

# Views of parents, adults born preterm and professionals on linkage of real-world data of preterm babies

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#### **ABSTRACT**

**Objective** To explore views of parents of preterm babies, adults born preterm and professionals, on the linkage of real-world health and education data for research on improving future outcomes of babies born preterm.

**Design** Three-stage mixed-methods participatory design involving focus groups, a national survey and interviews. Survey participants who expressed uncertainty or negative views were sampled purposively for invitation to interview. Mixed methods were used for data analysis. **Setting and participants** All data collection was online. Participants were: focus groups—17 parents; survey—499 parents, 44 adults born preterm (total 543); interviews—6 parents, 1 adult born preterm, 3 clinicians, 2 teachers.

Results Three key themes were identified: (1) Data linkage and opt-out consent make sense for improving future outcomes. We found clear demand for better information on long-term outcomes and strong support for data linkage with opt-out consent as a means of achieving this. (2) Information requirements—what, how and when. There was support for providing information in different formats and discussing linkage near to, or following discharge from, the neonatal unit, but not sooner. (3) Looking to the future; the rights of young people. We identified a desire for individuals born preterm to be consulted in the future on the use of their data.

**Conclusion** With appropriate information provision, at the right time, parents, adults born preterm and professionals are supportive of data linkage for research, including where temporary identifiers and opt-out consent are used. Resources are being co-produced to improve communication about routine data linkage.

#### **BACKGROUND**

Survival of the most preterm babies born before 26 weeks has improved over time, <sup>1</sup> but rates of disability remain unchanged. <sup>2</sup> Cognitive impairment is the most prevalent disability and contributes to poor educational attainment. Over half of surviving extremely preterm infants require educational support; <sup>4</sup> 23% have mental health problems such as autism, attention deficit and emotional disorders. <sup>5</sup> There is high risk of rehospitalisation and mortality in infancy <sup>6</sup> and of asthma. <sup>7</sup> In later life, there is increased risk of type 2 diabetes and cardiometabolic problems. <sup>8</sup> Long-term outcomes data are needed to evaluate the impact of neonatal

#### WHAT IS ALREADY KNOWN ON THIS TOPIC

- Data on long-term outcomes are important to patients and their families, and necessary to evaluate the impact of neonatal care and intervention.
- ⇒ Linkage of routinely collected data sets is a feasible and cost-efficient method to obtain long-term outcome data.

#### WHAT THIS STUDY ADDS

- ⇒ Parents of children born preterm, adults born preterm and professionals are very supportive of data linkage with opt-out consent.
- Most stakeholders are supportive based on basic written information; a minority who are not, become supportive when additional information is provided in an accessible format.

### HOW THIS STUDY MIGHT AFFECT RESEARCH, PRACTICE OR POLICY

⇒ Information on the use of data should be provided at or after discharge from neonatal care, and consultation of children beyond school age, regarding linkage of their data, is a priority for further research.

care and interventions,9 however, tracking longterm outcomes following hospital discharge is complex and expensive, with high attrition. As such, there is paucity of population-level long-term outcomes for very preterm babies born in the UK since 2006. 10 The National Institute for Health and Care Research-funded NeoWONDER research programme<sup>11</sup> will address this by linking the UK National Neonatal Research Database<sup>12</sup> to other health and education data sets to obtain information on the later status of preterm babies born in England and Wales 2007-2020. Patient and family perspectives on outcomes following preterm birth are increasingly used to inform research priorities. Mixed-methods approaches have previously been used in a study on parental perspectives on health outcomes in preterm birth, to provide both breadth and depth of data.<sup>13</sup> Patient and parent group involvement in study design beyond outcome selection is crucial in ensuing acceptability, and maximising participation. In this study, which is part of NeoWONDER, we explored the views of parents of preterm babies, adults born preterm and



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Figure 1 Flow diagram of methods and timeline.

professionals, on the acceptability of linkage of routine data for research.

#### **METHODS**

We used a three-stage mixed-methods design involving focus groups, a national survey and interviews (figure 1). Participants were recruited through the NeoWONDER patient and parent group group. This group of approximately 600 parents of preterm babies and adults born preterm signed up to the NeoWONDER 'Get Involved' page (www.neowonder.org.uk) launched in September 2020. All data collection was conducted online due to the COVID-19 pandemic.<sup>14</sup> The principles of participatory research<sup>15</sup> and the Consolidated Criteria for Reporting Qualitative research guided our work.<sup>16</sup>

#### Focus groups and co-designed national survey

The aim of the focus groups was to co-design a national survey to seek the views of parents and adults born preterm on linking routine health and education records with neonatal records. Focus group participants were recruited through an online advert circulated to the 600 members of the patient and parent group (figure 1). Prior to the focus group meeting, they were provided with a draft version of the survey. CB (clinician), MS (qualitative researcher) and a preterm-born peer researcher facilitated three 1 hour focus groups involving 17 participants. The groups discussed survey content, format and dissemination, and ways to ensure inclusion of harder-to-reach groups. Only one participant who signed up to the focus group did not attend.

These discussions strongly influenced the survey design. In particular, questions capturing demographic data were perceived as intrusive so were minimised.

The final online survey (online supplemental file S1) was constructed using the Imperial College Qualtrics platform and piloted with three parent collaborators before being advertised on social media, and via posters in 15 neonatal units. Survey responses were de-identified prior to analysis using Microsoft Excel.

#### **Semi-structured interviews**

The aim of interviews with parents and an adult born preterm was to enable more in-depth understanding of issues of interest in the survey data. We identified information-rich cases from the survey where uncertainty or negative views were expressed, then sampled purposively from this subset ensuring maximum

possible diversity of (self-reported) demographic criteria in the sample (online supplemental file S2).<sup>17</sup> Data collected so far suggested that opposition was unusual and for a narrow range of reasons. We anticipated, therefore, that six to eight interviews would provide sufficient insights, with scope for more if data saturation was not achieved. Semi-structured interviews were conducted with diagrams offered to interviewees to explain data flow (online supplemental file S3). The interviews were conducted by CB and MS.

The aim of interviews with professionals (carried out by EvB) was to learn more about their views on data linkage. We purposively sampled for professionals who care for preterm-born children in health and education settings; some were known contacts, others recruited using snowballing techniques, were unknown<sup>18</sup>

Interviews were recorded and transcribed, with thematic analvsis conducted manually. All three investigators participated in a systematic coding process. Survey and interview findings were triangulated to identify meta-themes across them. 19 Regular team discussions considered discrepancies within and across data sets, and agreed final themes.

#### **RESULTS**

#### Survey and interview participants

Of a total of 543 survey respondents, the majority (87%) were mothers of preterm-born children. The current ages of pretermborn children of parent respondents ranged from 'currently in neonatal unit' to 32 years. Participating adults born preterm were aged 20-68 years. Parents frequently reported their preterm child as having ongoing health needs (41%) or educational needs (26%). Interview participants included one adult born preterm, four mothers and two fathers, all of whom expressed negative or uncertain views on aspects of data linkage in the survey (see table 1 and online supplemental table S4 for survey and interview participant characteristics). None declined to be interviewed. Five professionals were interviewed: a neonatologist; a disability paediatrician; a professional with digital health expertise; two teachers. One clinician had experience as a parent to a preterm child. One neonatologist and one neonatal nurse declined due to scheduling conflicts. All interviews lasted 20-30 min. Data saturation appeared to be reached following the interviews.

#### **Themes**

From the survey and interviews we identified three themes.

#### Theme 1: Data linkage and opt-out consent make sense for improving future outcomes

The overwhelming majority (>98%) of survey respondents felt better information on long-term outcomes of preterm babies to be important. This high level of support was consistent for: (1) Health, (2) Behavioural, personal, social and emotional development, and (3) Education (table 2).

Interview and free-text survey data illustrated powerfully the challenges posed by the current lack of available information.

'The difficulty in having a preterm baby is having no gauge, comparison or reference' (Survey participant—mother of 5-year-old born 24 weeks)

"... probably my biggest worry now is what's going to happen in the future and it is just an unknown which makes it much worse' (interviewee—mother of 1-year-old born 25 weeks)

#### Original research

Table 1     Characteristics of survey respondents			
Characteristic	Category	Number n=543	% (to one decimal place)
Relationship to preterm birth*	Mother to a child born preterm	474	87.3
	Father to a child born preterm	25	4.6
	Adult born preterm	44	8.1
Gestation (if	<25 weeks	59	10.9
multiple children	25-27+6 weeks	138	25.4
then youngest gestation)*	28-31+6 weeks	189	34.8
gestation	32-36+6 weeks	152	28.0
	Missing	5	0.9
Ethnicity of	Asian	11	2.0
participant*	Black	9	1.7
	Chinese	1	0.2
	Mixed	7	1.3
	White	509	93.7
	Other	3	0.6
	Missing	3	0.6
Region of majority	East	23	4.2
of neonatal care*	Midlands	73	13.4
	Northern Ireland	5	0.9
	London	76	14.0
	North-East, Yorkshire	93	17.1
	and Humber		
	North-West	36	6.6
	Scotland	28	5.2
	South-East	54	9.9
	South-West	122	22.5
	Wales	16	2.9
	Other	17	3.1
	Missing	0	0.0
Parent reports	None	270	54.1
ongoing health needs†	Yes—a few	186	37.3
	Yes—a lot	19	3.8
	Other	14	2.8
	Blank/prefer not to say	10	2.0
Parent reports	None	281	56.3
additional educational	Yes—a few	101	20.2
needs†	Yes—a lot	27	5.4
	Other	78	15.6
	Blank/prefer not to say	12	2.4
How many	1	350	70.1
preterm-born children cared for	2	139	27.9
by parent†	3	7	1.4
.,,,	4	0	0.0
	4+	3	0.6
Current age of preterm-born child	Currently in neonatal unit	11	2.2
(eldest if more	< 1 year	74	14.8
than one preterm- born child)†	1–5 years	226	45.3
	6–10 years	109	21.8
	11–15 years	45	9.0
	16–20 years	20	4.0
	> 20 years	14	2.8
Current age	20–29	18	41.0
of adult born	30–39	19	43.2
preterm (years)‡	40+	7	16.0
			Continued

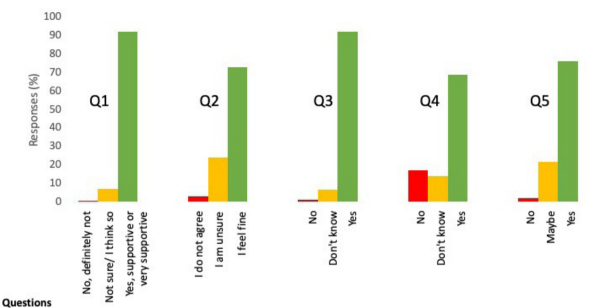
Table 1 Cont	inued		
Characteristic	Category	Number n=543	% (to one decimal place)
†Questions posed	to all respondents: N = 4 to adults born pretern	99.	

Most survey respondents (92%) supported linkage of anonymised existing records, including 'sensitive' information such as special educational needs or free school meals (figure 2). When asked for views on temporary use of identifiers for linkage purposes, such as child's name and postcode, 88% of adults born preterm remained supportive of linkage. However, support among parents fell to 74%, with the remainder either unsure and wanting more information (24%) or opposed (3%).

A majority of survey respondents were happy with data linkage for research purposes to be made routine (76%). Again,

 
 Table 2
 Survey results: questions on importance of long-term data
 and until what point linkage should be carried out

Question	Response options	Number	%
How important is having	Very important	418	80.2
better information on how preterm children develop as they grow up (ie, their long-term	Quite important	99	19.0
	Not important	1	0.2
	Not sure	3	0.6
outcomes)? (n=543)	No response	22	4.0
How important is	Very important	456	87.5
information on longer-	Quite important	60	11.5
term health of preterm babies? (n=543)	Not important	2	0.4
Dables? (II=343)	Not sure	3	0.6
	No response	22	4.0
How important	Very important	465	87.5
is information on	Quite important	52	10.0
behavioural, personal, social and emotional	Not important	1	0.2
development of preterm	Not sure	3	0.6
babies? (n=543)	No response	22	4.0
How important	Very important	432	82.9
is information on	Quite important	79	15.2
educational progress for	Not important	5	1.0
preterm babies? (n=543)	Not sure	5	1.0
	No response	22	4.0
How closely related	Very closely related	290	55.7
do you think a child's	Related	189	36.3
health and their learning/ educational progress is?	Possibly related	37	7.1
(n=543)	Not related	2	0.4
(11–343)	Not sure	3	0.6
	No response	22	4.0
Till when do you think	I don't think it is acceptable ever	7	1.5
it is acceptable to link preterm children's data? (n=471)	I agree with the principle of lifelong data linkage but would like the opportunity for my child to be consulted for their views when they are older (eg, 16 or above)	284	60.3
	Until they have completed education	41	8.7
	Until the age of 18 years as legally they will be an adult	70	14.9
	Lifelong	56	11.9
	Other	13	2.8
	No response	72	15.3



Q1 How do you feel about linking together existing records to find out what happens to these preterm children as they grow up? N = 471
Q2 How do you feel about using your child's name, date of birth, postcode, and other identifiers temporarily for linkage purposes? N = 471
Q3 Some information may be thought 'sensitive' like whether a child has special educational needs, an Education, Health and Care Plan (EHCP) or free school meals. Would you be happy for researchers to use your child's 'sensitive' information, if it was anonymous? N = 471
Q4 Would you be happy for researchers to use your child's existing electronic medical and school records, without consent, if they were always anonymous? N = 436
Q5 What do you think about making data linkage routine going forwards for preterm babies? N = 418

**Figure 2** Responses to data linkage and opt-out consent for obtaining long-term data.

the remainder were mainly unsure, with only 2% expressing negative views. Sixty-nine per cent were happy for the use of records without consent if they were always anonymous (see quote 1, table 3).

Survey responses by parents and adults born preterm were found to be similar (except where otherwise stated), with a slightly greater proportion of supportive responses from the preterm-born adult group throughout. The relatively high rates of support reflect an altruistic desire to 'give something back' and a commitment to helping individuals in the future.

'... if we can help to make life easier for any other parents going through the situation or to help with the care of premature babies then we're all up for it.' (Interviewee—mother of 7-year-old born 25 weeks)

All professionals interviewed were strongly supportive of data linkage and the use of identifiers with opt-out consent. One clinician felt that as data are necessary for an effective service, there is a moral argument that using data for research 'trumps' the rights of those who wish to use the service but have their data excluded. Another clinician argued that the study design is 'direct care, delivered retrospectively... to understand decisions already made'.

#### Theme 2: Information requirements—what, how and when

Interviews demonstrated that uncertainty or negativity in the survey generally stemmed from a lack, or misunderstanding, of information. Findings regarding the 'what', 'how' and 'when' of information provision are presented below.

Table 3	Additional quotes	
Reference number	Quote	Participant characteristics
1	"I think the opt out way is a good way of doing it because you're still (saying) 'yes', (but) people can say, "No" if they choose that they don't want their information to be used."	Interviewee—adult born preterm at 29 weeks
2	"You want to know who's got their hands on that sort of data…if it's in the right hands, in professional hands, then I'm happy."	Interviewee—father of 4-year-old twins born 27 weeks
3	"If it's as you're saying it (the data) is separated out and there's not one person that can see the whole picture then that's fine."	Interviewee—mother of 1-year-old born 26 weeks
4	"I think it probably needs to be done in a multitude of ways. So for some people they're visual learners so you know a video would explain it, but other people, they might benefit from having it written down."	Interviewee—mother of 10-year-old born 29 weeks
5	"Telling a story would be a better way."	Interviewee—father of 18-month-old twins born 26 weeks
6	"I'd be comfortable (with linkage) definitely until they're 16. But at that point I guess then you'd need to look at whether the child, as they're going into adolescence, whether they're happy to have their information shared."	Interviewee—mother of 7-year-old born 25 weeks

#### Original research

#### What

Our qualitative data demonstrated the following assurances were important to those uncertain about data linkage: (1) Only trust-worthy professional organisations would handle data, (2) Data would only be used for legitimate research purposes, (3) Data would be processed using secure, split-file methods, (4) Only routine data that already exist, and not new collections, would be linked (see quotes 2 and 3, table 3).

Explaining the following was also considered key to enabling parents to balance risk versus benefit: (1) The potential benefits of data linkage, (2) A requirement for opt-in consent will likely render data linkage unfeasible, not least because of resource implications. Professionals also thought these points were key to acceptability for parents.

#### How

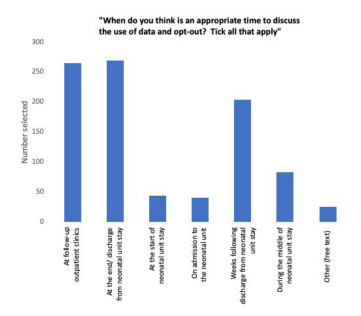
Only brief written information could be provided in the survey, while interviewees saw illustrations of data flows, including the secure split file component (online supplemental file S4), and had a discussion with an informed professional (CB). Having information in these formats led to all seven interviewees changing to full support of data linkage. One teacher also initially felt 'reluctant' and 'nervous' about using identifiers without consent. However, when the split file process was explained she no longer had these concerns.

'From what we've talked about (in the interview), my views are very different (to) what perhaps I said in the original survey.' (Interviewee—father of 18-month-old twins born 26 weeks).

There was strong support for information on data linkage, in the future, to be available in different formats (see quotes 4 and 5, table 3).

#### When

Quantitative survey results indicated that data linkage should only be discussed near to, or following discharge from the neonatal unit (figure 3). Qualitative data showed that earlier on, new parents are worried about their baby's survival and hence discussing data linkage into the future is unlikely to be appropriate.



**Figure 3** Preferred time to provide information on data linkage.

Definitely not in neonatal unit - too many emotions, concerns and unknowns while baby (is) in neonatal care (Survey free text—mother of 1-year-old born 30 weeks)

#### Theme 3: Looking to the future; the rights of young people

While the value of lifelong data linkage was acknowledged by participants, only 12% of survey respondents agreed this should be automatic (table 2). The majority of survey respondents (60%), and all interviewees, indicated that individuals born preterm should be consulted at an appropriate age on ongoing linkage of their data (see quote 6, table 3).

#### DISCUSSION

Using a mixed-methods approach to explore views of parents, adults born preterm and professionals on data linkage for research, we identified three themes. There was clear demand for better information on long-term outcomes and strong support for data linkage with opt-out consent as a method of achieving this (theme 1). The importance of the right content, timing and format of information was highlighted (theme 2) and there was a strong feeling that individuals born preterm should be consulted, in the future, about linking their data beyond school age (theme 3). Responses from parents and adults born preterm were similar.

Our findings of strong parental support for data-sharing and opt-out consent for research corroborate those from a previous study in 2011–2012.<sup>20</sup> This earlier study surveyed parents of preterm babies during admission to the neonatal unit; our study provides evidence of continued support into childhood and beyond. Both surveys identified small proportions of the target populations opposed to data sharing if identifiers were used, or explicit permission was not sought. We found that support for data linkage is likely to be maximised if differing requirements of individuals, for the content and format of information, are met. In terms of content, different levels of detail on key issues such as data security, were required. In terms of format, the availability of visual information and the opportunity for a discussion with an expert were important. Meeting these needs reduced concerns about the risks, and improved understanding of the benefits, of data linkage with use of temporary identifiers and opt-out consent.

There were two findings that it appears have not previously been explored. First, there was a clear message that data linkage discussions with parents would be insensitive during the neonatal stay, when survival is often the immediate concern. We will continue to explore ways to maximise communication following discharge, in community or hospital settings, as part of the NeoWONDER information dissemination programme. Second, our results clearly support children being consulted about linkage of their data beyond school age. These findings are important for researchers and policy makers involved in data linkage across the life course and support national initiatives facilitating dialogue and involvement from the public to build trust. <sup>21</sup>

Our work emphasises the importance of co-production of resources. Based on our findings, we have developed multiple versions of parent/patient information leaflets; a short version in an accessible Frequently Asked Question format, and a longer more detailed version. <sup>22</sup> Furthermore, we have co-produced a digital animation video to explain the complex data flows, <sup>23</sup> and have formed a young people's advisory group to co-lead future work consulting a larger group of individuals born preterm.

To our knowledge, the subject of data linkage has not been previously explored with this particular group of stakeholders. Other strengths of our work include our participatory and mixed-methods approaches which helped ensure our research design and methods were feasible and acceptable and provided breadth and depth of data, and our timely co-production of new resources. Additionally, we achieved diversity within our pool of interviewees in terms of gestation at birth, region of the UK where neonatal care was received, experience of multiple and singleton births, mothers and fathers (online supplemental file S3). However, we were only able to include one adult born preterm, and two from ethnic minority backgrounds in the interviews. This was because our sampling was limited to the small pool of participants who expressed negative or uncertain views (online supplemental file S3). Furthermore, despite efforts to include those with limited English proficiency in the study, we were unable to achieve this.

#### CONCLUSION

This mixed-methods study shows that with appropriate information, parents, adults born preterm and professionals are supportive of data linkage with opt-out consent. Resources are being co-produced with parents, adults and teenagers born preterm, and professionals to improve communication and understanding of routine data linkage.

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Patient consent for publication Not applicable.

**Ethics approval** This study involves human participants and was approved by The Yorkshire and The Humber-Leeds East Research Ethics Committee (Reference 20/YH/0330, IRAS ID 291612). Participants gave informed consent to participate in the study before taking part. The parent/patient information leaflets have been approved by the Confidential Advisory Group (CAG) and Research Ethics Approval Committees (REC) (reference 21/EM/0130, IRAS ID 293603, CAG 21/CAG/0081). These regulatory bodies have also granted permission for the study to use personal identifiers for data linkage without the need for explicit consent from parents.

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**Data availability statement** Data are available upon reasonable request.

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# Views of parents, adults born preterm and professionals on linkage of real-world data of preterm babies: Supplementary file

#### S1: Survey content

#### A) BACKGROUND DETAILS

These first quick questions will give us a bit of information about you; this helps us to know whether we have captured views from people with different experiences.

I am:

mother to a child born preterm

father to a child born preterm

an adult born preterm

other e.g. step parent, adult sibling: [free text box for further detail]

For parents only: How old is your preterm child/children (in completed years)? If more than one preterm child, please include all ages [free text]

For adults born preterm only: How old are you?

[free text]

For parents only: What gestation (weeks) was your preterm child/children born? [free text]

For adults born preterm only: What gestation were you born? [free text]

What ethnicity best describes you?

Asian

Black

Chinese

Mixed

White

Other

Which part of the country did you receive the majority of neonatal care?

London

Southwest

South East

East Midlands

West Midlands

Yorkshire and Humber

North East

North West

East

Other

Outside UK

For parents only: Does your preterm child/children have ongoing health needs? Yes, a lot

Yes, a few

No. none

Other

For parents only: Does your preterm child/children have educational needs?

Yes, a lot

Yes, a few

No, none

Other

#### B. IMPORTANCE OF LONG-TERM OUTCOMES

These next questions are to obtain your views on the general importance of long-term outcomes

How important do you think it is to have better information on how preterm children develop as they grow up (i.e. their long-term outcomes)?

Very important

Important

Quite important

Not important

Not sure

Please tell us a bit more about your answer

[Free text]

How important do you think information on longer term health is?

Very important

**Important** 

Quite important

Not important

Not sure

How important do you think information on behavioural, psychosocial and development is?

Very important

**Important** 

Quite important

Not important

Not sure

How important do you think information on educational progress is?

Very important

Important

Quite important

Not important

Not sure

How closely related do you think how healthy a child is and their learning or educational progress?

Very closely related

Related

Possibly related

Not related

Not sure

If you wish, please tell us a bit more about your answers [Free text]

#### C: DATA LINKAGE AND ANONYMISED DATA

Health records have NHS numbers, and education records have Unique Pupil numbers. As these are not the same, we would need to use personal identifiers such as name, postcode, date of birth to link records. The identifiers are only used for the linkage; once linked, the identifiers are removed and the data becomes anonymous. This means that anyone looking at the data cannot find out who the individual children are in the data set.

The next few questions ask for your views on the reality of linking a preterm child's records.

How do you feel about linking together existing records to find out what happens to these preterm children as they grow up?

Yes, very supportive Yes, supportive I think so No definitely not

Not sure

How do you feel about using your child's name, date of birth, postcode and other identifiers temporarily for linkage purposes?

I feel fine about this I am a bit unsure about this I do not agree with this

What are your concerns if any about this? [Free text]

Some information may be thought "sensitive" like whether a child has special educational needs, an Education, health and care plan (EHCP) or free school meals. Would you be happy for researchers to use your child's "sensitive information", if it was anonymous? Yes

No

Don't know

Whatever your answer, please explain why if you would like to [Free text]

Till when do you think it is acceptable to link preterm children's data? Lifelong

I agree with the principle of lifelong data linkage but would like the opportunity for my child to be consulted for their views

Until they have completed education

Until the age of 18 years as legally they will be an adult

I don't think it is acceptable ever

Other

D. USE OF ANONYMISED DATA WITHOUT CONSENT, NOTIFICATION AND OPTOUT/DISSENT

As it would be impossible and impractical to seek permission (i.e. consent) from 90,000 families to link these data, we will seek permission from the Confidential Advisory Group (CAG) to link these data without consent (assuming parents and ex-patients are supportive).

However, we would want to try our best to let parents of preterm babies and ex-patients know that this is what we were doing, and give them a chance to opt-out if they do not wish for their data to be linked.

Would you be happy for researchers to use your child's electronic medical and school records for this reason, without consent, if they were always anonymous?

Yes

No

Don't know

#### E. MAKING DATA LINKAGE ROUTINE IN THE FUTURE

The ideal situation is for data to be linked going forwards for babies born beyond 2018 as well. We want to make sure we provide information at the most appropriate time. We know that too much information can be overwhelming, especially on arrival and during your time in the neonatal unit when there is lots going on. We want to understand your views about how and when discussion about data should take place.

What do you think about making data linkage routine going forwards for preterm babies? Yes, should be done routinely

Maybe

No, I don't think this should happen

What are your concerns if any about this? [Free text]

When do you think is an appropriate time to discuss the use of data and opt-out? Tick all that apply:

On admission to the neonatal unit

At the start of neonatal unit stay

During the middle of neonatal unit stay

At the end/discharge from neonatal unit stay

Weeks following discharge from neonatal unit stay

At follow-up outpatient clinics

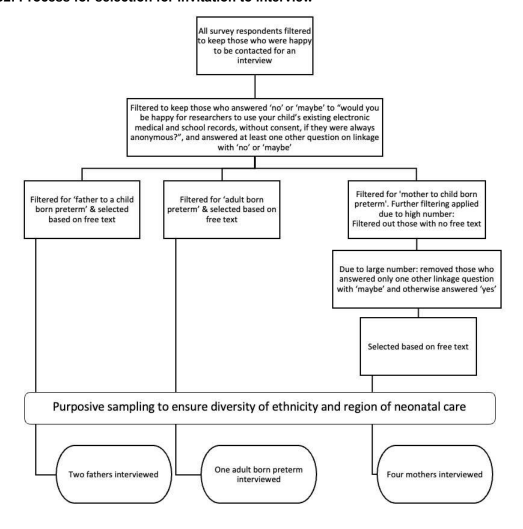
Please tell us a bit more about your answers [Free text]

We would like to interview a small number of parents to further our understanding. Is this something you would be happy doing?

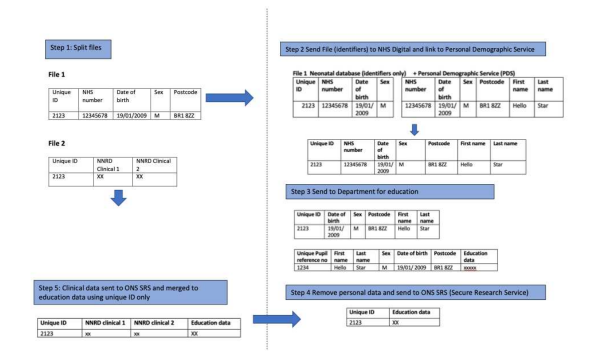
If so, please provide your email. Please be aware that if you provide your email address you potentially make your survey answers identifiable. We can assure you that no one other than the small research team will see your initial submission and we will remove your email address and save it separately to your survey as soon as we receive it.

If you would like us to inform you of the outcomes of this research or any outputs related to this research, please provide your email address.

#### S2: Process for selection for invitation to interview



#### S3: Flow diagrams used to explain data flows in interviews



## S4: Characteristics of interviewees (parents of preterm-born children and adult born preterm)

Characteristic	Category	Number N=7
Relationship to preterm	Mother to a child born preterm	4
birth	Father to a child born preterm	2
	Adult born preterm	1
Singleton or multiple birth	Singleton	6
	Multiple	1
Gestation (if multiple	25 – 27 + 6 weeks	5
children then earliest	28 – 31 + 6 weeks	1
gestation)	32 – 36 + 6 weeks	1
Ethnicity of participant	Asian	2
	White	5
Region of majority of	East	1
neonatal care	Midlands	3
	London	2
	Wales	1
Current age of preterm-born	< 1 year	2
child (eldest if more than	1 – 5 years	3
one preterm-born child) <sup>‡</sup>	6 – 10 years	1
Current age of adult born preterm (years)§	20 – 29	1

 $<sup>^{\</sup>ddagger}n = 6$