds Mon-Friday - Weekly multi-professional team meeting to have holistic oversight of progress ongoing care planning Named consultants for each surgical neonateneonatal and surgical - to lead with care, communication and decision making. Named consultants meets with family for updates Name boards at cot-side so families know who named consultants are
			Parents encouraged to attend ward rounds Audit on parent presence on ward rounds Nursing staff advocate for families and request meeting with surgeon/neonatologist if family request Poster with details and picture of staff uniforms and job roles to help families understand different staff roles Poster shared with staff with helpful 'top tips for good communication'. This was shared in Neonews (our weekly neonatal newsletter)

Conflicting information presents a large problem	"I saw 4 consultants on different days, and everybody wanted to do something"	Need to improve internal communication for consistency	As previous
	"If you change the plan every single day, how will you know if it was working or not?	If there are different opinions or the plan needs to change then we need to explain why	
	"The teams did not really communicate well. They would tell us one thing and then 5 hours later they told us they were not going to do it anymore"		
	"It was a bit cross-wired, because someone had said he had had a good night, but then it turned out he had not had a good night and was sick a lot"		
Specific			
Parents were aware of the counsellor but low uptake of	"might have been easier if it was in person. Talked to the staff instead"	Needs to be more visible and accessible	Psychology service now embedded on NNU
service	"the nurses' job is even harder now, where you have to be additional emotional support because	Nurses seen to be very supportive, but particularly important for families when isolated from other support	Hospital Chaplaincy team starting to facilitate weekly coffee morning to support families
	the parents don't have it from anywhere else"		Lots of resource posters with QR codes to help signpost families to support
			Hospital charity funds available to apply for families needing financial support
Bereavement care	"The stuff they did before X died and afterwards, it were never too much. Like we left some of his	Feedback to staff	Funding for a bereavement nurse- not in post yet
	clothes here and they posted them out to us I don't think I could have been as strong as I was if they weren't as strong as they were as well"	Encourage ongoing training with butterfly trust	Close links established with regional hospices
			Bereavement clinical room available
Video links are helpful in reducing anxiety	"a God-send and helped us through For any parent, mum or dad, it is the best thing"	Extend the utilisation of virtual platforms to support families	Bagdernet video diaries used frequently to connect families with their baby when not on the unit

Supplementary Table 5. Information and support on the neonatal unit

	Impact of C	COVID	
COVID-19 inform	ation		
Themes	Quotes	Actions	Service Improvements
Parents generally felt well informed and accepted the rues	"there were signs everywhere. I was told in advance when I had to get my covid test, when dad was supposed [to get his]" "Don't really believe in Covid but accept restrictions"	Be transparent regarding uncertainty or when things have changed	Regular communication and letters given to all parents in relation to any infection prevention and control (IPC) issues
Frustration about rule			, , ,
changes	"The rules kept on changing all the time You would ask someone questions about our appointments regarding COVID-19 regulations, and nobody really knew the answer"		
COVID-19 worries	5		
Themes	Quotes	Actions	Service Improvements
The potential risk to the baby was a constant additional worry	"Covid always at the back of the mind"	Be mindful of the impact of covid and how worrying it is	The COVID-19 team at Leeds Children's Hospital provided written information
			More psychological support available
What families the	ought went well		
Themes	Quotes	Actions	Service Improvements
Feeling safe that the staff followed rules and visitors were not allowed	"tested regularly - what more could they do - Really good, doing everything" "I did not feel unsafe at any time. I have seen the staff cleaning their hands, changing their aprons, gloves etc"	If any further waves we should be reassured our measures made parents feel safe Acknowledge previously unrecognised	More information about testing
	"I think the pandemic made it kind of safer"	positive impact of the infection control rules eg some parents enjoy stricter visiting policies	
Frustration that there were sometimes double standards – social distancing for parents	"told off for not social distancing but then the staff were all round the work station — looks like work most of the time but not always! Didn't like that"	Staff must follow rules	Written information was created which acknowledged that rules may change over the course of the pandemic. This recognised the

but staff congregate	"It was just strange that the medical staff were not being	Be open to people pointing out	variability between centres and
around station	swabbed"	discrepancy and be able to explain or	over time.
	"There did seem to be a little discrepancy regarding family	correct	
Discrepancies between	visits"		Written information also
hospitals		Need consistency of rules between	explained that beyond COVID-19
	"We were so happy that the rules in Leeds were more lenient	different wards and units or	there are differences between
	towards that"	explanation for differences	hospitals, and it is not possible
		·	and not always appropriate for
		Comparison with other centres needs	centres to be the same
		feedback to network real time to	
		identify useful differences	Paediatric Grand Rounds provided
			a time to highlight any concerns
The restrictions caused	"Sibling only sees on facetime/photo. Doesn't talk about baby"	Nurses very engaged in helping	'Sibling packs' created to ensure
stress. Parents worried		siblings bond – positive feedback for	siblings that were unable to visit
about their other	"siblings didn't see twin before he died"	build a bear	had activities to involve them in
children not bonding			care
with the baby and	"these are Newborns the first few weeks of their lives are critical	Acknowledge impact on siblings and	
worried about the	for bonding"	offer advice	Understanding of the importance
development of the			of relatives visiting for end-of-life
baby who is only	"they learn a lot from facial expressions"	In end-of-life cases we should	care led to more visiting being
interacting with people		reconsider sibling visiting. The siblings	allowed in end-of-life care as the
hidden behind personal	"the only people who were touching her were medical staff. For	weren't able to visit a dying sibling,	pandemic progressed
protective equipment	5 weeks we did not hold our baby, which made the bond we	they had to wait until he had already	
	initially had disappear only people touching her were medical	died and was in a hospice. This seems	
	staff for interventions"	very difficult and needs addressing	
	"It must be horrible for babies to just feel rubber and plastic all		
	the time"		
	"the nurses got [sibling] a Build-a-Bear, a storybook, which was		
	brilliant, and helped us massively. [Sibling] is 5 years old. He		
	realises he has a brother, and is able to give him little presents,		
	and keeps being interested in all the updates, but has never		
	managed to see him in real life"		
Supplementary Table 6 Im		l	

Supplementary Table 6. Impact of COVID-19

Discharge Preparedness for leaving the unit				
		Ensure family utilise Family Integrated Care documentation re discharge planning Consider wider family in discharge planning- utilise virtual forums for training/information Need clear discharge criteria and communicate this to family. If discharge is 'goal based' then families will be perhaps less focussed on an agreed time that may then change Reality of discharge different than expected – we need to work on this transition and managing expectations	Service Improvements Multi-disciplinary discharge meetings arranged for more complex infants Family integrated care imbedded in the ethos of care delivery. Discharge planning document utilised Weekly 'discharge hudddle' to discuss patient flow and what needs to be completed and any family needs, teaching, information etc plus weekly MDT ward round to highlight discharge needs. Feedback to the neonatal surgical outreach team the positive feedback on their involvement Extra BLS training sessions for staff so more staff able to support parental training and reduce delays on discharge Implementation of criteria led discharge-helps manage parental expectations and reduce delays on day of discharge	
Managing bereavement of a newborn	"There is nothing I can imagine that could have helped us more than was already done for us after the baby's passing"	Continuity of care important for end-of-life care		

Adjusting to life at home	Adjusting to life at home				
Theme	Quotes	Action	Service improvements		
Re-admittance can be unexpected	"in safe hands" "We felt prepared to go home. We just had not anticipated that he would have to be readmitted	Continue to signpost clear communication channels for families at home	Current service review of Neonatal surgical outreach provision and what service needs are		
and difficult - some happy if needed to come back in, some not as prepared as they thought	to hospital. I do not think anyone could have known"	Acknowledge that readmissions emotionally difficult - need preparation	Aim to improve parent contacts and availability for specialist discharge planning and education		
Many parents had built up a high level of trust in the hospital team, and sometimes worried about a perceived lack of expertise in their local hospital/ health visitor when home	"Even now that we have been discharged, we are always assured to have contact with the ward in case something goes wrong or for more questions"	Professional conversations with family needed to reduce lack of trust in other organisations	Outreach nurses go to district general hospitals (DGHs) to facilitate earlier repatriation and joined up care with DGH and tertiary centre. DGH team invited to MDT virtually if there is		
			a surgical neonate returned to local centre		
The difficulty of adjusting to life at home	"We had to get used to life out of hospital. All the alarms, beeping and those noises"	Feedback to community teams	Feedback to community team		
Feeling isolated at home on discharge	"It is a strange feeling to have your baby at home after long invasive support, which is why still	Make time to answer all questions prior to discharge	"Unexpected Beginnings- The Neonatal Unit" podcast episode 'Going home' explores issues around getting reading to go home,		
	having access in the community is essential"	Need to signpost sources of support for parents including outreach, GP, health	adapting to life away from the neonatal unit, managing the associated anxiety and the		
Appreciation for community teams	"I just have to be more careful and really watch my baby for any signs"	visitors, mental health support	role of the transitional unit in the process		
	"quiet but struggling"		Good coordination of discharge and communication facilitates. Needs some review and improvement		
	"There's always those things in the back of your mind and you don't want to sound an idiot to the doctors asking what if, what if, what if?"		·		
Conglessorters Table 7 Disabases	"We haven't really seen anyone since we came home It has just been us in our own little bubble. It is difficult some days"				

Supplementary Table 7. Discharge from hospital