Perinatal Management of Extreme Preterm Birth before 27 weeks of gestation

A Framework for Practice - Appendices

October 2019

in collaboration with
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Appendix 1: Outcomes for extremely preterm babies

Survival/Mortality

International studies indicate incremental improvements in survival for the most premature babies over the last 1-2 decades. There is wide variation in survival estimates of live born babies (for example, from 3 – 22% at 22 weeks and 39 - 70% at 24 weeks of gestation), influenced by cohort selection, place of birth and variation in provision of active obstetric and neonatal treatment. The largest changes in outcome appear to be at the lowest gestational ages. In particular, at 22 weeks of gestation, recent cohort studies from US, Sweden and Germany indicate that approximately 30% of live born babies who receive active treatment survive to discharge.

The latest data on outcome for extremely premature babies in the UK are presented in Table 1 (source: MBRRACE-UK). Survival has increased steadily since 2006 and currently active respiratory care is offered to 88% of babies at 23 weeks and 23% of births at 22 weeks of gestation. Survival at 22 weeks of gestation is based on small numbers of babies and thus the confidence limits are wider than at other gestational weeks. It is probable that selection of babies for active treatment is biased towards those with best outlook, and so expected survival following active (survival focused) management for all infants born at 22 weeks of gestation is likely to be lower than the reported survival figures. Figure 3 displays graphically the estimated survival rates at different gestations for babies who are provided with active care in the UK compared to 3 recent publications; recent MBRRACE-UK findings are consistent with those in other settings.
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Figure 3: Estimated survival if active (survival focused) care is provided, comparing UK with recent international studies (8,9,11,12)
### Table 1: Number and percentage of births, including births where the fetus was alive at onset of labour, live births, births receiving active care, admissions for neonatal care and survival to 1 year of age for births in 2016 in the UK. Recording of active care on the MBRRACE-UK database commenced during 2016 and thus rates are inferred from recording of a total of only 292 deaths.

<table>
<thead>
<tr>
<th>Gestational Week</th>
<th>22 weeks</th>
<th>23 weeks</th>
<th>24 weeks</th>
<th>25 weeks</th>
<th>26 weeks</th>
</tr>
</thead>
<tbody>
<tr>
<td>All births</td>
<td>486</td>
<td>510</td>
<td>656</td>
<td>664</td>
<td>832</td>
</tr>
<tr>
<td>Births alive at onset of labour</td>
<td>290</td>
<td>362</td>
<td>497</td>
<td>508</td>
<td>674</td>
</tr>
<tr>
<td>Live births</td>
<td>183</td>
<td>301</td>
<td>456</td>
<td>486</td>
<td>662</td>
</tr>
<tr>
<td>% live births (of those alive at onset of labour)</td>
<td>63%</td>
<td>83%</td>
<td>92%</td>
<td>96%</td>
<td>98%</td>
</tr>
<tr>
<td></td>
<td>57 to 69</td>
<td>79 to 87</td>
<td>90 to 94</td>
<td>94 to 98</td>
<td>97 to 99</td>
</tr>
<tr>
<td>Delivery room deaths</td>
<td>155</td>
<td>78</td>
<td>26</td>
<td>19</td>
<td>16</td>
</tr>
<tr>
<td>% deaths before admission</td>
<td>85%</td>
<td>26%</td>
<td>6%</td>
<td>4%</td>
<td>2%</td>
</tr>
<tr>
<td></td>
<td>80 to 90</td>
<td>21 to 31</td>
<td>4 to 8</td>
<td>2 to 6</td>
<td>1 to 3</td>
</tr>
<tr>
<td>Live births receiving active care</td>
<td>43</td>
<td>264</td>
<td>449</td>
<td>486</td>
<td>662</td>
</tr>
<tr>
<td>% receiving active care (of all live births)</td>
<td>23%</td>
<td>88%</td>
<td>98%</td>
<td>100%</td>
<td>100%</td>
</tr>
<tr>
<td>Admitted for neonatal care</td>
<td>28</td>
<td>223</td>
<td>430</td>
<td>467</td>
<td>646</td>
</tr>
<tr>
<td>% admitted for neonatal care (of births receiving active care)</td>
<td>65%</td>
<td>85%</td>
<td>96%</td>
<td>96%</td>
<td>98%</td>
</tr>
<tr>
<td></td>
<td>51 to 79</td>
<td>81 to 89</td>
<td>94 to 98</td>
<td>94 to 98</td>
<td>97 to 99</td>
</tr>
<tr>
<td>Deaths &lt; 1 year</td>
<td>13</td>
<td>122</td>
<td>160</td>
<td>108</td>
<td>106</td>
</tr>
<tr>
<td>Survivors to 1 year</td>
<td>15</td>
<td>101</td>
<td>270</td>
<td>359</td>
<td>540</td>
</tr>
<tr>
<td>Survival</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Of those alive in labour</td>
<td>5%</td>
<td>28%</td>
<td>54%</td>
<td>71%</td>
<td>80%</td>
</tr>
<tr>
<td></td>
<td>2 to 8</td>
<td>23 to 33</td>
<td>50 to 58</td>
<td>67 to 75</td>
<td>77 to 83</td>
</tr>
</tbody>
</table>
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<table>
<thead>
<tr>
<th>Of live births receiving active care</th>
<th>35%</th>
<th>38%</th>
<th>60%</th>
<th>74%</th>
<th>82%</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>21 to 49</td>
<td>32 to 44</td>
<td>55 to 65</td>
<td>70 to 78</td>
<td>79 to 85</td>
</tr>
<tr>
<td>Of those admitted to intensive care</td>
<td>54%</td>
<td>45%</td>
<td>63%</td>
<td>77%</td>
<td>84%</td>
</tr>
<tr>
<td></td>
<td>36 to 72</td>
<td>38 to 52</td>
<td>58 to 68</td>
<td>73 to 81</td>
<td>81 to 87</td>
</tr>
</tbody>
</table>

Severe impairment

The impact of a particular impairment has ramifications for functioning in many areas and is captured in the WHO International Classification of Functioning, Disability and Health (ICF) \(^50\). Individual perception of the impact of impairment on functioning in society (i.e. disability) is highly personal and varies from family to family, dependent on their experience, knowledge and attitudes, and the support available to them. Indeed, it also varies between neonatal health professionals \(^51,52\). What for one individual or family may be an acceptable outcome may not be so for another.

The criteria for categorisation of impairments in neonatal studies also vary. In 1992, a working party sponsored by the National Perinatal Epidemiology Unit and Oxford Health Authority defined health status at 2 years into severe disability, likely to result in high levels of dependency on others with reduced chances of independent living or other or no disability. This was refined in the BAPM working group document published in 2008 \(^26\). Other authorities have used a profound category, a subgroup of severe disability, on which to base counselling \(^53\). In the absence of regularly updated national data on the prevalence of profound impairments after extremely preterm birth, the working group recommended that the well-established “severe impairment” category as defined by the BAPM Working group be used to inform parents when discussing risk following extremely preterm birth. This has been used in several large population-based studies and gives a reliable estimate of risk not available from local data, where the small number of survivors introduces significant uncertainty into estimates. Generally, for extremely preterm babies, as mortality risk decreases, the risk of severe disability among survivors also decreases.

The severe impairment category includes any of:

- severe cognitive impairment with an IQ lower than 55 (<-3 standard deviation); this will usually result in the need for educational support and require supervision in daily activities
- severe cerebral palsy – classified as Gross Motor Function Classification System (GMFCS) grade 3 or greater (see Box 2)
Estimated prevalence rates of severe impairment in four major studies are shown in Figure 4, which may be summarised as:

- **22**° - **22**° weeks: 1-in-3 survivors has severe impairment
- **23**° - **23**° weeks: 1-in-4 survivors has severe impairment
- **24**° - **25**° weeks: 1-in-7 survivors has severe impairment
- **26**° - **26**° weeks: 1-in-10 survivors has severe impairment

![Figure 4 – Prevalence of severe neurodevelopmental impairment in England (2006) compared with rates reported in recent international publications using similar classifications](image)

Note: Data from reference [55] were kindly reanalysed by the NICHD NRN to match the UK classification.

**Level III**: Children walk using a hand-held mobility device in most indoor settings. When seated, children may require a seat belt for pelvic alignment and balance. Sit-to-stand and floor-to-stand transfers require physical assistance of a person or support surface. When travelling long distances, children use some form of wheeled mobility. Children may walk up and
down stairs holding onto a railing with supervision or physical assistance. Limitations in walking may necessitate adaptations to enable participation in physical activities and sports including self-propelling a manual wheelchair or powered mobility.

**Level IV:** Children use methods of mobility that require physical assistance or powered mobility in most settings. Children require adaptive seating for trunk and pelvic control and physical assistance for most transfers. At home, children use floor mobility (roll, creep, or crawl), walk short distances with physical assistance, or use powered mobility. When positioned, children may use a body support walker at home or school. At school, outdoors and in the community, children are transported in a manual wheelchair or use powered mobility. Limitations in mobility necessitate adaptations to enable participation in physical activities and sports, including physical assistance and/or powered mobility.

**Level V:** Children are transported in a manual wheelchair in all settings. Children are limited in their ability to maintain antigravity head and trunk postures and to control arm and leg movements. Assistive technology is used to improve head alignment, seating, standing, and/or mobility but limitations are not fully compensated by equipment. Transfers require complete physical assistance of an adult. At home, children may move short distances on the floor or may be carried by an adult. Children may achieve self-mobility using powered mobility with extensive adaptations for seating and control access. Limitations in mobility necessitate adaptations to enable participation in physical activities and sports including physical assistance and using powered mobility.

**Box 2: GMFCS Grades 3-5 description:**
Appendix 2: Situations of uncertainty and potential conflict

Uncertain gestational age

If gestational age is uncertain, (i.e. no dating ultrasound scan) but thought to be $\geq 22^{+0}$ weeks, an ultrasound scan by an experienced sonographer should be carried out if time permits. If the fetal heart is heard during labour, a professional experienced in stabilisation of extremely preterm babies should attend the birth. The baby should be delivered into a plastic bag and an estimate made of gestation. Unless the baby is clearly $< 22^{+0}$ weeks of gestation, and/or estimated (or weighed) at $< 350$ g, stabilisation and supported transition with lung inflation, using an appropriately sized facemask, should begin, usually after one minute of deferred cord clamping. Subsequent management will be dictated by the clinical condition of the baby, the response to stabilisation manoeuvres and parental wishes and expectations. In this scenario, it is likely that the parents will have had little, if any, time to consider the situation and so it may be appropriate to proceed with initiating active (survival focused) neonatal management and to reassess the situation in the ensuing minutes, hours and days. It is noted that assessment of either gestation or risk of poor outcome based on condition at birth is not reliable (44).

Rapid birth without time for counselling

Preterm labour often progresses rapidly, and there may be insufficient time for detailed discussion with the parents before the baby is born. In such a scenario, a decision about management at birth will need to be made based on the available clinical information and informed by the most recent management plan, if any. When risk is unclear (for example gestation is uncertain), and particularly if there has not been time for full discussion with parents, it would usually be reasonable to embark on a provisional plan of stabilisation +/- resuscitation – providing potentially life-sustaining treatment at delivery, but redirecting to palliative care if the baby appears very immature or responds poorly to stabilisation (for example remains severely bradycardic despite intubation and intermittent positive pressure ventilation).

Baby born in unexpectedly good condition

In the rare circumstance where palliative (comfort focused) care has been agreed, but a baby is born in unexpectedly good condition, attending midwifery and/paediatric medical staff should discuss with parents whether the estimated gestation and prognosis were accurate and whether the planned palliative approach is still appropriate. Stabilisation should not be delayed if deemed in the baby’s best interests.

Baby born in unexpectedly poor condition

Supplementary material

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When active neonatal management has been agreed, but the baby is born in unexpectedly poor condition, it is the responsibility of the most senior attending neonatal professional to decide if ongoing attempts at stabilisation and/or resuscitation are in the baby's best interests. This should be conveyed sympathetically but unambiguously to parents, and palliative care offered.

**Parents request a second opinion**

If the parents wish, they should have the opportunity to discuss outcomes with a second senior member of the perinatal team. When parents do not agree with the perinatal team, recently published RCPCH guidance around dealing with conflict may be helpful (57).

**Threatened birth before 22+0 weeks of gestation**

Where gestational age is certain and is below 22+0 weeks, it would be considered in the best interests of the baby, and standard practice, not to offer neonatal intensive care. If it is possible that the birth may be delayed to a point where active care of the baby would be planned, transfer of the mother to a maternity unit adjacent to a neonatal intensive care unit should be considered.
Appendix 3: Communication: Guidance for professionals consulting with families at risk of extreme preterm delivery.

This Appendix is designed for use by all staff caring for families at risk of extreme preterm birth, to facilitate the sharing of consistent and accurate information. It should be used to support conversations about decision making with parents, in conjunction with written information such as the suggested Template Parental Information - Helping parents to understand extreme preterm birth (Appendix 4). Written information should never be used as a stand-alone information sharing tool.

In order properly to involve parents as equal partners in care and decision-making for their babies, all parents facing potential extreme preterm birth need to understand the risks associated with their baby’s birth, and possible treatment options.

In the context of the different risk scenarios outlined in this Framework, health professionals need to consider how their approach to consulting with parents may differ, depending on the individual circumstances:

- **In extremely high risk** cases, parents should be provided with relevant information about the risk to their baby, and the recommendation that it would be best for their baby to provide palliative (comfort focused) obstetric and neonatal management. Parents should be told about the role they can play in caring for their baby and memory making after birth, and should be fully involved in decisions about how and where palliative management takes place.

- **In high risk** cases, the role of the consultation is critical in supporting parents together with professionals to decide on the right pathway for their baby. Parents should be provided with as much information as is available and should have as much time as possible, ideally over a number of discussions, to work through the different options available in order to agree with professionals what the right option is for their family.

- **In moderate risk** cases, parents should be provided with relevant information about the risk to their baby, and the recommendation that it will be best for their baby to provide active management both antenatally and after birth. Parents should be told about likely interventions and what may happen next, and be fully involved in decisions about how active treatment is managed.

Thus, consultation is most critical where delivery would be associated with a high risk of unacceptably poor outcome (see main text). In such situations, parents need support to make an informed choice about the provision of either active or palliative management; such situations demand the greatest care and sensitivity. Consultation should not be directive, but professionals...
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should seek to determine when gentle guidance around what is likely to be in the baby’s best interests would be helpful for the family.

When is the right time?

Evidence suggests that parents find that, where time allows, consultation is most useful at the earliest opportunity, both to allow time for information processing, discussion, and decision making, and to minimise the effects of labour and medications on cognition. Follow up consultations allowing ongoing dialogue are highly valued by families\(^{(58,59)}\) and should be offered at any point, acknowledging the challenging nature of the information that parents are being asked to receive, the time this may take to process and the decisions that need to be made.

Who should be involved?

Consultation with parents should ideally be provided by the most experienced members of the perinatal team involved in care of the mother and her baby. Continuity of care is essential and, whenever possible, consultation should be delivered as a joint obstetric, neonatal and midwifery approach, ensuring transparency and consistent, clear communication. The presence of members of the multidisciplinary team (particularly nurses and midwives) during such conversations is highly valued by families, and may provide opportunity for clarification and ongoing conversation outside the formality of such settings\(^{(59,60)}\). Parents may also find the advice and support of their family, friends, spiritual advisers and/or voluntary organisations to be of great value at this time.

Structuring the Consultation

1. **Exploring the parents’ prior knowledge and understanding** can be a useful way to open the consultation. Establishing parents’ own understanding about the risks of their situation, their prior experience and knowledge, as well as their expectations of the conversation is important, both to generate trust and to ensure that the consultation meets their individual needs. Parents’ hopes, priorities and expectations of the care that they and their baby will receive should be explored with sensitivity, honesty and compassion in a realistic way.

2. **Balanced Information** - Studies suggest that conveying solely negative information to parents is not well received. Providing balanced information with honesty seems to be most useful to parents. Respecting parents’ perspectives and the importance of hope, even in the most difficult of situations, is highly valued. Exploring parental hopes, wishes and fears in each scenario can help to do this, and to build trust and rapport.
with the clinical team. Where survival is not possible, or is extremely unlikely, parental hopes relating to spending time with their baby, involving family members, and memory making should be explored.

3. **Conveying Risk** - Categorisation of risk to the baby of death or survival (with or without impairment) in a given scenario should be conveyed sympathetically and with clarity. Parents may find it useful to see this displayed graphically (see Appendix 4). Gestation-based risk should be explained within the context of other risk modifiers (such as birth weight, gender, multiplicity, etc.). It is important to convey information accurately, in the appropriate context. While the most relevant statistic for parents is usually the chance of survival if active stabilisation and neonatal intensive care is attempted, parents should be helped to understand that not all babies survive labour, and so outcome data depend upon the stage at which parents are being counselled. Outcome data are, of course, also highly influenced by intention to treat at delivery and it is likely that current published outcomes are skewed towards those fetuses/babies in the best condition at birth. Not all parents find percentage figures easy to understand. It can be helpful to explain in terms of odds e.g. 1 in 4, or 1 in 10.

To avoid framing bias, we suggest interpreting risk neutrally. For example, “Given what we know about the situation for your baby, there is a 30% chance of your baby surviving. This means that for every 10 babies treated actively (with intensive care) in situations like this, three would survive while sadly seven would not”.

4. **Discussing Poor Outcomes** - There is not a simple definition of a ‘poor’ outcome – the interpretation of this is likely to vary greatly between clinicians, parents, and families. Published data generally refer to scoring systems and classification of motor and cognitive dysfunction, but also often include children with profound vision or hearing loss. Some of these terms may not be meaningful to families, and families’ views may differ on the outcome that they would regard as unacceptably poor. Therefore, discussions should always include exploration of the parents’ views and values relating to an acceptable outcome.

Conveying the concept of severe disability in childhood, and the possible implications for future quality of life, is difficult. Some helpful phrases may include:

- Not being able to walk or move independently
- The possibility of being unable to speak
- Difficulties with swallowing or feeding safely
- The possibility of not being able to understand the world around them in a meaningful way
• Not being able to see or hear properly
• Not being able to live independently
• Having a lot of health care needs with frequent visits to hospital
• Needing extra educational support (or needing to attend a special school)

It is important also to highlight the potential for longer term health issues, including chronic lung disease and consequences of necrotising enterocolitis as well as milder neurodisability, behavioural problems and issues with educational achievement.

5. **Discussing Palliative Care** – Where appropriate, the practicalities of commencing, withholding and withdrawing intensive care and the positive role of palliative care should be described to the parents. This will help prepare them for possible outcomes after the birth. It can be useful to speak about memory making, exploring parents’ hopes and wishes. We suggest referring to guidance from Together for Short Lives and the National Bereavement Care Pathway.


6. **Decision making** – A shared decision making process is vital, especially in situations of moderate to high risk of unacceptably poor outcome. Support and guidance should be tailored to the needs of each family. Parents should be helped to understand that, even taking all available information into account, babies may be born in unexpectedly poor or unexpectedly good condition, and that this may impact upon what care at birth would be best for their baby.

7. **Parental involvement in care** - Evidence suggests parents find it very useful to hear how they can be involved in care for their baby. “Family-centred care means supporting parents to be involved in their baby’s care”[^61][^62] – this should start before birth.

Where it is planned to offer active care to the baby, and time allows, parents should be given an opportunity to visit the neonatal unit and to meet staff, and should receive information and support regarding expressing breast milk and the other ways that they could be involved in the hands-on care of their baby if s/he is admitted to the neonatal unit.
8. **Documentation and follow up** - Communication and agreed plans should be documented in full in the clinical record and plans revised regularly if pregnancy continues and/or depending upon the condition of the baby at birth and in the early days after birth. If *in utero* transfer is undertaken, the content and results of previous conversations should be clearly communicated (verbally, and in writing) with the receiving centre. Evidence suggests that parents find it very useful to receive supplemental information such as written information, visual aids and links to other resources. We include a suggested template for this information below.
Appendix 4: Helping parents to understand extreme preterm birth.

Who is this information for?

You have been given this information because your healthcare team think that you may have your baby extremely early (prematurely). You and your family need to know what is likely to happen for you and your baby if this occurs. The maternity team and neonatal (specialist baby doctors and nurses) team will talk to you about this in detail as well as giving you this information and you will have the opportunity to ask any questions that you wish.

What does this mean?

A pregnancy usually lasts for about 40 weeks. How many weeks you are along in your pregnancy (gestation) is usually worked out from an ultrasound scan at around 12 weeks (your dating scan).

Babies born before 22 weeks are so small and fragile that they do not survive. Their lungs and other organs are not ready for them to live outside the womb. Such tiny babies may show signs of life for a short time after birth but even with the very best neonatal care they cannot survive for more than a few minutes or hours.

Babies born from 22 weeks sometimes are not strong enough to survive labour and either vaginal (normal) or caesarean birth. If they are born alive, they may be able to survive if they receive intensive medical treatment. However, some extremely premature babies sadly die despite this treatment. The earlier the baby is born, the less likely it is that they will be able to survive.

Babies who are born extremely early are also at increased risk of problems with health and development as they grow up. These risks get higher the earlier (more prematurely) a baby is born, and are especially common in those children born before 25 weeks of gestation. Health problems may include breathing difficulties, gut problems (including difficulties with feeding) and eye problems. Developmental problems may include problems with movement, learning and behaviour that can range from mild to very severe; such problems are described on the following page.

The doctors and midwives will talk to you about what they expect for your baby. In some situations, there are difficult decisions to be made about how to care for your baby before and after birth. The right thing to do can be different for different families. That is why it is important that you are fully informed and feel able to let the doctors and midwives know your wishes for your baby.
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Outcome for babies born alive between 22 & 26 weeks’ gestation

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**Survival**

- **22 weeks**
  - 7 in 10 babies die (51 to 79%)
  - 3 in 10 babies survive

- **23 weeks**
  - 6 in 10 babies die (56 to 68%)
  - 4 in 10 babies survive

- **24 weeks**
  - 4 in 10 babies die (35 to 45%)
  - 6 in 10 babies survive

- **25 weeks**
  - 3 in 10 babies die (22 to 30%)
  - 7 in 10 babies survive

- **26 weeks**
  - 2 in 10 babies die (15 to 21%)
  - 8 in 10 babies survive

**Severe disability**

- **22 weeks**
  - 1 in 3 babies has severe disability (24 to 43%)
  - 2 in 3 do not**

- **23 weeks**
  - 1 in 4 babies has severe disability (16 to 33%)
  - 3 in 4 do not**

- **24 weeks**
  - 1 in 7 babies has severe disability (11 to 24%)
  - 6 in 7 do not**

- **25 weeks**
  - 1 in 7 babies has severe disability (10 to 21%)
  - 6 in 7 do not**

- **26 weeks**
  - 1 in 10 babies has severe disability (6 to 14%)
  - 9 in 10 do not**

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The survival percentages are for babies who are born alive and receive active stabilisation.

*Some babies born this prematurely cannot survive labour and birth

* The lower and upper figures indicate how certain we are of the true survival rate.

** Up to a quarter of children without severe disability may nonetheless have milder forms of disability such as learning difficulty, mild cerebral palsy or behavioural problems.
‘Outcome’

These pictures are based on what we know about the small number of babies born extremely prematurely in the UK. They show how many babies survive out of every 10 babies born alive this early, and of those who do survive, how many are likely to have a ‘severe disability’ when they grow up.

The majority of babies grow up without severe disability. A proportion of these children will develop other problems as they grow up which may mean, for example, that they need extra help in school or have problems with walking or moving around. Some may have social and emotional problems. The frequency with which children have these problems is greatest the earlier they are born, and problems are most common in children born at 22 to 24 weeks of gestation.

The chance for your baby depends on a number of different things. As well as how early they are born, it also matters how much your baby weighs when it is born, whether it is a boy or girl, whether it is a multiple birth and also how well you and your baby are around the time of birth.

What does ‘severe disability’ mean?

Disability can mean different things to different people. When talking about babies who have been born extremely prematurely, the term severe disability includes problems such as:

- Not being able to walk or even get around independently (this includes conditions such as severe cerebral palsy)
- Being unable to talk, or see or hear properly
- Difficulties with swallowing or feeding safely
- Having multiple health problems with frequent visits to hospital
- Needing to attend separate school for children with special educational needs
- Being unable to care for themselves or live independently as they grow up

What does this mean for your baby?

We don’t know exactly the future for your baby. Every baby is different and it is important to talk with your doctors and midwife. They will give you specific information about your own and your baby’s condition.

What can parents do?

What is right for your baby and your family is very individual to you. Your doctors will talk with you about your situation and
seek to understand what is important for you and your family. They will support and guide you and involve you in making
decisions about treatment for your baby. Thinking about your hopes, your wishes, and your fears about your baby can help the
team to support you in the best way possible.

What may happen with my baby?

**Stillbirth:** Some babies who are born this early do not survive labour and delivery. If this happens your baby will be given to you
to hold for as long as you would like. You will have the opportunity to spend as much time with them as you would like and to make memories with them. Under UK law only babies born after 24 completed weeks of gestation can be registered as stillborn.

**Neonatal Intensive Care:** You and the team may decide that starting neonatal intensive care would be best for your baby. This
will mean you will need some extra treatments before your baby is born. You will be given steroids to help the baby’s lungs and
brain and magnesium which also helps to protect your baby’s brain. You may need to be transferred to a specialist centre, ideally
before you have your baby, but there may not be time to do this safely. The team will also talk to you about the treatment that
will be given to your baby immediately after birth and what may happen next depending on how your baby reacts to this
treatment.

If you and the team decide that intensive care is best for your baby, you should be offered the opportunity to be shown around
the neonatal unit (if there is time for this) as it may help to see the neonatal unit and meet the people that work there before
your baby is born. You can also talk to staff about expressing breast milk, as this makes such a big difference for premature
babies.

**Comfort Care:** You and the team may decide that it will be best to provide comfort care to your baby, either because there is an
extremely high risk that your baby will not survive or he/she is likely to suffer from life-long disability even with the very best
treatment. Comfort care is also known as palliative care and is special care for babies whose time is precious but short. It means
providing treatments that will make their time as comfortable as possible. We will help you to be part of this care if you would
like. Holding your baby close to you and talking to your baby may be very comforting.

More information about comfort care or ‘palliative care’ for babies is available from [Together for Short Lives](https://togetherforshortlives.org.uk).

**What if my baby doesn’t come now?**

If your baby does not come in the next few days their chances may improve. Ideally, they will stay in the womb for as long as
possible (depending on the health of you and your baby).
If that happens there may be different options for you and your baby around the time of birth. That will depend on when your baby comes and on other things that affect the baby’s chances of responding to treatment. If this is the case, your healthcare team will continue the conversation with you about what has changed and what different options may be available depending on when your baby is likely to be born, and you will be able to discuss and revise your agreed plans accordingly.

**What might my baby look like?**

Babies born this early can weigh less than half a kilogram (1 small packet of sugar) and can look quite different to how we imagine a newborn baby. Their skin is shiny and thin and covered with fine hair. Sometimes babies can be quite bruised from the birth. If the baby has died before being born, they will usually be still. Occasionally, where babies have died very close to being born, they may make brief reflex movements that disappear very quickly.

If your baby is born alive, they may take a breath and make a small cry or they may not breathe. Their eyes may not be able to open yet. The baby’s colour is often purple or blue to start with.

**Transfer to a different hospital**

When you have decided with the obstetric and neonatal care teams that starting neonatal intensive care would be best for your baby, research shows that for babies born before 27 weeks of gestation it is best, whenever possible, to be born in a specialist maternity unit with a specialist Neonatal Intensive Care Unit (sometimes called a ‘Level 3 NICU’). If a baby born before 27 weeks of gestation is born in a maternity unit (or at home) where there is not a specialist NICU, then we know that the baby will generally do better if moved to a specialist NICU after birth.

If your hospital does not have a specialist NICU, this may mean that you will be offered transfer to one of these centres before your baby is born. We understand that this can be a very anxious time and that you may be moved quite some distance from home. It can be very difficult to predict which mothers will deliver early and so some mothers may be moved to another hospital and their baby not born early.

It may also be the case that you are considered too unwell or too far on in labour to be safely moved to another hospital before your baby is born. When it is not possible to transfer you before the baby has been born your baby may be transferred by a specialist Neonatal Transport Team after the birth. Your own health needs may mean you will be unable to travel immediately with your baby but your local maternity team will do everything they can to move you to the same unit as your baby as soon as it
is safe to do so.

We appreciate that moving to another hospital can be distressing for you and your family, especially if you are separated from your baby for a while. We will talk to you about this in more detail if it is decided that this is the best option for your family.

**What if I have more questions?**

This information has been provided to you as part of the conversation that your healthcare team will have with you about your baby. If you have any other questions do make sure you ask your doctors and nurses to answer them, so you have all the information you need about your situation and the options available to you. Your healthcare team want to work with you make the best decision for your baby and for your family.

This space is for the healthcare team who are discussing this with you to write extra details about your baby or babies.

You may want to use this space to write down some questions to discuss with the team.

Many families find it useful to have follow-up discussions, so please ask to speak to the neonatal and maternity team again at any point.
Useful contact details:

**Bliss** - Premature and sick baby charity

http://www.bliss.org.uk/

**Together for Short Lives** - Charity for babies and children with life-limiting conditions

https://www.togetherforshortlives.org.uk/

Helpline: 0808 8088 100

**Sands** - Stillbirth and neonatal death charity

https://www.uk-sands.org/

Helpline: 0808 1643332

Email helpline@sands.org.uk
Appendix 5: Example scenarios

Case 1.

A mother has been admitted to a local maternity unit in preterm labour at $24^{1/3}$ weeks of gestation. The singleton male fetus is very small with an estimated weight of 450 grams. The mother has not yet received antenatal steroids. The on-call paediatric team is asked to provide counselling and attend the birth. As per the Framework, the first step is to assess the risk for the baby if delivery occurs. At a gestation of $24^{1/3}$ weeks, the average survival rate for liveborn babies in the UK (if active treatment is provided) would be approximately 60%, with a 1 in 7 risk of severe impairment among survivors. However, in this case, the very low birth weight for the gestational age in a male fetus increases the risk. It is difficult to quantify this risk, but the baby’s prognosis is worse than average for 24 weeks of gestation, and within the “high” risk category. Given the risk for the infant, counselling should, if possible, be provided by an experienced senior trainee or consultant neonatologist in conjunction with the obstetric team. If possible, this counselling should take place after the local team has discussed the case with the nearest NICU. It would be appropriate to provide active obstetric and neonatal management if that were desired by the parents. However, it would also be appropriate to provide palliative care, if that was felt by the parents to be in the baby’s best interests.

In this case, the parents decide after consultation that they wish the baby to receive palliative (comfort focused) care. Labour progresses and a live-born baby is delivered weighing 460 grams. He is bruised and floppy with a heart rate of 50 beats per minute. The paediatric team attend to support provision of palliative care. The baby is wrapped and given to his parents to hold. He dies at approximately 30 minutes of age.

Case 2.

A mother presents to her local maternity unit (SCBU) at $22^{40}$ weeks of gestation with bulging membranes and active preterm labour. No antenatal steroids have been given, and the estimated weight of the male fetus is 510g. There is an extremely high risk of poor outcome for this fetus if delivery occurs within a short period of time. The mother and her partner are informed of the likely outcome, and advised that active (survival focused) management is not considered to be appropriate. The mother receives palliative obstetric management, and the infant is stillborn.

Case 3.
A mother presents in preterm labour to a level 2 centre at 22\textsuperscript{+3} weeks of gestation. The fetus was conceived by IVF and gestation is certain. The fetus is female and has an estimated fetal weight of 480 grams. The mother has not yet received antenatal steroids. The paediatric team is asked to provide counselling and attend the birth. Risk assessment in this case indicates that if birth occurs imminently, there would be an extremely high risk of the baby dying or of surviving with severe impairment. If labour progresses, it would be usual to provide palliative care at birth. The neonatal team may attend the birth to provide support for palliative care, but not to provide resuscitation. However, there are potentially modifiable risk factors in this case. In the absence of evidence of chorioamnionitis, it may be possible to delay preterm birth with tocolysis and so, if the parents desire, an active approach to management, antenatal transfer and corticosteroids could be provided.

Two days later, at 22\textsuperscript{+5} weeks, following transfer to a maternity unit co-located with NICU and administration of steroids, labour progresses. Given the advance in gestation, availability of specialised neonatal intensive care, and anticipated effect of corticosteroids, the baby’s risk is now judged to fall in the “high” category. Accordingly, after further consultation with parents, it would be appropriate to provide active management if this is what parents wish.

In this case, the parents decide that they wish the baby to receive active neonatal care, magnesium sulphate is given, and a live baby is born weighing 490 grams. The baby is intubated, receives surfactant and is transferred to neonatal intensive care.

**Case 4.**

A mother presents to her local maternity unit at 25\textsuperscript{+2} weeks of gestation in early labour. She has a well grown female fetus. She is offered, and accepts, antenatal steroids, but declines transfer to the nearest NICU (1 hour’s journey away) where a cot is available. Her partner asks the obstetric team what would be best for the baby, and it is clear to the midwife that parents disagree about transfer.

**Recommended action:** as part of the consultation with parents, the reasons why transfer is being recommended should be clearly explained. Data show that for the most preterm babies, prognosis (both survival and neurodevelopmental outcome) is better if they are delivered in a maternity unit adjacent to a NICU. It can be difficult to predict preterm labour, and so early transfer is preferred. It would be important to explore reasons why the mother does not wish to be transferred and to address all of her concerns. If the mother still refuses transfer she cannot be moved, but she should be offered magnesium sulphate and counselled that the baby will be moved after delivery. In this instance, the best interests of the child would be served by early care in a NICU. It would be prudent to alert both the NICU and the local transport team, as well as to think about who will be...
available to stabilise the baby after birth. The mother should also be informed that, while every effort will be made to move her
to a maternity facility adjacent to the NICU, her transfer may need to be delayed if she is unwell after delivery.

Case 5.

A woman is transferred to a level 3 centre at 23+6 weeks of gestation in preterm labour following premature rupture of
membranes two days earlier. She had received steroids prior to transfer. The female fetus appears well grown and there are no
signs of fetal compromise. After discussion with the neonatal team, and being informed about the outcomes of preterm delivery,
the mother expresses that she is very concerned about the possibility of the baby surviving with severe disability. She requests
no active obstetric management, and palliative care of the baby at delivery.

The neonatal team advises the mother that, taking all factors into account, her baby would have a moderate risk of dying or of
severe disability. It would be usual to provide active management of the baby in this situation, with the knowledge that if
complications develop in the neonatal intensive care unit, there would be the option of later withdrawal of life-prolonging
treatment. The mother agrees to this plan and the baby is born a few hours later, receives stabilisation in the delivery room and
is transferred to the neonatal intensive care unit.
Appendix References

NB references 1-49 are contained in the main paper


Perinatal management of extreme preterm birth before 27 weeks of gestation
A BAPM Framework for Practice - Appendices

61. https://www.bliss.org.uk/health-professionals/bliss-baby-charter