Caesarean section

National Collaborating Centre for Women’s and Children’s Health

Commissioned by the National Institute for Health and Clinical Excellence

Draft following stakeholder consultation – September 2011
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1 Guideline summary

1.1 Guideline development group membership, NCC-WCH staff and acknowledgements

Original (2004) version

<table>
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<th>Name</th>
<th>Role</th>
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<tbody>
<tr>
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<td>General Practitioner and Group Leader</td>
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<td>Consumer</td>
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<td>Consumer</td>
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<td>Midwife</td>
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<td>Obstetrician</td>
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<td>Neonatologist</td>
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Updated (2011) version

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1.2 Foreword

This guidance is a partial update of NICE clinical guideline 13 (published April 2004) and will replace it.

New and updated recommendations have been included on the diagnosis and management of morbidly adherent placenta; the care of women with HIV; the appropriate decision-to-delivery interval for unplanned CS; the timing of antibiotic prophylaxis provision; the risks and benefits of CS and vaginal birth; the risks and benefits of vaginal birth following a previous CS; and the appropriate care pathway for women requesting a CS in the absence of an obstetric or medical indication.

Where recommendations are shaded in grey and end [2004] the evidence has not been updated since the original guideline. Yellow shading in these recommendations indicates where wording changes have been made for the purposes of clarification only.

You are invited to comment on the new and updated recommendations in this guideline only. These are marked as [2011] if the evidence has been reviewed but no change has been made to the recommendation or [new 2011] if the evidence has been reviewed and the recommendation has been added or updated.

Appendix J contains recommendations from the 2004 guideline that NICE proposes deleting in the 2011 update. This is because the evidence has been reviewed and the recommendation has been updated or because NICE has updated other relevant guidance and has replaced the original recommendations. Where there are replacement recommendations, details are provided. Where there is no replacement recommendation, an explanation for the proposed deletion is given. You are invited to comment on the deleted recommendations as part of the consultation on the 2011 update.

A grey bar down the side of the page indicates those sections of the guideline which are new or have been updated. Material from the original guideline which has been deleted can be found in appendix I.

The original NICE guideline and supporting documents are available from www.nice.org.uk/guidance/CG13
1.3 Algorithm

Planning a CS

Making the decision for CS

Offer pregnant women evidence-based information and support. This information should include indications for CS, what the procedure involves, associated risks and benefits, and implications for future pregnancies and birth after CS. Take into account the woman’s circumstances, concerns and priorities. Acknowledge the information available in tables 4.5 & 4.6 (see recommendations 1, 2 & 4)

Communication and information should be provided in a form that is accessible. It should take into account the needs of minority communities and women whose first language is not English or who cannot read, together with the needs of women with disabilities or learning difficulties (see recommendation 3)

Consent for CS should be requested after providing pregnant women with evidence-based information (see recommendation 6)

A pregnant woman is entitled to decline the offer of treatment such as CS, even when the treatment would benefit her or her baby’s health (see recommendation 7)

Give women with HIV information about the risks and benefits of HIV treatment options and mode of birth (see recommendation 24)

When a decision is made to perform a CS, record all the factors that influence the decision, and which of these is the most influential (see recommendation 8)

<table>
<thead>
<tr>
<th>Do not routinely offer planned CS to women with:</th>
<th>Offer planned CS to women with:</th>
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<tr>
<td>An uncomplicated twin pregnancy at term where the first twin is cephalic (see recommendation 11)</td>
<td>A singleton breech presentation at term, for whom external cephalic version is contraindicated or has been unsuccessful (see recommendation 10)</td>
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<td>A placenta that partly or completely covers the internal cervical os (minor or major placenta praevia) (see recommendation 15)</td>
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<td>HIV receiving HAART therapy with a viral load less than 400 copies per ml (see recommendation 25)</td>
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<td>HIV receiving any retroviral therapy with a viral load less than 50 copies per ml (see recommendation 25)</td>
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<td>HIV with hepatitis C virus (see recommendation 31)</td>
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<td>Hepatitis C virus (see recommendation 30)</td>
<td>Primary genital herpes simplex virus (HSV) infection occurring in the third trimester of pregnancy (see recommendation 32)</td>
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<tr>
<td>Recurrent genital herpes at term (see recommendation 33)</td>
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<td>A BMI of over 50 (and no other risk factors) (see recommendation 5)</td>
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Consider either a vaginal birth or CS for women on anti-retroviral therapy (ART) with a viral load 50-400 copies/ml (see recommendation 26)

Planning place of birth

Inform healthy pregnant women with anticipated uncomplicated pregnancies that:
- Planning a home birth reduces the likelihood of CS
- Birth in a ‘midwifery-led’ unit does not reduce the likelihood of CS (see recommendations 40 & 41)

Predicting ‘failure to progress’

Do not use pelvimetry in decision making about mode of birth (see recommendation 22)

Do not use shoe size, maternal height or estimations of fetal size to predict ‘failure to progress’ during labour (see recommendation 23)

Caesarean section: full guideline DRAFT (September 2011)
May be reduced after a planned CS | May be reduced after planned vaginal birth | No difference | Studies show conflicting findings
--- | --- | --- | ---
Perineal and abdominal pain during birth | Perineal and abdominal pain 4 months postpartum | Perineal and abdominal pain | Maternal death
Perineal and abdominal pain 3 days postpartum | Injury to bladder/ureter | Injury to bladder/ureter | Deep vein thrombosis
Injury to vagina | Injury to cervix | Injury to cervix | Blood transfusion
Early postpartum haemorrhage | Iatrogenic surgical injury | Iatrogenic surgical injury | Postpartum haemorrhage
Obstetric shock | Pulmonary embolism | Pulmonary embolism | Hysterectomy
Length of hospital stay | Wound infection | Wound infection | Anaesthetic complications
Hysterectomy due to postpartum haemorrhage | Intraoperative trauma | Intraoperative trauma | Neonatal mortality
Cardiac arrest | Assisted ventilation or intubation | Assisted ventilation or intubation | 5 min Apgar score < 7
NICU admission | Acute renal failure | Acute renal failure |

(See Tables 4.5 and 4.6 for further details)

### Diagnosing morbidly adherent placenta

After confirmation of low lying placenta at 32-34 weeks in women who have had a previous CS, use colour – flow Doppler ultrasound to test for morbidly adherent placenta (see recommendation 16).

If the ultrasound scan suggests morbidly adherent placenta:

- Discuss the improved accuracy of MRI in addition to ultrasound to help diagnose morbidly adherent placenta and clarify the degree of invasion.
- Explain what to expect during the procedure.
- Inform the woman that experience suggests that MRI is safe but there is a lack of evidence about any long term risks to the baby.
- Offer MRI if acceptable to the woman (see recommendation 17).

Discuss the interventions available for delivery including cross-matching of blood and planned CS with a consultant obstetrician present (see recommendation 18).

### CS for morbidly adherent placenta

Ensure that:

- a consultant obstetrician and anaesthetist are present.
- an experienced paediatrician is present.
- a senior haematologist is on notice that they may need to be available for advice.
- there is access to a critical care bed.
- there is sufficient cross matched blood and blood products readily available (see recommendation 19).

When performing a CS for women suspected to have morbidly adherent placenta, determine which other healthcare professionals need to be consulted or present (see recommendation 20).

### Timing of planned CS

Do not routinely carry out planned CS before 39 weeks (see recommendation 52).

### Maternal request

Explore, discuss, and record specific reasons for the request (see recommendation 34).

If a woman requests a CS when there is no other indication, discuss the overall benefits and risks of CS and vaginal birth and record that the discussion has taken place. Facilitate a discussion with other members of the obstetric team if necessary, to ensure the woman has accurate information (see recommendation 35).

For women requesting a CS because of anxiety about childbirth, offer referral to a healthcare professional with expertise in providing perinatal mental health support (see recommendation 36).

The healthcare professional providing perinatal mental health support should have access to the planned place of birth during the antenatal period (see recommendation 37).

For all women requesting a CS, if after discussion and offer of support, a vaginal birth is still not an acceptable offer a planned CS (see recommendation 38).

An obstetrician can decline a woman’s request for a CS. In this instance they should refer the woman to an NHS obstetrician in the same unit who will carry out the CS (see recommendation 39).
Pregnancy and childbirth after CS

When advising about mode of birth after a previous CS, take into account maternal preferences and priorities, the risks and benefits of repeat CS and the risks and benefits of planned vaginal birth after CS including the risk of unplanned CS (see recommendation 119).

Offer women planning a vaginal birth following a CS electronic fetal monitoring during labour and care during labour in a unit where there is immediate access to CS and on-site blood transfusion services (see recommendation 121).

During induction of labour, women who have had a previous CS should be monitored closely, with access to electronic fetal monitoring and with immediate access to CS, because they are at increased risk of uterine rupture (see recommendation 122).

Inform women with up to and including 4 CS that their risk of fever, bladder injuries and surgical injuries does not vary with planned mode of birth, and that uterine rupture is very rare (see recommendation 120).

Inform women with both previous CS and a previous vaginal birth that they have an increased likelihood of achieving a vaginal birth than women with a previous CS but no previous vaginal birth (see recommendation 123).

No influence on likelihood of CS

- walking in labour
- non-supine position during the second stage of labour
- immersion in water during labour
- epidural analgesia during labour
- the use of raspberry leaves
- active management of labour or early amniotomy to augment the progress of labour (see recommendations 47 & 49)

Interventions which may reduce the rate of CS

Involve consultant obstetricians in the decision making for CS (see recommendation 45).

Offer external cephalic version if breech at 36 weeks (exceptions include women in labour, women with a uterine scar or abnormality, fetal compromise, ruptured membranes, vaginal bleeding or medical conditions) (see recommendation 9).

Facilitate continuous support during labour from women with or without prior training (see recommendation 42).

Offer induction of labour beyond 41 weeks (see recommendation 43).

Perform fetal blood sampling before CS for abnormal cardiotocograph in labour if it is technically possible and there are no contraindications (see recommendation 46).

Use a partogram with a 4-hour action line for women in spontaneous labour with an uncomplicated singleton pregnancy at term (see recommendation 44).

Decision to delivery interval

Classification of urgency

Determine the urgency of CS using the following standardised scheme:

1. immediate threat to the life of the woman or fetus
2. maternal or fetal compromise which is not immediately life-threatening
3. no maternal or fetal compromise but needs early delivery
4. delivery timed to suit woman or staff (see recommendation 53)

Decision to delivery (DDI) interval for unplanned CS

Perform category 1 and 2 CS as quickly as possible (see recommendation 54).

Perform category 2 CS in most situations within 75 minutes of the decision (see recommendation 55).

Take into account the condition of the woman and the unborn baby when making decisions about rapid delivery (see recommendation 56).
## Preoperative testing and preparation for CS

For pregnant women having CS for antepartum haemorrhage, abruption, uterine rupture and placenta praevia conduct the CS at a maternity unit with on-site blood transfusion services (see recommendation 59).

For healthy women with an uncomplicated pregnancy do not routinely offer the following tests:

- grouping and saving of serum
- cross-matching of blood
- a clotting screen
- preoperative ultrasound for localisation of the placenta (see recommendation 60)

Offer haemoglobin assessment to identify women with anaemia (see recommendation 58).

Offer prophylaxis for thromboembolism (choice of prophylaxis should be based on risk of thromboembolic disease) (see recommendation 93).

Offer women prophylactic antibiotics at CS before skin incision (one dose of first generation cephalosporin or ampicillin – do not use co-amoxiclav) (see recommendations 90, 91 & 92).

In women having CS with regional anaesthesia use an indwelling urinary catheter (see recommendation 61).

## Anaesthesia for CS

Offer regional anaesthesia (see recommendation 63).

Women having induction of regional anaesthesia for CS should be cared for in theatre (see recommendation 64).

Offer women having a CS under regional anaesthesia intravenous ephedrine or phenylephrine, and volume pre-loading with crystalloid or colloid (see recommendation 65).

Offer antacids and drugs (such as H₂ receptor antagonists or proton pump inhibitors) to reduce gastric volumes and acidity (see recommendation 67).

Offer anti-emetics (see recommendation 68).

General anaesthesia for unplanned CS should include preoxygenation, cricoid pressure and rapid sequence induction (see recommendation 69).

Each maternity unit should have a drill for failed intubation during obstetric anaesthesia (see recommendation 66).

Discuss post-CS analgesia options (see recommendation 62).

Intravenous ephedrine or phenylephrine should be used in the management of hypotension during CS (see recommendation 70).

The operating table for CS should have a lateral tilt of 15° (see recommendation 71).

## Surgical techniques for CS

Wear double gloves for CS for women are HIV-positive (see recommendation 72).

Use a transverse lower abdominal incision (Joel-Cohen incision) (see recommendations 74 & 75).

When there is a well formed lower uterine segment use blunt extension of the uterine incision (see recommendation 77).

Use oxytocin 5 IU by slow intravenous injection (see recommendation 80).

Remove the placenta using controlled cord traction (see recommendation 81).

Undertake intraperitoneal repair of the uterus at CS (see recommendation 82).

Suture the uterine incision with two layers (see recommendation 83).

If a midline abdominal incision is used, use mass closure with slowly absorbable continuous sutures (see recommendation 85).

Perform umbilical artery pH after all CS for suspected fetal compromise (see recommendation 89).

Accommodate women’s preferences for the birth (such as music playing in theatre) where possible (see recommendation 94).

Only use forceps if there is difficulty delivering the baby’s head (see recommendation 79).

Do not exteriorise the uterus (see recommendation 82).

Do not manually remove the placenta (see recommendation 81).

Do not use separate surgical knives to incise the skin and the deeper tissues (see recommendation 76).

Do not suture the visceral or the parietal peritoneum (see recommendation 84).

Do not routinely close the subcutaneous tissue space unless the woman has more than 2 cm subcutaneous fat (see recommendation 86).

Do not use superficial wound drains (see recommendation 87).
**Post-CS care**

**Postoperative monitoring**

After CS, a trained staff member should observe the woman on a one-to-one basis until she has regained airway control, cardiorespiratory stability and is able to communicate (see recommendation 100).

After recovery from anaesthesia continue observations (respiratory rate, heart rate, blood pressure, pain and sedation) every half hour for 2 hours and hourly thereafter. If observations are not stable, observe more frequently (see recommendation 101).

For women who have had intrathecal opioids, observe respiratory rate, sedation and pain scores hourly for at least 12 hours for diamorphine and 24 hours for morphine (see recommendation 102).

For women who have had epidural opioids or patient-controlled analgesia with opioids, monitor respiratory rate, sedation and pain scores hourly throughout treatment and for at least 2 hours after discontinuation of treatment (see recommendation 103).

**Care of the baby born by CS**

A practitioner skilled in the resuscitation of the newborn should be present at CS performed under general anaesthesia or where there is evidence of fetal compromise (see recommendation 95).

Thermal care should be in accordance with good practice for thermal care of the newborn baby (see recommendation 96).

Encourage and facilitate early skin-to-skin contact between the woman and her baby (see recommendation 97).

Offer additional support to help women start breastfeeding as soon possible after the birth of their baby (see recommendation 98).

**Recovery following CS**

Provide general postnatal care, plus specific post-CS care, and management of pregnancy complications (see recommendation 111).

Encourage women to take regular analgesia for postoperative pain (see recommendation 112).

CS wound care should include:

- removing the dressing 24 hours after the CS
- specific monitoring for fever
- assessing the wound for signs of infection (such as increasing pain, redness or discharge), separation or dehiscence
- encouraging the woman to wear loose, comfortable clothes and cotton underwear
- gently cleaning and drying the wound daily
- if needed, planning the removal of sutures or clips (see recommendation 113)

Inform women they can resume activities (such as driving, exercise) once they have fully recovered from the CS (see recommendation 117).

Inform women that after CS they are not at increased risk of difficulties with breastfeeding, depression, post-traumatic stress symptoms, dyspareunia and faecal incontinence (see recommendation 118).

**Care of the woman after CS**

Offer diamorphine (0.3–0.4 mg intrathecally) or epidural diamorphine (2.5–5 mg) for intra- and postoperative analgesia (see recommendation 104).

Offer patient-controlled analgesia using opioid analgesics (see recommendation 105).

Offer non-steroidal anti-inflammatory drugs as an adjunct to other analgesics (if there is no contraindication) (see recommendation 106).

Women who are recovering well and who have no complications can eat or drink when they feel hungry or thirsty (see recommendation 107).

Remove the urinary bladder catheter once a woman is mobile after a regional anaesthetic and not sooner than 12 hours after the last epidural ‘top up’ dose (see recommendation 108).

Do not offer routine respiratory physiotherapy to women after a CS under general anaesthesia (see recommendation 109).

Discuss the reasons for the CS and implications for the child or future pregnancies before discharge from hospital. Provide information in verbal and printed formats. If the woman prefers, provide this at a later date (see recommendation 124).

Offer early discharge (after 24 hours) with follow up at home to women who are recovering well, are afebrile and do not have complications (see recommendation 110).

**Consider CS complications:**

Endometritis if heavy and/or irregular vaginal bleeding present (see recommendation 115).

Thromboembolism if chest or leg symptoms present (see recommendation 116).

Urinary tract infection, stress incontinence or urinary tract injury if urinary symptoms present (see recommendation 114).
## 1.4 Key priorities for implementation

<table>
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<th>Number</th>
<th>Recommendation</th>
<th>See section</th>
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<td></td>
<td><strong>Morbidly adherent placenta</strong></td>
<td>5.6</td>
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| 17     | If a colour-flow doppler ultrasound scan result suggests morbidly adherent placenta:  
  - discuss with the woman the improved accuracy of magnetic resonance imaging (MRI) in addition to ultrasound to help diagnose morbidly adherent placenta and clarify the degree of invasion  
  - explain what to expect during an MRI procedure  
  - inform the woman that current experience suggests that MRI is safe, but that there is a lack of evidence about any long-term risks to the baby  
  - offer MRI if acceptable to the woman. [new 2011]                                                                 |             |
|        | **Mother-to-child transmission of HIV**                                                                                                                                                                         | 5.8         |
| 25     | Do not offer a CS on the grounds of HIV status to prevent mother-to-child transmission of HIV to:  
  - women on highly active anti-retroviral therapy (HAART) with a viral load of less than 400 copies per ml or  
  - women on any anti-retroviral therapy with a viral load of less than 50 copies per ml.  

Inform women that in these circumstances the risk of HIV transmission is the same for a CS and a vaginal birth. [new 2011] |             |
|        | **Maternal request for CS**                                                                                                                                                                                      | 5.9         |
| 36     | When a woman requests a CS because she has anxiety about childbirth, offer referral to a healthcare professional with expertise in providing perinatal mental health support to help her address her anxiety in a supportive manner. [new 2011] |             |
| 38     | For all women requesting a CS, if after discussion and offer of support (including perinatal mental health support for women with anxiety about childbirth), a vaginal birth is still not an acceptable option, offer a planned CS. [new 2011] |             |
| 39     | An obstetrician has the right to decline a woman’s request for a CS. If this happens, they should refer the woman to an NHS obstetrician in the same unit who will carry out the CS. [new 2011] |             |
|        | **Decision-to-delivery interval for unplanned CS**                                                                                                                                                             | 7.3         |
| 57     | Use the following decision to delivery intervals to measure the overall performance of an obstetric unit:  
  - 30 minutes for category 1 CS\(^1\)  
  - both 30 and 75 minutes for category 2 CS.  

Use these as audit standards only and not to judge multidisciplinary team performance for any individual CS. [new 2011] |             |
|        | **Timing of antibiotic administration**                                                                                                                                                                       | 7.6         |
| 90     | Offer women prophylactic antibiotics at CS before skin incision.                                                                                                                                               |             |

\(^1\) Category 1 CS is when there is immediate threat to the life of the woman or fetus, and category 2 CS is when there is maternal or fetal compromise which is not immediately life-threatening.
Inform them that this reduces the risk of maternal infection more than prophylactic antibiotics given after skin incision, and that no effect on the baby has been demonstrated. [new 2011]

91 Women having a CS should be offered prophylactic antibiotics, such as a single dose of first-generation cephalosporin or ampicillin, to reduce the risk of postoperative infections (such as endometritis, urinary tract and wound infection), which occur in about 8% of women who have had a CS. [A] [2004]

92 Do not use co-amoxiclav when giving antibiotics before skin incision. [new 2011]

### Pregnancy and childbirth after CS

Inform women who have had up to and including four CS that the risk of fever, bladder injuries and surgical injuries does not vary with planned mode of birth and that uterine rupture is very rare. [new 2011]

### Recovery following CS

While women are in hospital after having a CS, give them the opportunity to discuss with healthcare professionals the reasons for the CS and provide both verbal and printed information about birth options for any future pregnancies. If the woman prefers, provide this at a later date. [new 2011]

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### 1.5 Recommendations

<table>
<thead>
<tr>
<th>Number</th>
<th>Recommendation</th>
<th>See section</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Pregnant women should be offered evidence-based information and support to enable them to make informed decisions about childbirth. Addressing women's views and concerns should be recognised as being integral to the decision-making process. [C] [2004]</td>
<td>4.1</td>
</tr>
<tr>
<td>2</td>
<td>Give pregnant women evidence-based information about CS during the antenatal period, because about one in four women will have a CS. Include information about CS, such as: • indications for CS (such as presumed fetal compromise, 'failure to progress' in labour, breech presentation) • what the procedure involves • associated risks and benefits • implications for future pregnancies and birth after CS. [new 2011]</td>
<td>4.1</td>
</tr>
<tr>
<td>3</td>
<td>Communication and information should be provided in a form that is accessible to pregnant women, taking into account the information and cultural needs of minority communities and women whose first language is not English or who cannot read, together with the needs of women with disabilities or learning difficulties. [GPP] [2004]</td>
<td>4.1</td>
</tr>
</tbody>
</table>
Planning mode of birth

Discuss the risks and benefits of CS and vaginal birth with the woman (see box A and recommendation 118), taking into account her circumstances, concerns, priorities and plans for future pregnancies (including the risks of placental problems with multiple CS). [new 2011]

Box A Planned caesarean section compared with planned vaginal birth for women with an uncomplicated pregnancy and no previous caesarean section

Planned caesarean section may reduce the risk of the following in women:
- perineal and abdominal pain during birth and 3 days postpartum
- injury to vagina
- early postpartum haemorrhage
- obstetric shock.

Planned caesarean section may reduce the risk of the following in babies:
- neonatal intensive care unit admission.

Planned caesarean section may increase the risk of the following in women:
- longer hospital stay
- hysterectomy caused by postpartum haemorrhage
- cardiac arrest.

Please refer to tables 4.5 and 4.6 for full details, including the absolute and relative risks for each effect.

Do not use a body mass index (BMI) of over 50 alone as an indication for planned CS. [new 2011]

Consent for CS should be requested after providing pregnant women with evidence-based information and in a manner that respects the woman’s dignity, privacy, views and culture, while taking into consideration the clinical situation. [G] [2004]

A pregnant woman is entitled to decline the offer of treatment such as CS, even when the treatment would clearly benefit her or her baby’s health. Refusal of treatment needs to be one of the woman’s options. [D] [2004]

When a decision is made to perform a CS, a record should be made of all the factors that influence the decision, and which of these is the most influential. [GPP] [2004]

Breech presentation

Women who have an uncomplicated singleton breech pregnancy at 36 weeks’ gestation should be offered external cephalic version. Exceptions include women in labour and women with a uterine scar or abnormality, fetal compromise, ruptured membranes, vaginal bleeding or medical conditions. [A] [2004]

Pregnant women with a singleton breech presentation at term, for whom external cephalic version is contraindicated or has been unsuccessful, should be offered CS because it reduces
perinatal mortality and neonatal morbidity. [A] [2004]

### Multiple pregnancy

11 In otherwise uncomplicated twin pregnancies at term where the presentation of the first twin is cephalic, perinatal morbidity and mortality is increased for the second twin. However, the effect of planned CS in improving outcome for the second twin remains uncertain and therefore CS should not routinely be offered outside a research context. [C] [2004]

12 In twin pregnancies where the first twin is not cephalic the effect of CS in improving outcome is uncertain, but current practice is to offer a planned CS. [GPP] [2004]

### Preterm birth and CS

13 Preterm birth is associated with higher neonatal morbidity and mortality. However, the effect of planned CS in improving these outcomes remains uncertain and therefore CS should not routinely be offered outside a research context. [C] [2004]

### Small for gestational age

14 The risk of neonatal morbidity and mortality is higher with ‘small for gestational age’ babies. However, the effect of planned CS in improving these outcomes remains uncertain and therefore CS should not routinely be offered outside a research context. [C] [2004]

### Placenta praevia

15 Women with a placenta that partly or completely covers the internal cervical os (minor or major placenta praevia) should be offered CS. [D] [2004]

### Morbidly adherent placenta

16 If low-lying placenta is confirmed at 32-34 weeks in women who have had a previous CS, offer colour-flow doppler ultrasound as the first diagnostic test for morbidly adherent placenta. [new 2011]

17 If a colour-flow doppler ultrasound scan result suggests morbidly adherent placenta:

- discuss with the woman the improved accuracy of magnetic resonance imaging (MRI) in addition to ultrasound to help diagnose morbidly adherent placenta and clarify the degree of invasion
- explain what to expect during an MRI procedure
- inform the woman that current experience suggests that MRI is safe, but that there is a lack of evidence about any long-term risks to the baby
- offer MRI if acceptable to the woman. [new 2011]

18 Discuss the interventions available for delivery with women suspected to have morbidly adherent placenta, including cross matching of blood and planned CS with a consultant obstetrician present. [new 2011]

19 When performing a CS for women suspected to have morbidly adherent placenta, ensure that:

- a consultant obstetrician and a consultant anaesthetist are present
• an experienced paediatrician is present
• a senior haematologist is available for advice
• a critical care bed is available
• sufficient cross-matched blood and blood products are readily available. [new 2011]

20 When performing a CS for women suspected to have morbidly adherent placenta the consultant obstetrician should decide which other healthcare professionals need to be consulted or present. [new 2011]

21 All hospitals should have a locally agreed protocol for managing morbidly adherent placenta that sets out how these elements of care should be provided. [new 2011]

Predicting CS for cephalopelvic disproportion in labour

22 Pelvimetry is not useful in predicting ‘failure to progress’ in labour and should not be used in decision making about mode of birth. [A] [2004]

23 Shoe size, maternal height and estimations of fetal size (ultrasound or clinical examination) do not accurately predict cephalopelvic disproportion and should not be used to predict ‘failure to progress’ during labour. [B] [2004]

Mother-to-child transmission of maternal infections

24 As early as possible give women with HIV information about the risks and benefits for them and their child of the HIV treatment options and mode of birth so that they can make an informed decision. [new 2011]

25 Do not offer a CS on the grounds of HIV status to prevent mother-to-child transmission of HIV to:

• women on highly active anti-retroviral therapy (HAART) with a viral load of less than 400 copies per ml or
• women on any anti-retroviral therapy with a viral load of less than 50 copies per ml.

Inform women that in these circumstances the risk of HIV transmission is the same for a CS and a vaginal birth. [new 2011]

26 Consider either a vaginal birth or a CS for women on anti-retroviral therapy (ART) with a viral load 50–400 copies per ml because there is insufficient evidence that a CS prevents mother-to-child transmission of HIV. [new 2011]

27 Offer CS to women with HIV who:

• are not receiving any anti-retroviral therapy or
• are receiving any anti-retroviral therapy and have a viral load of 400 copies per ml or more. [new 2011]

28 Researchers and national bodies responsible for the collection of UK population data should continue to collect data about HIV diagnoses in pregnant women, including treatment, mode of birth, and mother-to-child transmission rates. [new 2011]

29 Mother-to-child transmission of hepatitis B can be reduced if the baby receives immunoglobulin and vaccination. In these situations pregnant women with hepatitis B should not be offered a planned CS because there is insufficient evidence that this reduces mother-to-child transmission of hepatitis B virus.
Women who are infected with hepatitis C should not be offered a planned CS because this does not reduce mother-to-child transmission of the virus. [C] [2004]

Pregnant women who are co-infected with hepatitis C virus and HIV should be offered planned CS because it reduces mother-to-child transmission of both hepatitis C virus and HIV. [C] [2004]

Women with primary genital herpes simplex virus (HSV) infection occurring in the third trimester of pregnancy should be offered planned CS because it decreases the risk of neonatal HSV infection. [C] [2004]

Pregnant women with a recurrence of HSV at birth should be informed that there is uncertainty about the effect of planned CS in reducing the risk of neonatal HSV infection. Therefore, CS should not routinely be offered outside a research context. [C] [2004]

Maternal request for CS

When a woman requests a CS explore, discuss and record the specific reasons for the request. [new 2011]

If a woman requests a CS when there is no other indication, discuss the overall risks and benefits of CS compared with vaginal birth (see box A on page 14) and record that this discussion has taken place. Include a discussion with other members of the obstetric team (including the obstetrician, midwife and anaesthetist) if necessary to explore the reasons for the request, and to ensure the woman has accurate information. [new 2011]

When a woman requests a CS because she has anxiety about childbirth, offer referral to a healthcare professional with expertise in providing perinatal mental health support to help her address her anxiety in a supportive manner. [new 2011]

Ensure the healthcare professional providing perinatal mental health support has access to the planned place of birth during the antenatal period in order to provide care. [new 2011]

For all women requesting a CS, if after discussion and offer of support (including perinatal mental health support for women with anxiety about childbirth), a vaginal birth is still not an acceptable option, offer a planned CS. [new 2011]

An obstetrician has the right to decline a woman’s request for a CS. If this happens, they should refer the woman to an NHS obstetrician in the same unit who will carry out the CS. [new 2011]

Place of birth

During their discussions about options for birth, healthy pregnant women with anticipated uncomplicated pregnancies should be informed that planning a home birth reduces the likelihood of CS. [B] [2004]

During their discussions about options for birth, healthy pregnant women with anticipated uncomplicated pregnancies should be informed that planned childbirth in a ‘midwifery-led unit’ does not
### Factors reducing the likelihood of CS

42 Women should be informed that continuous support during labour from women with or without prior training reduces the likelihood of CS. [A] [2004]

43 Women with an uncomplicated pregnancy should be offered induction of labour beyond 41 weeks because this reduces the risk of perinatal mortality and the likelihood of CS. [A] [2004]

44 A partogram with a 4-hour action line should be used to monitor progress of labour of women in spontaneous labour with an uncomplicated singleton pregnancy at term, because it reduces the likelihood of CS. [A] [2004]

45 Consultant obstetricians should be involved in the decision making for CS, because this reduces the likelihood of CS. [C] [2004]

46 Electronic fetal monitoring is associated with an increased likelihood of CS. When CS is contemplated because of an abnormal fetal heart rate pattern, in cases of suspected fetal acidosis, fetal blood sampling should be offered if it is technically possible and there are no contraindications. [B] [2004]

### No influence on likelihood of CS

47 Women should be informed that the following interventions during intrapartum care have not been shown to influence the likelihood of CS, although they may affect other outcomes that are outside the scope of this guideline:

- walking in labour
- non-supine position during the second stage of labour
- immersion in water during labour
- epidural analgesia during labour
- the use of raspberry leaves. [A] [2004]

48 Women should be informed that the effects on the likelihood of CS of complementary therapies used during labour (such as acupuncture, aromatherapy, hypnosis, herbal products, nutritional supplements, homeopathic medicines, and Chinese medicines) have not been properly evaluated and further research is needed before such interventions can be recommended. [D] [2004]

### ‘Failure to progress’ in labour and CS

49 The following aspects of intrapartum care have not been shown to influence the likelihood of CS for ‘failure to progress’ and should not be offered for this reason, although they may affect other outcomes which are outside the scope of this guideline:

- active management of labour
- early amniotomy. [A] [2004]

### Eating during labour

50 Women should be informed that eating a low-residue diet during labour (toast, crackers, low-fat cheese) results in larger gastric volumes, but the effect on the risk of aspiration if anaesthesia is
Women should be informed that having isotonic drinks during labour prevents ketosis without a concomitant increase in gastric volume. [A] [2004]

Timing of planned CS

The risk of respiratory morbidity is increased in babies born by CS before labour, but this risk decreases significantly after 39 weeks. Therefore planned CS should not routinely be carried out before 39 weeks. [B] [2004]

Classification of urgency

The urgency of CS should be documented using the following standardised scheme in order to aid clear communication between healthcare professionals about the urgency of a CS:

1. immediate threat to the life of the woman or fetus
2. maternal or fetal compromise which is not immediately life-threatening
3. no maternal or fetal compromise but needs early delivery
4. delivery timed to suit woman or staff. [C] [2004]

Decision-to-delivery interval for unplanned CS

Perform category 1 and 2 CS\(^2\) as quickly as possible after making the decision, particularly for category 1. [new 2011]

Perform category 2 CS\(^2\) in most situations within 75 minutes of making the decision. [new 2011]

Take into account the condition of the woman and the unborn baby when making decisions about rapid delivery. Remember that rapid delivery may be harmful in certain circumstances. [new 2011]

Use the following decision-to-delivery intervals to measure the overall performance of an obstetric unit:

- 30 minutes for category 1 CS\(^2\)
- both 30 and 75 minutes for category 2 CS.

Use these as audit standards only and not to judge multidisciplinary team performance for any individual CS. [new 2011]

Preoperative testing and preparation for CS

Pregnant women should be offered a haemoglobin assessment before CS to identify those who have anaemia. Although blood loss of more than 1000 ml is infrequent after CS (it occurs in 4-8% of CS) it is a potentially serious complication. [C] [2004]

Pregnant women having CS for antepartum haemorrhage, abruption, uterine rupture and placenta praevia are at increased risk of blood loss of more than 1000 ml and should have the CS carried out at a maternity unit with on-site blood transfusion services. [C] [2004]

\(^2\) Category 1 CS is when there is immediate threat to the life of the woman or fetus, and category 2 CS is when there is maternal or fetal compromise which is not immediately life-threatening.
Pregnant women who are healthy and who have otherwise uncomplicated pregnancies should not routinely be offered the following tests before CS:

- grouping and saving of serum
- cross-matching of blood
- a clotting screen
- preoperative ultrasound for localisation of the placenta, because this does not improve CS morbidity outcomes (such as blood loss of more than 1000 ml, injury of the infant, and injury to the cord or to other adjacent structures). [C] [2004]

Women having CS with regional anaesthesia require an indwelling urinary catheter to prevent over-distension of the bladder because the anaesthetic block interferes with normal bladder function. [GPP] [2004]

**Anaesthesia for CS**

Pregnant women having a CS should be given information on different types of post-CS analgesia so that analgesia best suited to their needs can be offered (see recommendation 104). [GPP] [2004]

Women who are having a CS should be offered regional anaesthesia because it is safer and results in less maternal and neonatal morbidity than general anaesthesia. This includes women who have a diagnosis of placenta praevia. [A] [2004]

Women who are having induction of regional anaesthesia for CS should be cared for in theatre because this does not increase women’s anxiety. [B] [2004]

Women who are having a CS under regional anaesthesia should be offered intravenous ephedrine or phenylephrine, and volume pre-loading with crystalloid or colloid to reduce the risk of hypotension occurring during CS. [A] [2004]

Each maternity unit should have a drill for failed intubation during obstetric anaesthesia. [D] [2004]

To reduce the risk of aspiration pneumonitis women should be offered antacids and drugs (such as H₂ receptor antagonists or proton pump inhibitors) to reduce gastric volumes and acidity before CS. [B] [2004]

Women having a CS should be offered antiemetics (either pharmacological or acupressure) to reduce nausea and vomiting during CS. [A] [2004]

General anaesthesia for unplanned CS should include preoxygenation, cricoid pressure and rapid sequence induction to reduce the risk of aspiration. [GPP] [2004]

Intravenous ephedrine or phenylephrine should be used in the management of hypotension during CS. [A] [2004]

The operating table for CS should have a lateral tilt of 15°, because this reduces maternal hypotension. [A] [2004]
<table>
<thead>
<tr>
<th>Section</th>
<th>Text</th>
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</thead>
<tbody>
<tr>
<td>72</td>
<td>Healthcare professionals should wear double gloves when performing or assisting at CS on women who have tested positive for HIV, to reduce the risk of HIV infection of healthcare professionals during surgery. [A] [2004]</td>
</tr>
<tr>
<td>73</td>
<td>General recommendations for safe surgical practice should be followed at CS to reduce the risk of HIV infection of staff. [C] [2004]</td>
</tr>
<tr>
<td>74</td>
<td>CS should be performed using a transverse abdominal incision because this is associated with less postoperative pain and an improved cosmetic effect compared with a midline incision. [B] [2004]</td>
</tr>
<tr>
<td>75</td>
<td>The transverse incision of choice should be the Joel Cohen incision (a straight skin incision, 3 cm above the symphysis pubis; subsequent tissue layers are opened bluntly and, if necessary, extended with scissors and not a knife), because it is associated with shorter operating times and reduced postoperative febrile morbidity. [A] [2004]</td>
</tr>
<tr>
<td>76</td>
<td>The use of separate surgical knives to incise the skin and the deeper tissues at CS is not recommended because it does not decrease wound infection. [B] [2004]</td>
</tr>
<tr>
<td>77</td>
<td>When there is a well formed lower uterine segment, blunt rather than sharp extension of the uterine incision should be used because it reduces blood loss, incidence of postpartum haemorrhage and the need for transfusion at CS. [A] [2004]</td>
</tr>
<tr>
<td>78</td>
<td>Women who are having a CS should be informed that the risk of fetal lacerations is about 2%. [C] [2004]</td>
</tr>
<tr>
<td>79</td>
<td>Forceps should only be used at CS if there is difficulty delivering the baby's head. The effect on neonatal morbidity of the routine use of forceps at CS remains uncertain. [C] [2004]</td>
</tr>
<tr>
<td>80</td>
<td>Oxytocin 5 IU by slow intravenous injection should be used at CS to encourage contraction of the uterus and to decrease blood loss. [C] [2004]</td>
</tr>
<tr>
<td>81</td>
<td>At CS, the placenta should be removed using controlled cord traction and not manual removal as this reduces the risk of endometritis. [A] [2004]</td>
</tr>
<tr>
<td>82</td>
<td>Intraperitoneal repair of the uterus at CS should be undertaken. Exteriorisation of the uterus is not recommended because it is associated with more pain and does not improve operative outcomes such as haemorrhage and infection. [A] [2004]</td>
</tr>
<tr>
<td>83</td>
<td>The effectiveness and safety of single layer closure of the uterine incision is uncertain. Except within a research context, the uterine incision should be sutured with two layers. [B] [2004]</td>
</tr>
<tr>
<td>84</td>
<td>Neither the visceral nor the parietal peritoneum should be sutured at CS because this reduces operating time and the need for postoperative analgesia, and improves maternal satisfaction. [A] [2004]</td>
</tr>
</tbody>
</table>
In the rare circumstances that a midline abdominal incision is used at CS, mass closure with slowly absorbable continuous sutures should be used because this results in fewer incisional hernias and less dehiscence than layered closure.  
[B] [2004]

Routine closure of the subcutaneous tissue space should not be used, unless the woman has more than 2 cm subcutaneous fat, because it does not reduce the incidence of wound infection.  
[A] [2004]

Superficial wound drains should not be used at CS because they do not decrease the incidence of wound infection or wound haematoma.  
[A] [2004]

Obstetricians should be aware that the effects of different suture materials or methods of skin closure at CS are not certain.  
[C] [2004]

Umbilical artery pH should be performed after all CS for suspected fetal compromise, to allow review of fetal wellbeing and guide ongoing care of the baby.  
[B] [2004]

Offer women prophylactic antibiotics at CS before skin incision. Inform them that this reduces the risk of maternal infection more than prophylactic antibiotics given after skin incision, and that no effect on the baby has been demonstrated [new 2011]

Women having a CS should be offered prophylactic antibiotics, such as a single dose of first-generation cephalosporin or ampicillin, to reduce the risk of postoperative infections (such as endometritis, urinary tract and wound infection), which occur in about 8% of women who have had a CS.  
[A] [2004]

Do not use co-amoxiclav when giving antibiotics before skin incision [new 2011]

Women having a CS should be offered thromboprophylaxis because they are at increased risk of venous thromboembolism. The choice of method of prophylaxis (for example, graduated stockings, hydration, early mobilisation, low molecular weight heparin) should take into account risk of thromboembolic disease and follow existing guidelines.  
[D] [2004]

Women’s preferences for the birth, such as music playing in theatre, lowering the screen to see the baby born, or silence so that the mother’s voice is the first the baby hears, should be accommodated where possible. [GPP] [2004]

Presence of paediatrician at CS

An appropriately trained practitioner skilled in the resuscitation of the newborn should be present at CS performed under general anaesthesia or where there is evidence of fetal compromise.  
[C] [2004]

Thermal care for babies born by CS

Babies born by CS are more likely to have a lower temperature, and thermal care should be in accordance with good practice for thermal care of the newborn baby.  
[GPP] [2004]
Maternal contact (skin-to-skin)

Early skin-to-skin contact between the woman and her baby should be encouraged and facilitated because it improves maternal perceptions of the infant, mothering skills, maternal behaviour, and breastfeeding outcomes, and reduces infant crying. [A] [2004]

Breastfeeding

Women who have had a CS should be offered additional support to help them to start breastfeeding as soon as possible after the birth of their baby. This is because women who have had a CS are less likely to start breastfeeding in the first few hours after the birth, but, when breastfeeding is established, they are as likely to continue as women who have a vaginal birth. [A] [2004]

High dependency unit/intensive therapy unit admission

Healthcare professionals caring for women after CS should be aware that, although it is rare for women to need intensive care following childbirth, this occurs more frequently after CS (about 9 per 1000). [B] [2004]

Routine monitoring after CS

After CS, women should be observed on a one-to-one basis by a properly trained member of staff until they have regained airway control and cardiorespiratory stability and are able to communicate. [D] [2004]

After recovery from anaesthesia, observations (respiratory rate, heart rate, blood pressure, pain and sedation) should be continued every half hour for 2 hours, and hourly thereafter provided that the observations are stable or satisfactory. If these observations are not stable, more frequent observations and medical review are recommended. [GPP] [2004]

For women who have had intrathecal opioids, there should be a minimum hourly observation of respiratory rate, sedation and pain scores for at least 12 hours for diamorphine and 24 hours for morphine. [GPP] [2004]

For women who have had epidural opioids or patient-controlled analgesia with opioids, there should be routine hourly monitoring of respiratory rate, sedation and pain scores throughout treatment and for at least 2 hours after discontinuation of treatment. [GPP] [2004]

Pain management after CS

Women should be offered diamorphine (0.3–0.4 mg intrathecally) for intra- and postoperative analgesia because it reduces the need for supplemental analgesia after a CS. Epidural diamorphine (2.5–5 mg) is a suitable alternative. [A] [2004]

Patient-controlled analgesia using opioid analgesics should be offered after CS because it improves pain relief. [GPP] [2004]

Providing there is no contraindication, non-steroidal anti-inflammatory drugs should be offered post-CS as an adjunct to other analgesics, because they reduce the need for opioids.
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<tbody>
<tr>
<td>Early eating and drinking after CS</td>
<td>107</td>
<td>Women who are recovering well after CS and who do not have complications can eat and drink when they feel hungry or thirsty. [A] [2004]</td>
</tr>
<tr>
<td>Urinary catheter removal after CS</td>
<td>108</td>
<td>Removal of the urinary bladder catheter should be carried out once a woman is mobile after a regional anaesthetic and not sooner than 12 hours after the last epidural ‘top up’ dose. [D] [2004]</td>
</tr>
<tr>
<td>Respiratory physiotherapy after CS</td>
<td>109</td>
<td>Routine respiratory physiotherapy does not need to be offered to women after a CS under general anaesthesia, because it does not improve respiratory outcomes such as coughing, phlegm, body temperature, chest palpation and auscultatory changes. [A] [2004]</td>
</tr>
<tr>
<td>Length of hospital stay and readmission to hospital</td>
<td>110</td>
<td>Length of hospital stay is likely to be longer after a CS (an average of 3–4 days) than after a vaginal birth (average 1–2 days). However, women who are recovering well, are apyrexial and do not have complications following CS should be offered early discharge (after 24 hours) from hospital and follow-up at home, because this is not associated with more infant or maternal readmissions. [A] [2004]</td>
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<tr>
<td>Recovery following CS</td>
<td>111</td>
<td>In addition to general postnatal care, women who have had a CS should be provided with:</td>
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<td></td>
<td></td>
<td>• specific care related to recovery after CS</td>
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<td>• care related to management of other complications during pregnancy or childbirth. [GPP] [2004]</td>
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<td></td>
<td>112</td>
<td>Women who have a CS should be prescribed and encouraged to take regular analgesia for postoperative pain, using:</td>
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<td>• for severe pain, co-codamol with added ibuprofen</td>
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<td>• for moderate pain, co-codamol</td>
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<td></td>
<td></td>
<td>• for mild pain, paracetamol. [D] [2004]</td>
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<tr>
<td></td>
<td>113</td>
<td>CS wound care should include:</td>
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<td>• removing the dressing 24 hours after the CS</td>
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<td>• specific monitoring for fever</td>
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<td>• assessing the wound for signs of infection (such as increasing pain, redness or discharge), separation or dehiscence</td>
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<td>• encouraging the woman to wear loose, comfortable clothes and cotton underwear</td>
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<td>• gently cleaning and drying the wound daily</td>
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<td>• if needed, planning the removal of sutures or clips. [D] [2004]</td>
</tr>
<tr>
<td></td>
<td>114</td>
<td>Healthcare professionals caring for women who have had a CS and who have urinary symptoms should consider the possible...</td>
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</table>
diagnosis of:

- urinary tract infection
- stress incontinence (occurs in about 4% of women after CS)
- urinary tract injury (occurs in about 1 per 1000 CS).

[D] [2004]

115 Healthcare professionals caring for women who have had a CS and who have heavy and/or irregular vaginal bleeding should consider that this is more likely to be due to endometritis than retained products of conception. [D] [2004]

116 Women who have had a CS are at increased risk of thromboembolic disease (both deep vein thrombosis and pulmonary embolism), so healthcare professionals need to pay particular attention to women who have chest symptoms (such as cough or shortness of breath) or leg symptoms (such as painful swollen calf). [D] [2004]

117 Women who have had a CS should resume activities such as driving a vehicle, carrying heavy items, formal exercise and sexual intercourse once they have fully recovered from the CS (including any physical restrictions or distracting effect due to pain). [GPP] [2004]

118 Healthcare professionals caring for women who have had a CS should inform women that after a CS they are not at increased risk of difficulties with breastfeeding, depression, post-traumatic stress symptoms, dyspareunia and faecal incontinence. [D] [2004]

Pregnancy and childbirth after CS

119 When advising about the mode of birth after a previous CS consider:

- maternal preferences and priorities
- the risks and benefits of repeat CS
- the risks and benefits of planned vaginal birth after CS, including the risk of unplanned CS. [new 2011]

120 Inform women who have had up to and including four CS that the risk of fever, bladder injuries and surgical injuries does not vary with planned mode of birth and that uterine rupture is very rare. [new 2011]

121 Offer women planning a vaginal birth who have had a previous CS:

- electronic fetal monitoring during labour
- care during labour in a unit where there is immediate access to CS and on-site blood transfusion services. [GPP] [2011]

122 During induction of labour, women who have had a previous CS should be monitored closely, with access to electronic fetal monitoring and with immediate access to CS, because they are at increased risk of uterine rupture. [GPP] [2004]

For more information see ‘Induction of labour’ (NICE clinical guideline 70).
Pregnant women with both previous CS and a previous vaginal birth should be informed that they have an increased likelihood of achieving a vaginal birth than women who have had a previous CS but no previous vaginal birth. [B] [2004]

While women are in hospital after having a CS, give them the opportunity to discuss with healthcare professionals the reasons for the CS and provide both verbal and printed information about birth options for any future pregnancies. If the woman prefers, provide this at a later date. [new 2011]
<table>
<thead>
<tr>
<th>Effects around the time of birth</th>
<th>Finding for planned CS</th>
<th>Finding for planned vaginal birth (including % unplanned CS in planned vaginal birth group)</th>
<th>Absolute effect</th>
<th>Relative effect (95% confidence interval)</th>
<th>Evidence quality and reference</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Studies suggest may be reduced after a planned CS</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Perineal and abdominal pain during birth (score/10 – higher scores indicate higher pain levels)</td>
<td>Median score 1.0</td>
<td>Median score 7.3 (10.3%)</td>
<td>6.3 lower</td>
<td>NC</td>
<td>Very low</td>
</tr>
<tr>
<td>Perineal and abdominal pain 3 days postpartum (score/10 – higher scores indicate higher pain levels)</td>
<td>Median score 4.5</td>
<td>Median score 5.2 (10.3%)</td>
<td>0.7 lower</td>
<td>NC</td>
<td>Very low</td>
</tr>
<tr>
<td>Injury to vagina</td>
<td>0.0%</td>
<td>0.56% (14.7%)</td>
<td>6 fewer per 1000 (from 6 fewer to 2 fewer)</td>
<td>NC</td>
<td>Very low</td>
</tr>
<tr>
<td>Early postpartum haemorrhage</td>
<td>1.1% (35%)</td>
<td>6.0%</td>
<td>49 per 1000 (from 4 fewer to 56 fewer)</td>
<td>OR 0.23 (0.06 to 0.94)</td>
<td>Low</td>
</tr>
<tr>
<td></td>
<td>3.9% (8.3%)</td>
<td>6.2%</td>
<td>23 fewer per 1000 (from 35 fewer to 6 fewer)</td>
<td>RR 0.06 (0.4 to 0.9)</td>
<td>Very low</td>
</tr>
<tr>
<td>Obstetric shock</td>
<td>0.006% (8.2%)</td>
<td>0.018%</td>
<td>12 fewer per 100,000 (from 17 fewer to 0.1 fewer)</td>
<td>RR 0.33 (0.11 to 0.99)</td>
<td>Very low</td>
</tr>
<tr>
<td><strong>Studies suggest may be reduced after planned vaginal birth</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Length of hospital stay</td>
<td>3.2 days (35%)</td>
<td>2.6 days</td>
<td>0.6 days longer</td>
<td>Mean difference 1.58 (1.27 to 2.17)</td>
<td>Low</td>
</tr>
</tbody>
</table>
## Caesarean Section

### 2011 Update

<table>
<thead>
<tr>
<th>Event</th>
<th>Control 1</th>
<th>Control 2</th>
<th>Adjusted Mean Difference</th>
<th>Risk Ratio</th>
<th>Confidence Interval</th>
<th>Rating</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hysterectomy due to post-partum haemorrhage</td>
<td>0.03%</td>
<td>0.01%</td>
<td>14 more per 100,000 (from 3 more to 33 more)</td>
<td>RR 2.31</td>
<td>(1.30 to 4.09)</td>
<td>Very low</td>
</tr>
<tr>
<td>Cardiac arrest</td>
<td>0.19%</td>
<td>0.03%</td>
<td>15 more per 10,000 (from 11.5 more to 19.5 more)</td>
<td>RR 4.91</td>
<td>(3.95 to 6.11)</td>
<td>Very low</td>
</tr>
</tbody>
</table>

### No difference found in studies

#### Perineal and abdominal pain 4 months postpartum (score / 10 – higher scores indicate higher pain levels)

<table>
<thead>
<tr>
<th>Median score</th>
<th>Median score 0.17</th>
<th>0.17 lower</th>
<th>Rating</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.0</td>
<td>0.17</td>
<td>NC</td>
<td>Very low</td>
</tr>
</tbody>
</table>

#### Injury to bladder/ureter

<table>
<thead>
<tr>
<th>Median score</th>
<th>0.14%</th>
<th>1 fewer per 1000 (from 2 fewer to 2 more)</th>
<th>NC</th>
<th>Very low</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.0%</td>
<td>(14.7%)</td>
<td>(14.7%)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

#### Injury to cervix

<table>
<thead>
<tr>
<th>Median score</th>
<th>0.28%</th>
<th>3 fewer per 1000 (from 3 fewer to 1 more)</th>
<th>NC</th>
<th>Very low</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.0%</td>
<td>(14.7%)</td>
<td>(14.7%)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

#### Iatrogenic surgical injury

<table>
<thead>
<tr>
<th>Median score</th>
<th>0.07%</th>
<th>7 fewer per 10,000</th>
<th>NC</th>
<th>Very low</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.00%</td>
<td>(14.7%)</td>
<td>(from 10 fewer to 30 more)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

#### Pulmonary embolism

<table>
<thead>
<tr>
<th>Median score</th>
<th>0.003%</th>
<th>2 fewer per 10,000 (from 2 fewer to 40 more)</th>
<th>NC</th>
<th>Very low</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.00%</td>
<td>(14.7%)</td>
<td>(14.7%)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

#### Wound infection

<table>
<thead>
<tr>
<th>Median score</th>
<th>0.00%</th>
<th>1 more per 10,000</th>
<th>p = 1.0</th>
<th>Low</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.01%</td>
<td>(35%)</td>
<td>(35%)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Median score</th>
<th>0.9%</th>
<th>6 fewer per 1000 (from 1 fewer to 19 more)</th>
<th>RR 1.7</th>
<th>(0.9 to 3.2)</th>
<th>Very low</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.5%</td>
<td>(8.3%)</td>
<td>(8.3%)</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

#### Intraoperative trauma

<table>
<thead>
<tr>
<th>Median score</th>
<th>0.3%</th>
<th>1 fewer per 1000 (from 3 fewer to 7 more)</th>
<th>RR 0.5</th>
<th>(0.1 to 3.5)</th>
<th>Very low</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.1%</td>
<td>(8.3%)</td>
<td>(8.3%)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Event</td>
<td>Rate 1</td>
<td>Rate 2</td>
<td>Adjusted Rate Difference</td>
<td>Risk Ratio (Confidence Interval)</td>
<td>Evidence Level</td>
</tr>
<tr>
<td>--------------------------------</td>
<td>--------</td>
<td>--------</td>
<td>--------------------------</td>
<td>---------------------------------</td>
<td>---------------</td>
</tr>
<tr>
<td>Uterine rupture</td>
<td>0.02%</td>
<td>0.03%</td>
<td>13 fewer per 100,000 (from 22 fewer to 2.2 more)</td>
<td>RR 0.51 (0.25 to 1.07)</td>
<td>Very low</td>
</tr>
<tr>
<td>Assisted ventilation or intubation</td>
<td>0.01%</td>
<td>0.005%</td>
<td>7 more per 100,000 (from 0 fewer to 22 more)</td>
<td>RR 2.21 (0.99 to 4.90)</td>
<td>Very low</td>
</tr>
<tr>
<td>Acute renal failure</td>
<td>0.004%</td>
<td>0.001%</td>
<td>2 more per 100,000 (from 9 fewer to 13 more)</td>
<td>RR 2.17 (0.58 to 8.14)</td>
<td>Very low</td>
</tr>
<tr>
<td>Conflicting findings from studies</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Maternal death</td>
<td>9/737</td>
<td>49/9133 (cases/controls)</td>
<td>NC</td>
<td>OR 2.28 (1.11 to 4.65)</td>
<td>Very low</td>
</tr>
<tr>
<td>Deep vein thrombosis</td>
<td>0.00%</td>
<td>0.03%</td>
<td>0.7 fewer per 1000 (from 0.2 fewer to 4 more)</td>
<td>NC</td>
<td>Very low</td>
</tr>
<tr>
<td></td>
<td>0.06%</td>
<td>0.03%</td>
<td>32 more per 100,000 (from 14 more to 59 more)</td>
<td>RR 2.20 (1.51 to 3.20)</td>
<td>Very low</td>
</tr>
<tr>
<td>Blood transfusion</td>
<td>1.7%</td>
<td>1.9%</td>
<td>2 fewer per 1000 (from 14 fewer to 34 more)</td>
<td>OR 0.87 (0.27 to 2.78)</td>
<td>Low</td>
</tr>
<tr>
<td></td>
<td>0.3%</td>
<td>0.3%</td>
<td>0 fewer per 1000 (from 2 fewer to 5 more)</td>
<td>RR 0.89 (0.20 to 3.99)</td>
<td>Very low</td>
</tr>
<tr>
<td></td>
<td>0.3%</td>
<td>0.4%</td>
<td>1 fewer per 1000 (from 2 fewer to 34 more)</td>
<td>RR 0.7 (0.2 to 2.7)</td>
<td>Very low</td>
</tr>
<tr>
<td>Procedure</td>
<td>Risk 1 (%)</td>
<td>Risk 2 (%)</td>
<td>Change (95% CI)</td>
<td>RR (95% CI)</td>
<td>Level</td>
</tr>
<tr>
<td>------------------------------------------</td>
<td>------------</td>
<td>------------</td>
<td>--------------------------------------------------------------------------------</td>
<td>-------------</td>
<td>-------</td>
</tr>
<tr>
<td>Caesarean section</td>
<td>0.02%</td>
<td>0.07%</td>
<td>fewer to 5 more</td>
<td>RR 0.20 (0.20 to 0.64)</td>
<td>Very low</td>
</tr>
<tr>
<td></td>
<td>(8.3%)</td>
<td>(8.2%)</td>
<td>41 fewer per 100,000 (from 53 fewer to 23 fewer)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Infection – wound and postpartum</td>
<td>1.1%</td>
<td>0.8%</td>
<td>3 more per 1000 (from 2 fewer to 11 more)</td>
<td>RR 1.36 (0.75 to 2.4)</td>
<td>Very low</td>
</tr>
<tr>
<td></td>
<td>(14.7%)</td>
<td>(8.2%)</td>
<td>390 more per 100,000 (from 323 more to 464 more)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hysterectomy</td>
<td>0.6%</td>
<td>0.1%</td>
<td>5 more per 1000</td>
<td>p = 0.13</td>
<td>Low</td>
</tr>
<tr>
<td></td>
<td>(35%)</td>
<td>(14.7%)</td>
<td>1 more per 1000 (from 0 more to 5 more)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>0.1%</td>
<td>0.01%</td>
<td>1 more per 1000 (from 0 more to 5 more)</td>
<td>RR 9.09 (1.36 to 60.33)</td>
<td>Very low</td>
</tr>
<tr>
<td></td>
<td>(14.7%)</td>
<td>(14.7%)</td>
<td>41 more per 100,000 (from 23.6 more to 68 more)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Anaesthetic complications</td>
<td>0.4%</td>
<td>0.3%</td>
<td>1 more per 1000 (from 2 fewer to 11 more)</td>
<td>RR 1.24 (0.34 to 4.59)</td>
<td>Very low</td>
</tr>
<tr>
<td></td>
<td>(14.7%)</td>
<td>(14.7%)</td>
<td>319 more per 100,000 (from 257 more to 389 more)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>0.53%</td>
<td>0.21%</td>
<td></td>
<td>RR 2.5 (2.22 to 2.86)</td>
<td>Very low</td>
</tr>
</tbody>
</table>
### Table 4.6 Summary effect on babies’ health of planned CS compared with planned vaginal birth for women with an uncomplicated pregnancy and no previous CS

<table>
<thead>
<tr>
<th>Effects around the time of birth</th>
<th>Finding for CS</th>
<th>Finding for vaginal birth</th>
<th>Absolute effect</th>
<th>Relative effect (95% confidence interval)</th>
<th>Evidence quality and reference</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Studies suggest may be reduced after planned vaginal birth</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>NICU admission</td>
<td>13.9% (35%)</td>
<td>6.3% (35%)</td>
<td>76 more per 1000 (from 31 more to 134 more)</td>
<td>RR 2.20 (1.4 to 3.18)</td>
<td>Low</td>
</tr>
<tr>
<td><strong>No difference found in studies</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hypoxic-Ischemic Encephalopathy (CNS depression, seizures, pH &lt; 7)</td>
<td>0.2% (14.7%)</td>
<td>0.2% (14.7%)</td>
<td>0 fewer per 1000 (from 2 fewer to 5 more)</td>
<td>RR 0.81 (0.22 to 3.00)</td>
<td>Very low</td>
</tr>
<tr>
<td>Intracranial haemorrhage</td>
<td>0.00% (14.7%)</td>
<td>0.01% (14.7%)</td>
<td>0.2 fewer per 1000 (from 0.4 fewer to 3 more)</td>
<td>NC</td>
<td>Very low</td>
</tr>
<tr>
<td>Neonatal respiratory morbidity</td>
<td>12.0% (14.7%)</td>
<td>11.5% (14.7%)</td>
<td>5 more per 1000 (from 14 fewer to 27 more)</td>
<td>RR 1.04 (0.88 to 1.23)</td>
<td>Very low</td>
</tr>
<tr>
<td><strong>Conflicting findings from studies</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Neonatal mortality</td>
<td>0.0% (14.7%)</td>
<td>0.1% (14.7%)</td>
<td>1 fewer per 1000 live births (from 1 fewer to 2 more)</td>
<td>NC</td>
<td>Very low</td>
</tr>
<tr>
<td></td>
<td>0.17% (7.9%)</td>
<td>0.07% (7.9%)</td>
<td>1 more per 1000 live births (from 1 more to 2 more)</td>
<td>RR 2.4 (2.20 to 2.65)</td>
<td>Very low</td>
</tr>
<tr>
<td>Apgar score at 5 mins &lt; 7</td>
<td>0.0% (14.7%)</td>
<td>0.5% (14.7%)</td>
<td>5 fewer per 1000 (from 5 fewer to 1 fewer)</td>
<td>NC</td>
<td>Very low</td>
</tr>
<tr>
<td></td>
<td>0.6% (35%)</td>
<td>1.2% (35%)</td>
<td>6 fewer per 1000 (from 9 fewer to 157 more)</td>
<td>RR 0.44 (0.07 to 2.51)</td>
<td>Very low</td>
</tr>
</tbody>
</table>
1.6 Key research recommendations

<table>
<thead>
<tr>
<th>Number</th>
<th>Research Recommendation</th>
<th>See section</th>
</tr>
</thead>
<tbody>
<tr>
<td>RR 1</td>
<td>Risks and benefits of CS</td>
<td>4.2</td>
</tr>
</tbody>
</table>

What are the medium- to long-term risks and benefits to women and their babies of planned CS compared with planned vaginal birth?

The main focus would be the outcomes in women which could be measured at 1 year (medium term) and 5-10 years (long term). These outcomes could include:

- urinary dysfunction
- gastrointestinal dysfunction
- dyspareunia
- breastfeeding
- psychological health.

Infant outcomes could include medical problems, especially ongoing respiratory and neurological problems.

Why this is important

Morbidities arising intraoperatively or in the days after a caesarean section have been reasonably well described in the literature. Much less is known, however, about physical and emotional outcome measures in the longer term.

The Confidential Enquiries into Maternal Death in the UK, most recently published as ‘Saving mothers’ lives 2006-2008’ (Cantwell R. et al., 2011), devote a significant proportion of their work to investigating ‘late’ causes of maternal death. These include events arising in the medium term, namely, up to 1 year after a woman has given birth, many of which originate from the preceding pregnancy. The infectious, psychiatric and other conditions arising in or related to pregnancy do not always cause death but are responsible for arguably a greater burden of morbidity in the medium and long term, long after the pregnancy is over.

To provide more meaningful information to women when they are choosing their mode of birth, there is a pressing need to document medium- to long-term outcomes in women and their babies after a planned CS or a planned vaginal birth. First, it should be possible to gather data using standardised questions (traditional paper-based questionnaires and face-to-face interviews) about maternal septic morbidities and emotional wellbeing up to 1 year after a planned CS in a population of women who have consented for follow-up. Internet-based questionnaires could also be devised, to achieve the high response rates required for a full interpretation of the data. Similarly, it would be important to collect high-quality data on infant morbidities after a planned CS compared with a planned vaginal birth. A long term morbidity evaluation (between 5 and
10 years after the CS) would use similar methodology but assess symptoms related to urinary and gastrointestinal function.

**Maternal request for CS**

What support or psychological interventions would be appropriate for women who have a fear of vaginal childbirth and request a CS?

Interventions for evaluation could include:
- support from a named member of the maternity team
- continuity of carer
- formal counselling
- cognitive behavioural therapy.

Outcomes could include:
- mode of birth planned at term
- psychological outcomes (postnatal depression, post-traumatic stress disorder, self-esteem, mother-infant bonding)
- breastfeeding.

**Why this is important**

Fear of vaginal childbirth may stem from:
- fear of damage to the maternal pelvic floor
- damage to the baby during childbirth
- self-doubt on the ability to physically achieve vaginal birth
- previous childbirth experience
- unresolved issues related to the genital area.

Currently there is a wide variation in practice and limited resources lead to limited availability of effective interventions. Interventions that may be appropriate include:
- antenatal clinics dedicated to providing care for women with no obstetric indications who request a CS
- referral to a psychologist or a mental health professional
- referral to an obstetric anaesthetist
- intensive midwifery support.

Continuity of healthcare professional support from the antenatal to the intrapartum periods and ‘one to one’ midwifery care during labour are also often lacking and may make a difference to women who are anxious or afraid.

All of these interventions have different resource implications and there is no clear evidence to suggest that any are of benefit. The proposed research would compare in a randomised controlled trial two or more of these interventions in women requesting a CS. In the absence of any evidence, there is a case for comparing these interventions with routine antenatal care (that is, no special intervention).

This research is relevant because it would help to guide the optimal use of these limited resources and future guideline recommendations.
Decision-to-delivery interval for unplanned CS

What factors influence the decision-to-delivery interval when there is a category 1 level of urgency for CS?

Factors to be investigated could include:

- staff grade/level of experience
- skill mix within the multidisciplinary team
- task allocation
- methods of communication
- time of day
- availability of ongoing staff training about emergency procedures and levels of attendance.

The research could be conducted using simulation methods and video observation to determine what factors influence the decision-to-delivery interval for category 1 CS. The videos could also be used to train staff.

Why this is important

‘Crash’ CS is a psychologically traumatic event for women and their partners and is also stressful for clinical staff. Staff and resources may have to be obtained from other areas of clinical care. This should be undertaken as efficiently and effectively as possible, minimising anxiety and ensuring the safety of the mother and her baby.

For category 1 CS there is a recognised urgency to deliver as quickly as is reasonably possible. The majority of research in this area is quantitative and looks at the impact of the decision-to-delivery interval on various aspects of fetal and maternal outcomes rather than the interplay of factors that can affect this time period itself. Much of this evidence is retrospective. Although some work has been conducted in the UK to examine where the systematic delays lie and how to avoid them (Tuffnell et al., 2001), more work is needed to determine how to optimise the decision-to-delivery interval. This work should use qualitative as well as quantitative research methods to assess which factors influence the decision-to-delivery interval for a category 1 CS. Evaluation of these factors could be used to inform future NICE guidance, for example specific guidance for management of category 1 CS. Such information could also be used by hospitals for maternity services planning and at a team level would assist with audit and ongoing evaluation and training of the multidisciplinary team.

A large amount of NHS and other state funding is used to provide continuing care for infants who are disabled as a result of birth asphyxia and in providing lifelong support for the child and their family. In addition, large sums of public money are spent on litigation and compensation in some of these cases through the Clinical Negligence Scheme for Trusts (CNST). If research helped to minimise the impact of birth asphyxia this would reduce the costs of continuing care to the state and the burden to the child, their family and the wider community.

More realistic and more relevant expectations for the decision-to-
delivery interval based on evidence would inform debate in the legal system and may help to reduce the cost to the state of related litigation.

RR 29

A prospective study to determine whether the decision-to-delivery interval has an impact on maternal and neonatal outcomes when there is a category 2 level of urgency for CS.

Important primary outcomes would be

- fetal wellbeing (such as cord blood gases, Apgar score at 5 minutes, hypoxic encephalopathy, neonatal respiratory problems, unanticipated admission to neonatal intensive care unit (NICU), duration of stay in the NICU)
- maternal wellbeing (such as haemoglobin levels on day 2, need for blood transfusion, duration of hospital stay controlled for prolonged neonatal stay and general health/wellbeing).

Valuable secondary outcomes could include:

- fetal trauma at delivery
- iatrogenic maternal bladder or bowel injury
- postoperative maternal infectious morbidity
- establishment of breast-feeding
- psychological outcomes for women, such as the development of postnatal depression/post-traumatic stress disorder.

Why this is important

This research is important to inform the ongoing debate about the management of category 2 CS. The ‘continuum of risk’ in this setting has been recognised. However, the majority of work in this area, looking at maternal and fetal outcomes, generally considers unplanned caesarean sections as a whole group without making any distinction between degrees of urgency. Furthermore much of this work is retrospective. The majority of women who undergo intrapartum CS fall into the category 2 level of urgency (Thomas et al., 2001) and therefore specific information for this group could affect and benefit many women and contribute to the delivery of equity of care.

Delay in delivery with a compromised fetus may result in major and long-term harm including cerebral palsy and other major long-term disability. The immediate and long-term effect on a family of the birth of a baby requiring life-long specialised care and support is enormous. If such harm could be avoided by appropriate haste this would be an important improvement in outcome. However, if such haste is of no benefit then any related risk of adverse maternal outcome needs to be minimised.

A large amount of NHS and other state funding is used to provide continuing care for infants who are disabled as a result of delay in delivery and in providing lifelong support for the child and their family. In addition, large sums of public money are spent on litigation and compensation in some of these cases through the Clinical Negligence Scheme for Trusts (CNST). If research helped to minimise the impact of delay in delivery this would reduce the costs of continuing care to the state and the burden to the child, their family and the wider community.
More realistic and more relevant expectations for the decision-to-delivery interval based on evidence would inform debate within the legal system and may help to reduce the cost to the state of related litigation.

RR 30 Repeat of the National Caesarean Section Sentinel Audit

The original CS guideline included a set of ‘auditable standards’. It would be a straightforward task to produce an updated set of auditable standards based on the important topics covered in the updated guideline. These could include:

- consent
- indications (including maternal request)
- procedural aspects
- maternal and fetal outcomes.

Many of the outcomes documented in a new CS audit would relate directly to recommendations in this CS guideline update. Researchers may also want to consider categorising different reasons underlying maternal request for CS such as previous poor childbirth experience, longstanding fear of childbirth, belief that CS is safer for the baby etc.

An additional useful feature of the audit would be to record key related data, such as the proportion of CS for a breech presentation that had an attempted external cephalic version.

Why this is important

During the 10 years since the National Caesarean Section Sentinel Audit was undertaken (2000–2001), many of the findings may have changed significantly. The audit examined who was having a CS and why, as well as the views of women having babies and the obstetricians looking after them. The audit found that a 20% CS rate was considered too high by 51% of obstetricians. UK CS rates now average about 25%.

A repeat of the CS Sentinel Audit would reveal any changes in indications and the views of women and obstetricians. The current literature does not adequately address the issue of maternal request for CS and this is one aspect the audit may address. Women’s views on maternal request for CS for when there are no obstetric indications are particularly relevant. Such requests may be on the rise and the reasons are not always clearly expressed or documented.

The methodology of the audit is established, making a repeat feasible. This should be given high priority because the benefit to the NHS would be significant.
# 1.7 Research recommendations

<table>
<thead>
<tr>
<th>Number</th>
<th>Research Recommendation</th>
<th>See section</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Risks and benefits of CS</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>RR 1</td>
<td>What are the medium- to long-term risks and benefits to women and their babies of planned CS compared with planned vaginal birth?</td>
<td>4.2</td>
</tr>
<tr>
<td>RR 2</td>
<td>Further evaluation is needed to determine the impact of demographic and clinical factors (such as ethnic group, increase in body mass index) and attitudinal factors on CS rates.</td>
<td>4.2</td>
</tr>
<tr>
<td><strong>Breech presentation</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>RR 3</td>
<td>Further research is needed to determine the effect of caesarean section compared with vaginal birth for women with:</td>
<td>5.1</td>
</tr>
<tr>
<td></td>
<td>• preterm breech</td>
<td></td>
</tr>
<tr>
<td></td>
<td>• a breech presentation that is diagnosed in the second stage of labour</td>
<td></td>
</tr>
<tr>
<td><strong>Multiple pregnancy</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>RR 4</td>
<td>RCTs are needed to evaluate the benefits and risks to mothers and babies of CS for delivery of twin and triplet pregnancies</td>
<td>5.2</td>
</tr>
<tr>
<td><strong>Preterm birth and CS</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>RR 5</td>
<td>RCTs are needed to evaluate the impact of CS on the benefits and risks to mothers and babies born preterm.</td>
<td>5.3</td>
</tr>
<tr>
<td><strong>Small for gestational age</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>RR 6</td>
<td>RCT evidence is needed to determine the effect of planned CS on neonatal mortality and morbidity for ‘small for gestational age’ babies.</td>
<td>5.4</td>
</tr>
<tr>
<td><strong>Morbidly adherent placenta</strong></td>
<td></td>
<td></td>
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<tr>
<td>RR 7</td>
<td>How accurate is 3D ultrasound compared with 2D ultrasound or MRI scanning for diagnosing morbidly adherent placenta?</td>
<td>5.6</td>
</tr>
<tr>
<td>RR 8</td>
<td>What is the effectiveness of procoagulant agents (such as recombinant factor VIIa, beriplex, tranexamic acid, fibrinogen concentrate) in reducing blood loss in women with morbidly adherent placenta?</td>
<td>5.6</td>
</tr>
<tr>
<td>RR 9</td>
<td>What is the effectiveness of point of care testing for haematological indices in women with an established postpartum haemorrhage and in cases of morbidly adherent placenta in reducing maternal morbidity?</td>
<td>5.6</td>
</tr>
<tr>
<td>RR 10</td>
<td>What is the effectiveness of the components of the package of care for morbidly adherent placenta such as imaging techniques (e.g. interventional radiology including balloon catheters), stenting of ureters, removal of the placenta, and cell salvage in reducing</td>
<td>5.6</td>
</tr>
</tbody>
</table>
morbidity associated with maternal blood loss?

RR 11 What is the appropriate gestational age of elective birth for babies of women with a morbidly adherent placenta

RR 12 What is the effectiveness of performing an elective hysterectomy to reduce morbidity associated with blood loss in women with morbidly adherent placenta?

Mother-to-child transmission of maternal infections

RR 13 RCTs are needed to evaluate the effect of planned CS in addition to immunoglobulin and vaccination on MTCT of hepatitis B.

RR 14 RCTs are needed to determine whether planned CS should be offered to prevent MTCT of HSV to women with recurrence of HSV at birth and in women in whom the primary HSV infection occurs in the first trimester of pregnancy.

Maternal request for CS

RR 15 What support or psychological interventions would be appropriate for women who have a fear of vaginal childbirth and request a CS?

RR 16 Medium to long term quality of life study comparing psychological and physical outcomes in women who have had a requested and given birth by CS compared with women who plan a vaginal birth.

RR 17 Qualitative and quantitative research should be carried out to look at the reasons that lead to pregnant women’s request for CS.

RR 18 The effect of counselling and other interventions such as second opinion and provision of support on the likelihood of CS for women who express a preference for CS need further evaluation.

Place of birth

RR 19 RCTs comparing planned birth in a stand-alone birthing centre to birth in conventional maternity facilities or midwifery led units.

RR 20 Qualitative research is needed to explore women’s opinions on place of birth and the impact of place of birth on their birth experiences.

RR 21 Further RCTs are needed to determine the effect of ‘delayed admission in labour’ on the likelihood of CS.

Factors reducing the likelihood of CS

RR 22 RCT evidence is needed to determine the impact of partograms based on different curves of labour on CS rates and morbidity outcomes.

No influence on likelihood of CS

RR 23 RCT evidence is required to evaluate the effect of parenteral analgesia (intramuscular and intravenous morphine based analgesia) used during childbirth on the likelihood of CS.

RR 24 RCTs are needed to establish the safety and efficacy of complementary therapies used during labour.

‘Failure to progress’ in labour and CS

RR 25 More RCTs are required to determine the effect of oxytocin augmentation as single interventions or as part of a package of interventions (such as “active management of labour”) on the likelihood of CS and other outcomes including women’s...
satisfaction with care.

RR 26 Further research on the short and longer term health impacts of CS during the second stage compared to operative vaginal delivery are needed.

Eating during labour

RR 27 RCTs that evaluate the effects of eating during labour compared with restricting intake on labour outcomes are needed. Cohort or case control studies on the risk factors for aspiration and other morbidities for women having CS are needed.

Decision-to delivery-interval for unplanned CS

RR 28 What factors influence the decision-to-delivery interval when there is a category 1 level of urgency for CS?

RR 29 A prospective study to determine whether the decision-to-delivery interval has an impact on maternal and neonatal outcomes when there is a category 2 level of urgency for CS.

RR 30 Repeat of the National Caesarean Section Sentinel Audit

Surgical techniques for CS

RR 31 RCTs are required to determine the effectiveness of adhesive drapes at CS in reducing blood spillage and cross infection and improving safety for staff in the operating room.

RR 32 RCTs are needed to evaluate the effectiveness of incisions made with diathermy compared with surgical knife in terms of operating time, wound infection, wound tensile strength, cosmetic appearance and women's satisfaction with the experience.

RR 33 RCTs are needed to determine the effect of delayed cord clamping on neonatal outcomes including transient tachypnoea of the newborn and risk of maternal fetal transfusion in rhesus negative women for term and preterm births.

RR 34 RCTs are required to determine the effectiveness of mass closure compared to layered closure of the abdominal wall incision at CS particularly for transverse abdominal incisions.

RR 35 Research is required to assess the effect of the various surgical techniques for CS on future surgery such as repeat CS and the incidence of complications during future surgery such as hysterectomy and urogynaecological procedures.

RR 36 More RCTs are needed to determine the effect of wound drainage of postoperative morbidity especially in women more at risk of this outcome such as obese women.

RR 37 More RCTs are needed to determine the effect of staples compared to subcuticular sutures for skin closure at CS on postoperative pain, cosmetic appearance and removal of sutures and staples.

RR 38 What is the most effective antibiotic to prevent maternal infectious morbidity post-CS when given prior to incision

RR 39 What is the physical, psychological and social impact of maternal infectious morbidity post-CS?

RR 40 More evaluation of interventions such as seeing baby born via a lowered screen; music playing in theatre; silence in theatre so mother's voice is the first baby hears and lowering the lights in
theatre during CS are needed.

**Neonatal encephalopathy and cerebral palsy**

**RR 41** Further evaluation of the long and short term risks and benefits of CS compared with vaginal birth for babies is required. 8.2

**Thermal care for babies born by CS**

**RR 42** Research is required to establish the thermal care requirements for babies born by CS. 8.4

**Pain management after CS**

**RR 43** Further research is needed to determine the effect of wound infiltration with local anaesthetic at CS on the need for post-CS analgesia. 9.2

**Respiratory physiotherapy after CS**

**RR 44** Research is needed to establish the thermal care requirements for babies born by CS. 9.5

**Debriefing for women after CS**

**RR 45** More RCT evidence is required to determine the effect of midwifery-led debriefing following CS. 9.6

**Pregnancy and childbirth after CS**

**RR 46** A comparison of the long term psychological and physical outcomes between women who have chosen and/or been advised towards a VBAC or a planned repeat CS. 11.2

**RR 47** An evaluation of the effectiveness of continuity of carer on the proportion of women planning and achieving a VBAC, and the short and long term psychological and physical outcomes of women following a planned VBAC. 11.2

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### 1.8 Other versions of the guideline

Details about the other versions of the guideline will be included here once the guideline is published.

### 1.9 Schedule for updating the guideline

Clinical guidelines commissioned by NICE are published with a review date 3 years from the date of publication. Reviewing may begin before 3 years have elapsed if significant evidence that affects guideline recommendations is identified sooner.
2 Introduction

2.1 Caesarean Section

Between 20-25% of births in the UK are undertaken by caesarean section (CS). The indications for the procedure vary. This evidence based guideline has been developed to help ensure consistency and quality of care experienced by women who:

- have had a CS in the past and are now pregnant again or,
- have a clinical indication for a CS or,
- are considering a CS in the absence of a clinical indication.

It provides evidence-based information for healthcare professionals and women about:

- the risks and benefits of both planned and unplanned CS
- specific indications for CS
- effective management strategies to avoid CS
- anaesthetic and surgical aspects of care
- interventions to reduce morbidity from CS
- organisational and environmental factors that affect CS rates.

For the update the following specific topics have been addressed:

- the risks and benefits of planned CS compared with planned vaginal birth
- care of women considered at risk of a morbidly adherent placenta
- appropriate care and choices for women who are HIV positive
- care of women requesting a CS in the absence of a clinical indication
- audit standards with respect to the decision-to-delivery interval
- timing of the administration of antibiotics for CS
- appropriate care and choices for those women who have previously had a CS

This guideline links with other relevant NICE guidelines such as Antenatal care (NCC-WCH, 2008), Induction of labour (NCC-WCH, 2008), Intrapartum care (NCC-WCH, 2007), Diabetes in pregnancy (NCC-WCH, 2008), Hypertension in pregnancy (NCC-WCH, 2010), and Postnatal care (NCC-PC, 2006), the published NICE interventional procedure Intra-operative blood salvage in obstetrics (NICE, 2005) the findings of the NSCSA\(^5\) and the National Service Framework for Children, Young People and Maternity Services: Maternity services (Department of Health, 2004).

In 1980, the CS rate in England was 9%, increasing to 13% in 1992 (Treffers PE & Pel M, 1993), 21% in 2000 (RCOG, 2001), 23% in 2004 (Quick Stats, 2005), and 24.8%\(^5\) in 2009 (Department of Health, 2009). Similar increases have been seen in all developed countries though the absolute rates vary; for example, they are about 14% in Nordic countries and over 40% in Italy (Quick Stats, 2005). The reverse is seen in developing countries where CS rates are generally less than 5% (Buekens et al. 2003). There are a number of possible reasons for the increased rates in developed countries including changes in socio-demographic factors, clinical practices and the attitudes of professionals.

\(^5\) This is based on all deliveries taking place in NHS hospitals (in England), but excludes home births and those taking place in independent sector hospitals.
and women to the procedure. Many of the factors contributing to CS rates are often poorly understood. The guideline has not sought to define acceptable CS rates. Rather the purpose of this guideline is to enable clinicians to give appropriate research-based advice to women and their families. This will enable the woman to make a properly informed decision.

### 2.2 For whom is this guideline intended

This guideline is of relevance to those who work in or use the National Health Service in England and Wales:

- primary, community and secondary healthcare professionals who are involved in the care of women during pregnancy, birth and in the postnatal period who may need or have had a CS
- those with responsibilities for commissioning and planning health services such as Primary Care Trust commissioners (UK), Welsh Assembly Government officers
- public health and trust managers
- pregnant women, their families, birth supporters and other carers.

### 2.3 Related NICE guidance

- Multiple pregnancy. Currently in development. Due to publish in September 2011
3 Guideline development methodology

3.1 Original (2004) methodology

The guideline was commissioned by NICE and developed in accordance with the guideline development process outlined in The Guideline Development Process – Information for National Collaborating Centres and Guideline Development Groups.13

Literature search strategy

The aim of the literature review was to identify and synthesise relevant evidence within the published literature, in order to answer specific clinical questions. Searches were performed using generic and specially developed filters, relevant medical subject heading terms and free-text terms. Details of all literature searches are available on application to the NCC-WCH.

The National Guidelines Clearinghouse database, the Turning Research into Practice database, and the Organising Medical Networked Information service on the Internet were searched for guidelines produced by other development groups. The reference lists in these guidelines were checked against our searches to identify any missing evidence.

Searches were carried out for each topic of interest. The Cochrane Library (up to Issue 4, 2003) was searched to identify systematic reviews (with or without meta-analyses) of randomised controlled (clinical) trials (RCTs) and individual RCTs. The electronic databases MEDLINE (Ovid version for the period January 1966 to January 2004), EMBASE (Ovid version for the period between 1988 to January 2004), the Cumulative Index to Nursing and Allied Health Literature, the British Nursing Index and PsychInfo were also searched, as was the Database of Abstracts and Reviews of Effectiveness.

There was no systematic attempt to search the ‘grey literature’ (conferences, abstracts, theses and unpublished trials). A preliminary scrutiny of titles and abstracts was undertaken and full papers were obtained if the research question addressed the Guideline Development Group’s question relevant to the topic. Following a further review of the full version of the study, articles that did not address the Group’s question were excluded. Studies that did not report relevant outcomes were also excluded.

Submitted evidence from stakeholders was included where the evidence was relevant to the Group’s clinical question and was of equivalent or better quality than the research identified in the literature searches.

The economic evidence presented in this guideline is not a systematic review of all the economic evidence around CS, but a review of evidence relating to specific aspects of CS in addition to the databases listed above, the Health Economic Evaluations Database and the NHS Economic Evaluations Database were searched for relevant economic studies.

The search strategies were designed to find any economic study related to CS. Relevant references in the bibliographies of reviewed papers were also identified. Abstracts and database reviews of papers found were reviewed by the health economists and were excluded if they appeared not to contain any cost data relevant to the UK setting or did not relate to the precise topic or question being considered. Studies were included if they focused on the appropriate clinical question and were generalisable to the England and Wales setting. The review of the evidence included cost-effectiveness studies, cost-consequence studies (cost of present and future costs only) and high quality systematic reviews of the evidence (see below).
**Clinical effectiveness**

For all subject areas, evidence from the study designs least subject to bias was included. Where possible, the highest levels of evidence were used, but all papers were reviewed using established guides.\(^{14-20}\) Published systematic reviews or meta-analyses were used where available. For subject areas where neither was available, other appropriate experimental or observational studies were sought.

Identified articles were assessed methodologically and the best available evidence was used to form and support the recommendations. The highest level of evidence was selected for each clinical question. The retrieved evidence was graded according to the evidence-level structure shown in Table 3.1.

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<table>
<thead>
<tr>
<th>Level</th>
<th>Evidence</th>
</tr>
</thead>
<tbody>
<tr>
<td>1a</td>
<td>Systematic review or meta-analysis of randomised controlled trials</td>
</tr>
<tr>
<td>1b</td>
<td>At least one randomised controlled trial</td>
</tr>
<tr>
<td>2a</td>
<td>At least one well-designed controlled study without randomisation</td>
</tr>
<tr>
<td>2b</td>
<td>At least one well-designed quasi-experimental study, such as a cohort study</td>
</tr>
<tr>
<td>3</td>
<td>Well-designed non-experimental descriptive studies, such as comparative studies, correlation studies, case–control studies, and case series</td>
</tr>
<tr>
<td>4</td>
<td>Expert committee reports, or opinions and/or clinical experience of respected authorities</td>
</tr>
</tbody>
</table>

The clinical question dictated the highest level of evidence that could be sought. For issues of therapy or treatment the highest possible level of evidence was a meta-analysis of RCTs or an individual RCT.

For issues of prognosis, a cohort study was the best possible level of evidence. This equates to a grade B recommendation (see below). However, this should not be interpreted as an inferior grade of recommendation because it represents the highest level of evidence attainable for that type of clinical question.

For diagnostic tests, test evaluation studies examining the performance of the test were used if the efficacy of the test was required, but where an evaluation of the effectiveness of the test in the management and outcome was required, evidence from RCTs or cohort studies was sought. For questions about women’s beliefs, attitudes and experiences of childbirth and CS, qualitative research was reviewed.

All retrieved articles were appraised methodologically using established guides. Where appropriate, if a systematic review, meta-analysis or RCT existed in relation to a topic, studies of a weaker design were excluded.

The evidence was synthesised using qualitative methods. These involved summarising the content of identified papers in the form of evidence tables and agreeing brief statements that accurately reflected the relevant evidence. Quantitative synthesis (meta-analysis) was performed where appropriate. Meta-analyses based on dichotomous outcomes are presented as relative risks with 95% confidence intervals.

For the purposes of this guideline, data are presented as absolute risks, relative risks or odds ratios where relevant (i.e. in RCTs and cohort studies). Where the data are statistically significant they are also presented as numbers needed to treat (for beneficial outcomes) or numbers need to harm (for adverse effects of treatment) if relevant.
Health economics

The purpose of including economic evidence in a clinical guideline is to allow recommendations to be made not just on the clinical effectiveness of different forms of care, but also on their cost effectiveness. The aim is to produce guidance that uses scarce health service resources efficiently, that is providing the best possible care within resource constraints.

There is economic literature that has considered the economic costs and consequences of different modes of birth. The economic evidence is focused around the cost of CS compared to vaginal birth. The economic evidence presented in this guideline is not a systematic review of all the economic evidence around CS. Specific topics were considered where it was thought that economic evidence would help them to inform decision making.

Topics for economic analysis were selected on the following basis by the guideline development group:

- Does the proposed topic have major resource implications?
- Is there a change of policy involved?
- Are there sufficient data of adequate quality to allow useful review or modelling?
- Is there a lack of consensus amongst clinicians?
- Is there a particular area with a large amount of uncertainty?

Where the above answers are "yes", this indicated that further economic analysis including modelling was more likely to be useful.

A simple economic model was developed for each of the specific topic areas for which the economic evidence was reviewed, in order to present the guideline development group with a coherent picture of the costs and consequences of the decisions based on the clinical and economic evidence. The health economist undertook the literature review in these specific areas and obtained cost data considered to be the closest to current UK opportunity cost (the value of the resources used, rather than the price or charge). The criteria for assessing the economic papers was based on that developed by Drummond et al (1997) and the format of the abstract follows that of the NHS Economic Evaluation Database (NHS EED) managed by the NHS Centre for Reviews and Dissemination (http://nhscr.d.york.ac.uk/).

Health economics evidence was available for the following areas:

- external cephalic version for breech presentation at term,
- CS in the management of women with breech presentation,
- HIV/AIDS,
- herpes simplex virus
- vaginal birth after CS
- maternal request for CS
- use of antibiotics at CS
- intrathecal diamorphine.

The economic evidence is based not only on the economic literature, but is also consistent with the clinical effectiveness evidence presented in the guideline.

Forming and grading recommendations

The Guideline Development Group was presented with the summaries (text and evidence tables) of the best available research evidence to answer each clinical question. Recommendations were based on, and explicitly linked to, the evidence that supported them. Where possible, the Group worked on an informal consensus basis. Formal consensus methods (the nominal group technique) were employed when required (e.g. grading recommendations and agreeing audit criteria).
The strength of evidence corresponding to each level of recommendation is shown in Table 3.2. The grading of recommendations follows that outlined in the Health Technology Assessment ‘How to develop cost conscious guidelines’.

Summary results are presented in the guideline text. More detailed results and other data are presented in the relevant evidence tables.

### Table 3.2 Grading of recommendations

<table>
<thead>
<tr>
<th>Grade</th>
<th>Strength of evidence</th>
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</thead>
<tbody>
<tr>
<td>A</td>
<td>Based directly based on level 1 evidence</td>
</tr>
<tr>
<td>B</td>
<td>Based directly on level 2 evidence or extrapolated from level 1 evidence</td>
</tr>
<tr>
<td>C</td>
<td>Based directly on level 3 evidence or extrapolated from level 1 or level 2 evidence</td>
</tr>
<tr>
<td>D</td>
<td>Based directly on level 4 evidence or extrapolated from level 1, level 2 or level 3 evidence</td>
</tr>
<tr>
<td>GPP</td>
<td>Good practice point based on the view of the Guideline Development Group</td>
</tr>
<tr>
<td>NICE TA</td>
<td>Recommendation taken from a NICE Technology Appraisal</td>
</tr>
</tbody>
</table>

### External review

The guideline has been developed in accordance with the NICE guideline development process. This has included the opportunity for registered stakeholders to comment on the scope of the guideline, the first draft of the full and summary guidelines and the second draft of all versions of the guideline.

In addition the drafts were reviewed by an independent Guideline Review Panel and the Patient Involvement Unit established by NICE. The summary of recommendations was reviewed the NICE Executive.

The comments made by the stakeholders, peer reviewers, the Guideline Review Panel and NICE were collated and presented anonymously for consideration by the Guideline Development Group. All comments were considered systematically by the Guideline Development Group and the resulting actions and responses were recorded.

### 3.2 Methodology for 2011 update

This guidance was commissioned by NICE and developed in accordance with the guideline development process outlined in the 2009 edition of The Guidelines Manual (www.nice.org.uk/guidelinesmanual).

In accordance with NICE’s Equality Scheme, ethnic and cultural considerations and factors relating to disabilities have been considered by the GDG throughout the development process and specifically addressed in individual recommendations where relevant. Further information is available from: www.nice.org.uk/aboutnice/howwework/NICEEqualityScheme.jsp.

### Developing review questions and protocols and identifying evidence

The GDG formulated review questions based on the scope (see Appendix A) and prepared a protocol for each review question (see Appendix D). These formed the starting point for systematic reviews of relevant evidence. Published evidence was identified by applying systematic search strategies (see Appendix E) to the following databases: Medline, Medline In-Process, Embase, Cumulative Index to Nursing and Allied Health Literature (CINAHL), and three Cochrane databases (Cochrane Central Register of Controlled Trials, Cochrane Database of Systematic Reviews, and the Database of Abstracts of Reviews of Effects). Searches to identify economic studies were undertaken using Medline, Embase, the Cochrane Central Register of Controlled Trials, the NHS Economic Evaluation Database (NHS EED), and the Health Technology Assessment (HTA) database. Dates of searching and database coverage are given with the details of the search strategies in Appendix E.
Where appropriate, review questions were grouped together for searching. Animal studies were excluded from Medline and both Medline and Embase were limited to English-language studies only. Searches designed to update sections of the existing guideline were limited to 2003 onwards; searches for new review areas were not limited by date. Scottish Intercollegiate Guidelines Network (SIGN) search filters were used to identify particular study designs, such as randomised controlled trials (RCTs). There was no systematic attempt to search grey literature (conference abstracts, theses or unpublished trials), nor was hand searching of journals not indexed on the databases undertaken.

Towards the end of the guideline development process, the searches were updated and re-executed to include evidence published and indexed in the databases by 17th March 2011.

Reviewing and synthesising evidence
Evidence relating to clinical effectiveness was reviewed and synthesised according to the Grading of Recommendations Assessment, Development and Evaluation (GRADE) approach (see http://www.gradeworkinggroup.org/index.htm). In the GRADE approach, the quality of the evidence identified for each outcome listed in the review protocol is assessed according to the factors listed below, and an overall quality rating (high, moderate, low, or very low) is assigned by combining the ratings for the individual factors.

- Study design (as an indicator of intrinsic bias; this determines the initial quality rating)
- Limitations in the design or execution of the study (including concealment of allocation, blinding, loss to follow up; these can reduce the quality rating)
- Inconsistency of effects across studies (this can reduce the quality rating)
- Indirectness (the extent to which the available evidence fails to address the specific review question; this can reduce the quality rating)
- Imprecision (this can reduce the quality rating)
- Other considerations (including large magnitude of effect, evidence of a dose-response relationship, or confounding variables likely to have reduced the magnitude of an effect; these can increase the quality rating in observational studies, provided no downgrading for other features has occurred)

The type of review question determines the highest level of evidence that may be sought. For issues of therapy or treatment, the highest possible evidence level is a well-conducted systematic review or meta-analysis of RCTs, or an individual RCT. In the GRADE approach, a body of evidence for a given outcome based entirely on such studies has an initial quality rating of high, and this may be downgraded to moderate, low, or very low if factors listed above are not addressed adequately. For issues of prognosis, the highest possible level of evidence is a controlled observational study (a cohort study or case–control study), and a body of evidence for a particular outcome based on such studies would have an initial quality rating of low, which might be downgraded to very low or upgraded to moderate or high, depending on the factors listed above.

For each review question the highest available level of evidence was sought. Where appropriate, for example, if a systematic review, meta-analysis or RCT was identified to answer a question directly, studies of a weaker design were not considered. Where systematic reviews, meta-analyses and RCTs were not identified other appropriate experimental or observational studies were sought. For diagnostic tests, test evaluation studies examining the performance of the test were used if the accuracy of the test was required, but where an evaluation of the effectiveness of the test in the clinical management of the condition was required, evidence from RCTs or cohort studies was optimal. For studies evaluating the accuracy of a diagnostic test, sensitivity, specificity, positive predictive value (PPV), negative predictive value (NPV) and likelihood ratios for positive and negative test results (LR+ and LR−, respectively), were calculated or quoted where possible (see Table 3.3).

The GRADE system described above covers studies of treatment effectiveness. However, it is less well established for studies reporting accuracy of diagnostic tests. For such studies, NICE recommends using the Quality Assessment of Studies of Diagnostic Accuracy (QUADAS) methodology checklist to assess study quality (see the NICE guidelines manual).
Some studies were excluded from the guideline reviews after obtaining copies of the corresponding publications because they did not meet inclusion criteria specified by the GDG and recorded in the protocol (see Appendix D). The characteristics of each included study were summarised in evidence tables for each review question (see Appendix G). Where possible, dichotomous outcomes were presented as relative risks (RRs) or odds ratios (ORs) with 95% confidence intervals (CIs), and continuous outcomes were presented as mean differences with 95% CIs or standard deviations (SDs).

The body of evidence identified for each review question (or part of a review question) was presented in the form of a GRADE evidence profile summarising the quality of the evidence and the findings (pooled relative and absolute effect sizes and associated CIs). Summary GRADE tables have been reported in the main text, with the full GRADE evidence profiles reported in appendix H. Where possible, the body of evidence corresponding to each outcome specified in the review protocol was subjected to quantitative meta-analysis. In such cases, pooled effect sizes were presented as pooled risk ratios (RRs), pooled odds ratios (ORs), or weighted mean differences. By default, meta-analyses were conducted by fitting fixed effects models, but where statistically significant heterogeneity was identified random effects models were used. Where quantitative meta-analysis could not be undertaken (for example, because of heterogeneity in the included studies) the range of effect sizes reported in the included studies was presented.

| Table 3.3 '2 x 2' table for calculation of diagnostic accuracy parameters |
|---------------------------------|---------------------------------|------------------|
| Index test result positive      | Reference standard positive     | Reference standard negative |
|                                 | a (true positive)               | b (false positive) |
| Index test result negative      | c (false negative)              | d (true negative)  |
| Total                           | a+c                             | b+d               |
|                                 | a+b+c+d = N (total number of tests in study) |

**Incorporating health economics**

The aims of the health economic input to the guideline were to inform the GDG of potential economic issues relating to CS, and to ensure that recommendations represented a cost effective use of healthcare resources. Health economic evaluations aim to integrate data on benefits (ideally in terms of quality adjusted life years (QALYs)), harms and costs of different care options.

The GDG prioritised a number of review questions where it was thought that economic considerations would be particularly important in formulating recommendations. Systematic searches for published economic evidence were undertaken for these questions. For economic evaluations, no standard system of grading the quality of evidence exists and included papers were assessed using a quality assessment checklist based on good practice in economic evaluation. Reviews of the (very limited) relevant published health economic literature are presented alongside the clinical effectiveness reviews.

Health economic considerations were aided by original economic analysis undertaken as part of the development process. For this guideline the areas prioritised for economic analysis were as follows:

- Diagnosis of morbidly adherent placenta (see section 5.6 for summary and 13.2 for full details)
- Maternal request for CS (see section 5.9 for summary and 13.3 for full details)
- Vaginal birth after CS (see section 11.2 for summary and 13.4 for full details)
Evidence to recommendations

For each review question recommendations for clinical care were derived using, and linked explicitly to, the evidence that supported them. In the first instance, short clinical and, where appropriate, cost effectiveness evidence statements were drafted by the technical team which were presented alongside the evidence profiles and agreed by the GDG. Statements summarising the GDG’s interpretation of the evidence and any extrapolation from the evidence used to form recommendations were also prepared to ensure transparency in the decision-making process. The criteria used in moving from evidence to recommendations are summarised in Table 3.4.

In areas where no substantial clinical research evidence was identified, the GDG considered other evidence-based guidelines and consensus statements or used their collective experience to identify good practice. The health economics justification in areas of the guideline where the use of NHS resources (interventions) was considered was based on GDG consensus in relation to the likely cost effectiveness implications of the recommendations. The GDG also identified areas where evidence to answer their review questions was lacking and used this information to formulate recommendations for future research.

Towards the end of the guideline development process formal consensus methods incorporating anonymous voting were used to consider all the clinical care recommendations and research recommendations that had been drafted previously. The GDG identified ten ‘key priorities for implementation’ (key recommendations) and five high-priority research recommendations. The key priorities for implementation were those recommendations thought likely to have the biggest impact on clinical care and outcomes in the NHS as a whole. The priority research recommendations were selected in a similar way.

<table>
<thead>
<tr>
<th>Criterion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Relative value placed on the outcomes considered</td>
</tr>
<tr>
<td>Consideration of clinical benefits and harms</td>
</tr>
<tr>
<td>Consideration of net health benefits and resource use</td>
</tr>
<tr>
<td>Quality of the evidence</td>
</tr>
<tr>
<td>Other considerations (including equalities issues)</td>
</tr>
</tbody>
</table>

Stakeholder involvement

This paragraph will be completed following the stakeholder consultation.
4 Woman-centred care

4.1 Provision of information

In 1993, the Expert Maternity Group from the Department of Health (DH) released the Changing Childbirth report which made explicit the right of women to be involved in decisions regarding all aspects of their care during pregnancy and childbirth. One of the priorities of the report is to enable women to make informed decisions about their care. To make these decisions women require access to evidence-based information so that they can take part in discussions with caregivers about these decisions.

In a survey, pregnant women were asked their views about childbirth. This included questions about the information they wanted or had received. About 40% of women reported that they had sufficient information on the risks and benefits of CS, however almost 50% reported that they would have liked more information on reasons for CS, what to expect and the risks and benefits of CS [evidence level 3]. Information from RCTs on antenatal education suggests that the provision of information is often seen as inadequate by women [evidence level 3]. About 1 in 4 pregnant women will be delivered by CS (Department of Health, 2009) therefore when planning the birth of their baby women need information on both vaginal birth and CS. For women who experience a fear of childbirth, it is possible that building up confidence during pregnancy in her ability to give birth has the potential to influence her choices for the birth of her baby and the interventions she receives during birth.

Information leaflets

An RCT assessed the impact of evidence-based leaflets to promote informed decision making among pregnant women. The leaflets were designed to be used in a conscious and controlled way (i.e. not left in a rack at an antenatal clinic or GP office) and the information provided was based on results of systematic reviews of the best available evidence and they were peer-reviewed. No differences were detected in the proportion of women who reported that they had exercised informed choice or among those who reported an ‘active’ decision making role during antenatal care between the women that received these leaflets and those that did not. However, satisfaction with the amount of information received was higher among women who had received the leaflets [evidence level 1b]. Qualitative assessment within the RCT of the use of the leaflets found that their potential as decision aids was reduced due to competing demands within the clinical environment. Time pressures limited discussion and the hierarchical nature of the relationship between healthcare professionals and patients determined which ‘choices’ were available. This meant that women complied with their carer’s choice rather than making an informed decision [evidence level 3].

Antenatal education

A systematic review based on six RCTs (n = 1443) assessed the effects of antenatal education on knowledge acquisition, anxiety, sense of control, pain, support, breastfeeding, infant care abilities, and psychological and social adjustment. The largest RCT (n = 1275) examined an educational intervention to increase vaginal birth after CS. The other five RCTs (combined n = 168, range RCT n = 10 to 67) included more general educational interventions; however the methodological quality of these RCTs is uncertain as they do not report randomisation procedures, allocation concealment or accrual/loss of participants. None of the RCTs included labour and birth outcomes, anxiety, breastfeeding success, or general social support. The effects on knowledge acquisition and infant care competencies were measured but interpretation is difficult because of the size and methodological quality of the RCTs [evidence level 1b]

Recommendations
Number  Recommendation

1  Pregnant women should be offered evidence-based information and support to enable them to make informed decisions about childbirth. Addressing women’s views and concerns should be recognised as being integral to the decision-making process.  
   [C] [2004]

2  Give pregnant women evidence-based information about CS during the antenatal period, because about one in four women will have a CS. Include information about CS, such as:
   - indications for CS (such as presumed fetal compromise, ‘failure to progress’ in labour, breech presentation)
   - what the procedure involves
   - associated risks and benefits
   - implications for future pregnancies and birth after CS.  
   [GPP] [new 2011]

3  Communication and information should be provided in a form that is accessible to pregnant women, taking into account the information and cultural needs of minority communities and women whose first language is not English or who cannot read, together with the needs of women with disabilities or learning difficulties.  
   [GPP] [2004]

4.2  Planning mode of birth

Risks and benefits of planned CS compared with planned vaginal birth for women with an uncomplicated pregnancy

Clinical question

What are the risks and benefits of planned CS compared with planned vaginal birth for both women and babies?

Updating the risks and benefits tables

The original Caesarean Section guideline reviewed evidence for this question that included women with an obstetric indication for CS as well as women from a general population including both high and low obstetric risk. Data from all studies were presented together and summarised in 2 tables (labelled Tables 3.1a and 3.1b in the original guideline). Moreover, these previous tables included data drawn from both RCTs and observational studies. For the RCTs, findings were presented on an “intention to treat basis” truly representing planned CS vs. planned vaginal birth. For all observational studies, data were presented depending on actual mode of birth rather than planned mode of birth. For outcomes of interest where the number of events is very small this makes a very big difference to the interpretation of the findings. This is illustrated in Appendix K where for 3 outcomes it has been possible to compare the findings calculated on an intention to treat basis with those presented according to actual mode of birth. It should be noted that the sources of observational data presented in Tables 3.1a and 3.1b in the original guideline were not referenced nor detailed in chapter 3 but appear throughout the guideline under the relevant topic headings according to outcome. The deleted tables 3.1a and 3.1b from the original guideline have been annotated to provide extra information regarding the source of the data and the population studied (see Appendix K).

For this update to the guideline, the GDG wanted to focus specifically on women without antenatal complications (either medical or obstetric) in order that the review provided information for women who were requesting a CS in the absence of a clinical indication. It was also considered to be important to ensure included studies presented findings on an intention to treat basis thus reflecting the risks appropriate during the antenatal period when a woman is planning mode of birth. The updated reviews were undertaken following protocols that reflected this important difference (see Appendix D). Again both RCTs and observational studies were considered for inclusion.
The reasons outlined above mean the data presented in the updated tables for risks and benefits associated with planned CS compared with planned vaginal birth are quite different from the data that appeared in the original guideline. However, the updated tables provide a more valid estimation of these risks for a woman with an uncomplicated pregnancy planning mode of birth.

Overview of evidence

Nine studies were included in this review. One study was conducted in the UK (Homer et al., 2011), three studies were conducted in the USA (MacDorman et al., 2008; Geller et al., 2010a & 2010b), three in Canada (Dahlgren et al., 2009; Allen et al., 2006; Liu et al., 2007), one in France (Deneux-Tharaux et al., 2006) and one in Austria (Schindl et al., 2003). Three studies (MacDorman, et al., 2008; Geller et al., 2010a & 2010b) employed “intention to treat” analysis despite the actual route of delivery in cases and controls. Two population based studies (Liu et al., 2007; Dahlgren, et al., 2009) included women who gave birth by planned CS because of breech presentation, as an appropriate surrogate for “maternal request”. The comparison group in these studies was planned vaginal birth (not vaginal breech birth). One study (Schindl et al., 2003) reported “caesarean section on demand” for cases and women with “planned vaginal birth” for controls. Two studies (Allen et al., 2006; Deneux-Tharaux et al., 2006) compared the birth outcomes of women undergoing low risk pre-labour CS with those women intending to have a vaginal birth. Eight studies considered primary CS only; however, one study did not report excluding women with a prior CS (Deneux-Tharaux et al., 2006). Two studies reported excluding women who underwent induction of labour (Allen et al., 2006; Dahlgren, et al., 2009).

Four of the studies included in the review solely looked at nulliparous women (Allen et al., 2006; Dahlgren, et al., 2009; Geller et al., 2010a & 2010b), while the remainder also included multiparous women. Four studies reported neonatal outcomes, of which three (Dahlgren et al., 2009; Geller et al., 2010b; MacDorman et al., 2009) only included women giving birth at 37 weeks or later. The fourth study (Homer et al., 2011) included babies born before 37 weeks. More details of this study are given in the paragraph below, and for all studies in the evidence tables in Appendix G.

The UK study (Homer et al., 2011) comprised a population of extremely obese women (BMI ≥ 50) including both nulliparous women (34%) and multiparous women. It also included women with medical complications (e.g. pre-eclampsia, diabetes). This study was included in the review as it focussed on a population of women that was identified in the scope as being of particular importance and downgraded due to the indirectness of the sample population. Findings for this study are presented separately, and not included in the summary risk-benefits tables (4.5 and 4.6).

All outcomes reported in the included papers are recorded here in order to make the table as comprehensive as possible. The GDG felt it was not appropriate to choose which outcomes would be most important as this will vary from woman to woman. By presenting all the data available it is hoped each individual woman will receive the information she needs to make a fully informed decision.

Evidence Profile

For maternal outcomes:

Evidence was identified from nine studies. It is important to underline that the findings presented are for planned CS vs. planned vaginal birth i.e. the comparison is based upon what was planned antenatally not the actual mode of birth. This reflects the risks and benefits as they appear to a woman planning birth and take into account that when planning a vaginal birth, the woman may give birth by CS (usually carried out during labour). The number and percentage of unplanned CS carried out in the planned vaginal birth group for each study is also included in the table.

All reported outcomes from the included studies are presented in the table in order to provide information that is as comprehensive as possible. The GDG felt it was not appropriate to limit the outcomes included as there will be considerable variation between individual women as to what information they use when deciding mode of birth.

The quality of the evidence was low and very low for all studies included.
### Table 4.1 GRADE summary of findings for risks of planned CS compared with planned vaginal birth for women with an uncomplicated pregnancy and no previous CS (maternal outcomes)

<table>
<thead>
<tr>
<th>Number of studies</th>
<th>Percentage of unplanned CS in planned vaginal birth group</th>
<th>Number of women</th>
<th>Effect</th>
<th>Quality</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Planned CS</td>
<td>Planned vaginal birth</td>
<td>Relative (95% CI)</td>
</tr>
<tr>
<td>Maternal death</td>
<td></td>
<td>9/737* (cases/ controls)</td>
<td>49/9133* (cases/ controls)</td>
<td>OR 2.28* (1.11 to 4.65)</td>
</tr>
<tr>
<td>1 study</td>
<td>Of maternal deaths occurring in the planned vaginal birth group</td>
<td>13/49 (26.5%) were women who gave birth by unplanned CS</td>
<td>49/737* cases/controls</td>
<td></td>
</tr>
<tr>
<td>(Deneux-Tharaux et al., 2006)</td>
<td></td>
<td>9/737*</td>
<td>49/9133*</td>
<td>OR 2.28*</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0/1046 (0%)</td>
<td>0/38021 (0%)</td>
<td>NC</td>
</tr>
<tr>
<td>1 study</td>
<td></td>
<td>5580/38021 (14.7%)</td>
<td>0/1046 (0%)</td>
<td>NC</td>
</tr>
<tr>
<td>(Dahlgren et al. 2009)</td>
<td></td>
<td>0/1046</td>
<td>0/38021</td>
<td>NC</td>
</tr>
<tr>
<td>1 study</td>
<td></td>
<td>187,978/2,292,420 (8.2%)</td>
<td>41/2,292,420 (0.02 per 1000)</td>
<td>NC</td>
</tr>
<tr>
<td>(Liu et al., 2007)</td>
<td></td>
<td>0/46766</td>
<td>41/2,292,420</td>
<td>NC</td>
</tr>
</tbody>
</table>

### Perineal and abdominal pain (during birth) (range of scores: 0-10; Better indicated by lower values)

<table>
<thead>
<tr>
<th>Number of studies</th>
<th>Percentage of unplanned CS in planned vaginal birth group</th>
<th>Number of women</th>
<th>Effect</th>
<th>Quality</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Planned CS</td>
<td>Planned vaginal birth</td>
<td>Relative (95% CI)</td>
</tr>
<tr>
<td>1 study</td>
<td>93/903 (10.3%)</td>
<td>Median score = 1.0*</td>
<td>Median score = 7.3*</td>
<td>Significantly lower for planned CS</td>
</tr>
<tr>
<td>(Schindl et al., 2003)</td>
<td></td>
<td>Median score = 1.0*</td>
<td>Median score = 7.3*</td>
<td></td>
</tr>
</tbody>
</table>

### Perineal and abdominal pain (3 days postpartum) (range of scores: 0-10; Better indicated by lower values)

<table>
<thead>
<tr>
<th>Number of studies</th>
<th>Percentage of unplanned CS in planned vaginal birth group</th>
<th>Number of women</th>
<th>Effect</th>
<th>Quality</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Planned CS</td>
<td>Planned vaginal birth</td>
<td>Relative (95% CI)</td>
</tr>
<tr>
<td>1 study</td>
<td>93/903 (10.3%)</td>
<td>Median score = 4.5*</td>
<td>Median score = 5.2*</td>
<td>Significantly lower for planned CS</td>
</tr>
<tr>
<td>(Schindl et al., 2003)</td>
<td></td>
<td>Median score = 4.5*</td>
<td>Median score = 5.2*</td>
<td></td>
</tr>
</tbody>
</table>

### Perineal and abdominal pain (4 months postpartum) (range of scores: 0-10; Better indicated by lower values)

<table>
<thead>
<tr>
<th>Number of studies</th>
<th>Percentage of unplanned CS in planned vaginal birth group</th>
<th>Number of women</th>
<th>Effect</th>
<th>Quality</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Planned CS</td>
<td>Planned vaginal birth</td>
<td>Relative (95% CI)</td>
</tr>
<tr>
<td>1 study</td>
<td>93/903 (10.3%)</td>
<td>Median score = 0.0*</td>
<td>Median score = 0.17*</td>
<td>No significant difference between groups</td>
</tr>
<tr>
<td>(Schindl et al., 2003)</td>
<td></td>
<td>Median score = 0.0*</td>
<td>Median score = 0.17*</td>
<td></td>
</tr>
</tbody>
</table>

### Injury to bladder/ureter

<table>
<thead>
<tr>
<th>Number of studies</th>
<th>Percentage of unplanned CS in planned vaginal birth group</th>
<th>Number of women</th>
<th>Effect</th>
<th>Quality</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Planned CS</td>
<td>Planned vaginal birth</td>
<td>Relative (95% CI)</td>
</tr>
<tr>
<td>1 study</td>
<td>5580/38021 (14.7%)</td>
<td>0/1046 (0%)</td>
<td>53/38021 (0.14%)</td>
<td>NC</td>
</tr>
<tr>
<td>(Dahlgren et al., 2009)</td>
<td></td>
<td>0/1046</td>
<td>53/38021</td>
<td>NC</td>
</tr>
</tbody>
</table>

2011 Update
<table>
<thead>
<tr>
<th>Injury to cervix</th>
<th>1 study (Dahlgren et al., 2009)</th>
<th>5580/38021 (14.7%)</th>
<th>0/1046 (0%)</th>
<th>108/38021 (0.28%)</th>
<th>NC</th>
<th>3 fewer per 1000* (from 3 fewer to 1 more)</th>
<th>Very Low</th>
</tr>
</thead>
<tbody>
<tr>
<td>Injury to vagina</td>
<td>1 study (Dahlgren et al., 2009)</td>
<td>5580/38021 (14.7%)</td>
<td>0/1046 (0%)</td>
<td>213/38021 (0.56%)</td>
<td>NC</td>
<td>6 fewer per 1000* (from 6 fewer to 2 fewer)</td>
<td>Very Low</td>
</tr>
<tr>
<td>Iatrogenic surgical injury</td>
<td>1 study (Dahlgren et al., 2009)</td>
<td>5580/38021 (14.7%)</td>
<td>0/1046 (0%)</td>
<td>27/38021 (0.07%)</td>
<td>NC</td>
<td>7 fewer per 10,000 (from 10 fewer to 30 more)*</td>
<td>Very Low</td>
</tr>
<tr>
<td>Hysterectomy</td>
<td>1 study (Geller et al., 2010a)</td>
<td>1340/3868 (35%)</td>
<td>0.6%</td>
<td>0.1%</td>
<td>p = 0.13</td>
<td>5 more per 1000</td>
<td>Low</td>
</tr>
<tr>
<td></td>
<td>1 study (Dahlgren et al., 2009)</td>
<td>5580/38021 (14.7%)</td>
<td>1/1046 (0.1%)</td>
<td>4/38021 (0.01%)</td>
<td>RR 9.09* (1.36 to 60.33)</td>
<td>1 more per 1000* (from 0 more to 5 more)</td>
<td>Very Low</td>
</tr>
<tr>
<td></td>
<td>1 study (Liu et al., 2007)</td>
<td>187,978/2,292,420 (8.2%)</td>
<td>27/46,766* (0.6 per 1000)</td>
<td>367/2,292,420* (0.2 per 1000)</td>
<td>RR 3.60* (2.44 to 5.31)</td>
<td>41 more per 100,000* (from 23.6 more to 68 more)</td>
<td>Very Low</td>
</tr>
<tr>
<td>Hysterectomy due to postpartum haemorrhage</td>
<td>1 study (Liu et al., 2007)</td>
<td>187,978/2,292,420 (8.2%)</td>
<td>12/46766 (0.3 per 1000)</td>
<td>254/2,292,420 (0.1 per 1000)</td>
<td>RR 2.31* (1.30 to 4.09)</td>
<td>14 more per 100,000 (from 3 more to 33 more)</td>
<td>Very Low</td>
</tr>
<tr>
<td>Deep vein thrombosis</td>
<td>1 study (Dahlgren et al., 2009)</td>
<td>5580/38021 (14.7%)</td>
<td>0/1046 (0.0%)</td>
<td>3/38021 (0.007%)</td>
<td>NC</td>
<td>0.7 fewer per 1000* (from 0.2 fewer to 4 more)</td>
<td>Very Low</td>
</tr>
<tr>
<td></td>
<td>1 study (Liu et al., 2007)</td>
<td>187,978/2,292,420 (8.2%)</td>
<td>28/46,766 (0.6 per 1000)</td>
<td>623/2,292,420 (0.3 per 1000)</td>
<td>RR 2.20* (1.51 to 3.20)</td>
<td>32 more per 100,000* (from 14 more to 59 more)</td>
<td>Very Low</td>
</tr>
<tr>
<td>Pulmonary embolism</td>
<td>1 study 5580/38021</td>
<td>0/1046</td>
<td>1/38021</td>
<td>NC</td>
<td>0.2 fewer per</td>
<td>Very</td>
<td></td>
</tr>
</tbody>
</table>
### Blood transfusion

<table>
<thead>
<tr>
<th>Study</th>
<th>Blood Transfusion Rate</th>
<th>1/1000</th>
<th>1/1000*</th>
<th>Very Low</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 study (Geller et al., 2010a)</td>
<td>1340/3868 (35%)</td>
<td>3/180 (1.7%)</td>
<td>74/3868 (1.9%)</td>
<td>OR 0.87 (0.27 to 2.78)</td>
</tr>
<tr>
<td>1 study (Dahlgren et al., 2009)</td>
<td>5580/38021 (14.7%)</td>
<td>3/1046 (0.3%)</td>
<td>123/38021 (0.3%)</td>
<td>RR 0.89* (0.20 to 3.99)</td>
</tr>
<tr>
<td>1 study (Allen et al., 2006)</td>
<td>1480/17714 (8.3%)</td>
<td>2/721 (0.3%)</td>
<td>73/17714 (0.4%)</td>
<td>RR 0.7* (0.2 to 2.7)</td>
</tr>
<tr>
<td>1 study (Liu et al., 2007)</td>
<td>187,978/2,292,420 (8.2%)</td>
<td>11/46,766 (0.2 per 1000)</td>
<td>1500/2,292,420 (0.7 per 1000)</td>
<td>RR 0.20* (0.20 to 0.64)</td>
</tr>
</tbody>
</table>

### Early post-partum haemorrhage

<table>
<thead>
<tr>
<th>Study</th>
<th>Early Post-Partum Haemorrhage Rate</th>
<th>1/1000</th>
<th>1/1000*</th>
<th>Very Low</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 study (Geller et al., 2010a)</td>
<td>1340/3868 (35%)</td>
<td>2/180 (1.1%)</td>
<td>231/3868 (6.0%)</td>
<td>OR 0.23** (0.06 to 0.94)</td>
</tr>
<tr>
<td>1 study (Allen et al., 2006)</td>
<td>1480/17714 (8.3%)</td>
<td>28/721 (3.9%)</td>
<td>1098/17714 (6.2%)</td>
<td>RR 0.06* (0.4 to 0.9)</td>
</tr>
</tbody>
</table>

### Infection (wound and postpartum)

<table>
<thead>
<tr>
<th>Study</th>
<th>Infection Rate</th>
<th>1/1000</th>
<th>1/1000*</th>
<th>Very Low</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 study (Dahlgren et al., 2009)</td>
<td>5580/38021 (14.7%)</td>
<td>11/1046 (1.1%)</td>
<td>293/38021 (0.8%)</td>
<td>RR 1.36* (0.75 to 2.4)</td>
</tr>
<tr>
<td>1 study (Liu et al., 2007)</td>
<td>187,978/2,292,420 (8.2%)</td>
<td>281/46766 (6.0 per 1000)</td>
<td>4833/2,292,420 (2.1 per 1000)</td>
<td>RR 2.85* (2.52 to 3.21)</td>
</tr>
</tbody>
</table>

### Infection (wound)

<table>
<thead>
<tr>
<th>Study</th>
<th>Infection Rate</th>
<th>1/1000</th>
<th>1/1000*</th>
<th>Very Low</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 study (Geller et al., 2010a)</td>
<td>1340/3868 (35%)</td>
<td>0.01%</td>
<td>0.00%</td>
<td>p = 1.0</td>
</tr>
</tbody>
</table>
1 study  
(Allen et al., 2006)  
1480/17714  
(8.3%)  
11/721  
(1.5%)  
157/17714  
(0.9%)  
RR 1.7*  
(0.9 to 3.2)  
6 fewer per 1000*  
(from 1 fewer to 19 more)  
Very Low

**Anaesthetic complication**

1 study  
(Dahlgren et al., 2009)  
5580/38021  
(14.7%)  
4/1046  
(0.4%)  
117/38021  
(0.3%)  
RR 1.24*  
(0.34 to 4.59)  
1 more per 1000*  
(from 2 fewer to 11 more)  
Very Low

1 study  
(Liu et al., 2007)  
187,978/2,292,420  
(8.2%)  
247/46,766  
(5.3 per 1000)  
4793/2,292,420  
(2.1 per 1000)  
RR 2.5*  
(2.22 to 2.86)  
319 more per 100,000*  
(from 257 more to 389 more)  
Very Low

**Intraoperative trauma**

1 study  
(Allen et al., 2006)  
1480/17714  
(8.3%)  
1/721  
(0.1%)  
51/17714  
(0.3%)  
RR 0.5  
(0.1 to 3.5)  
1 fewer per 1000  
(from 3 fewer to 7 more)  
Very Low

**Uterine rupture**

1 study  
(Liu et al., 2007)  
187,978/2,292,420  
(8.2%)  
7/46,766  
(0.2 per 1000)  
660/2,292,420  
(0.3 per 1000)  
RR 0.51*  
(0.25 to 1.07)  
13 fewer per 100,000*  
(from 22 fewer to 2.2 more)  
Very Low

**Length of hospital stay (mean/days)**

1 study  
(Geller et al., 2010a)  
1340/3868  
(35%)  
3.2  
(SD 0.7)  
2.6  
(SD 1.1)  
Mean difference 1.58  
(1.27 to 2.17)  
Absolute mean difference 0.6 days longer  
Low

1 study  
(Liu et al., 2007)  
187,978/2,292,420  
(8.2%)  
3.96 days  
(SD 1.36)  
2.56 days  
(SD 1.36)  
Adjusted mean difference 1.47  
(1.46 to 1.49)  
Absolute mean difference 1.4 days longer  
Very Low

**Assisted ventilation or intubations**

1 study  
(Liu et al., 2007)  
187,978/2,292,420  
(8.2%)  
6/46766  
(0.1 per 1000)  
133/2,292,420  
(0.05 per 1000)  
RR 2.21*  
(0.99 to 4.90)  
7 more per 100,000*  
(from 0 fewer to 22 more)  
Very Low

**Acute renal failure**

1 study  
(Liu et al., 2007)  
187,978/2,292,420  
(8.2%)  
2/46,766  
(0.04 per 1000)  
45/2,292,420  
(0.01 per 1000)  
RR 2.17*  
(0.58 to 8.14)  
2 more per 100,000*  
(from 9 fewer to 13 more)  
Very Low

**Cardiac arrest**

1 study  
(Liu et al., 2007)  
187,978/2,292,420  
(8.2%)  
89/46,766  
(1.9 per 1000)  
887/2,292,420  
(0.3 per 1000)  
RR 4.91*  
(3.95 to 6.11)  
151 more per 100,000  
Very Low

Caesarean section: full guideline DRAFT (September 2011)  
Page 56 of 275
### Obstetric shock

<table>
<thead>
<tr>
<th>Study</th>
<th>Percentage of unplanned CS in planned vaginal birth group</th>
<th>Number of women</th>
<th>Effect</th>
<th>Quality</th>
</tr>
</thead>
<tbody>
<tr>
<td>Liu et al., 2007</td>
<td>3/46,766 (0.06 per 1000)</td>
<td>435/2,292,420</td>
<td>RR 0.33 (0.11 to 0.99)</td>
<td>12 fewer per 100,000* (from 17 fewer to 0.1 fewer)</td>
</tr>
</tbody>
</table>

* Calculated by NCC–WCH technical team  
** OR adjusted for age, race, gestational age and prolonged rupture of membranes

### Table 4.2 GRADE summary of findings for risks of planned CS compared with planned vaginal birth for women with BMI ≥ 50 kg/m² and no previous CS (maternal outcomes)

<table>
<thead>
<tr>
<th>Number of studies</th>
<th>Percentage of unplanned CS in planned vaginal birth group</th>
<th>Number of women</th>
<th>Effect</th>
<th>Quality</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Infection (wound and postpartum)</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 study</td>
<td>127/417 (30.5%)</td>
<td>38/174 (22%)</td>
<td>33/127* (26%)</td>
<td>OR 0.79 (0.46 to 1.35)</td>
</tr>
<tr>
<td><strong>Anaesthetic complication</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 study</td>
<td>127/417 (30.5%)</td>
<td>18/174 (10.3%)</td>
<td>35/417 (8.4%)</td>
<td>OR 1.25 (0.69 to 2.29)</td>
</tr>
<tr>
<td><strong>Major maternal morbidity (composite score including one or more of : intraoperative or postpartum haemorrhage, thromboembolic event, septicaemia, septic shock and/or admission to intensive care unit)</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 study</td>
<td>127/417 (30.5%)</td>
<td>11/174 (6.3%)</td>
<td>18/417 (4.3)</td>
<td>OR 1.49 (0.69 to 3.23)</td>
</tr>
</tbody>
</table>

* Denominator is women who had a CS

### Table 4.3 GRADE summary of findings for risks of planned CS compared with planned vaginal birth for women with an uncomplicated pregnancy and no previous CS (neonatal outcomes)

<table>
<thead>
<tr>
<th>Number of studies</th>
<th>Percentage of unplanned CS in planned vaginal birth group</th>
<th>Number of babies (%)</th>
<th>Effect</th>
<th>Quality</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Neonatal mortality</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 study</td>
<td>n = 5580/38021</td>
<td>0/1046</td>
<td>38/38021</td>
<td>Not</td>
</tr>
<tr>
<td>Condition</td>
<td>Study Reference</td>
<td>Rate 1 (per 1000 live births)</td>
<td>Rate 2 (per 1000 live births)</td>
<td>RR (95% CI)</td>
</tr>
<tr>
<td>--------------------------------------------------------------------------</td>
<td>--------------------------</td>
<td>--------------------------------</td>
<td>--------------------------------</td>
<td>-------------</td>
</tr>
<tr>
<td><strong>Hypoxic-Ischemic Encephalopathy (Central Nervous System depression, Seizures, pH &lt; 7)</strong></td>
<td>Dahlgren et al., 2009</td>
<td>617/1687,755,236</td>
<td>469/271,179</td>
<td>RR 2.4 (2.20 to 2.65)</td>
</tr>
<tr>
<td><strong>Intracranial haemorrhage</strong></td>
<td>Dahlgren et al., 2009</td>
<td>5580/38021</td>
<td>0/1046</td>
<td>NC</td>
</tr>
<tr>
<td><strong>Neonatal respiratory morbidity (composite outcome of intermittent positive pressure ventilation, transient tachypnoea, endotracheal tube insertion and pneumonia)</strong></td>
<td>Dahlgren et al., 2009</td>
<td>5580/38021</td>
<td>126/1046</td>
<td>RR 1.04* (0.88 to 1.23)</td>
</tr>
<tr>
<td><strong>Composite of respiratory morbidity</strong></td>
<td>Geller et al., 2010b</td>
<td>1340/3868</td>
<td>5/180</td>
<td>RR 1.2* (0.51 to 2.85)</td>
</tr>
<tr>
<td><strong>NICU admission</strong></td>
<td>Geller et al., 2010b</td>
<td>1340/3868</td>
<td>25/180</td>
<td>RR 2.20* (1.4 to 3.18)</td>
</tr>
<tr>
<td><strong>Apgar at 5 mins &lt; 7</strong></td>
<td>Dahlgren et al., 2009</td>
<td>5580/38021</td>
<td>0/1046</td>
<td>NC</td>
</tr>
<tr>
<td>Number of studies</td>
<td>Percentage of unplanned CS in planned vaginal birth group</td>
<td>Number of babies (%)</td>
<td>Effect</td>
<td>Quality</td>
</tr>
<tr>
<td>-------------------</td>
<td>------------------------------------------------------------</td>
<td>----------------------</td>
<td>--------</td>
<td>---------</td>
</tr>
<tr>
<td></td>
<td>Planned CS</td>
<td>Planned vaginal birth</td>
<td>Relative (95% CI)</td>
<td>Absolute</td>
</tr>
<tr>
<td>Neonatal mortality</td>
<td>1 study</td>
<td>127/417 (30.5%)</td>
<td>1/174 (0.6%)</td>
<td>2/417 (0.5%)</td>
</tr>
<tr>
<td></td>
<td>1 study</td>
<td>127/417 (30.5%)</td>
<td>27/174 (15.5%)</td>
<td>34/417 (8.3%)</td>
</tr>
</tbody>
</table>

* When adjusted for gestation < 37, the difference was not statistically significant between the two groups

**Evidence statements**

**Maternal outcomes: women with an uncomplicated pregnancy**

**Maternal mortality rate**

One study found that the maternal mortality rate in women with a planned CS was higher in women with a planned vaginal birth. This finding was statistically significant. Two other studies did not find a statistically significant difference in the maternal mortality rate between the two groups. The evidence for this outcome was of very low quality.

**Pain**

One study found that the median pain level during birth and three days postpartum was lower in women who had a planned CS compared with those who had a planned vaginal birth. This finding was statistically significant. The same study did not find a statistical difference between the two groups at 4 months postpartum. The evidence for this outcome was of very low quality.

---

1. * Calculated by NCC–WCH technical team
2. **No further details given in study
3. Table 4.4 GRADE summary of findings for risks of planned CS compared with planned vaginal birth for women with BMI ≥ 50 kg/m² and no previous CS (neonatal outcomes)
4. * When adjusted for gestation < 37, the difference was not statistically significant between the two groups
Injury to bladder/ureter
One study did not find a statistically significant difference in the incidence of injury to the bladder/ureter for women with a planned CS compared with women with a planned vaginal birth. The evidence for this outcome was of very low quality.

Injury to cervix
One study did not find a statistically significant difference in the incidence of injury to the cervix for women with a planned CS compared with women with a planned vaginal birth. The evidence for this outcome was of very low quality.

Injury to vagina
One study found that the incidence of injury to the vagina was lower among women with a planned CS compared with women who had a planned vaginal birth. This finding was statistically significant. The evidence for this outcome was of very low quality.

Iatrogenic surgical injury
One study did not find a statistically significant difference in the rate of iatrogenic surgical for women with a planned CS compared with women with a planned vaginal birth. The evidence for this outcome was of very low quality.

Hysterectomy
Two studies of very low quality found that the rate of hysterectomies was higher in women with a planned CS than in women with a planned vaginal birth. These findings were statistically significant. The evidence for this outcome was of very low quality.

Hysterectomy due to postpartum haemorrhage
One study found that the rate of hysterectomy due to postpartum haemorrhage was higher among women who had a CS without labour compared with women who had a planned vaginal birth. This finding was statistically significant. The evidence for this outcome was of very low quality.

Deep vein thrombosis
One study found that the rate of deep vein thrombosis (DVT) was higher in women who had a planned CS than in women with a planned vaginal birth. This finding was statistically significant. Another study did not find a statistically significant difference in rates of DVT between women who had a pre-labour CS and those who had a planned vaginal birth. The evidence for this outcome was of very low quality.

Pulmonary embolism
One study did not find a statistically significant difference in the rate of pulmonary embolism for women with a planned CS compared with women with a planned vaginal birth. The evidence for this outcome was of very low quality.

Blood transfusion
One study found that fewer women with a planned CS required a blood transfusion than women with a planned vaginal birth. This finding was statistically significant. Three other studies did not find a statistically significant difference in the number of women requiring a blood transfusion in the two groups. The evidence for this outcome was of very low quality.

Early postpartum haemorrhage
Two studies found that fewer women who had a CS without labour experienced an early postpartum haemorrhage than women who had a planned vaginal birth. This finding was statistically significant. The quality of evidence for this outcome was low in one study and very low in the other.

Wound and postpartum infection
One study of very low quality found that the rate of wound and postpartum infection was higher in women who had CS without labour compared with women who had a planned vaginal birth. This finding was statistically significant. One other study did not find a statistically significant difference in the rate of wound and postpartum infection between women who had a planned CS and those who had a planned vaginal birth. The evidence for this outcome was of very low quality.
Wound infection

Two studies did not find a statistically significant difference in the rate of wound infection for women with a planned CS compared with women with a planned vaginal birth. The evidence for this outcome was of low and very low quality.

Anaesthetic complications

One study found that the number of women experiencing anaesthetic complications was higher amongst women who had CS without labour compared with women who had a planned vaginal birth. This finding was statistically significant. One other study found no statistically significant difference in anaesthetic complications between women who had a planned CS and those who had a planned vaginal birth. The evidence for this outcome was of very low quality.

Intraoperative trauma

One study did not find a statistically significant difference in the rate of intraoperative trauma for women with a planned CS compared with women with a planned vaginal birth. The evidence for this outcome was of very low quality.

Uterine rupture

One study did not find a statistically significant difference in the rate of uterine rupture for women with a planned CS compared with women with a planned vaginal birth. The evidence for this outcome was of very low quality.

Length of hospital stay

Two studies found that the length of hospital stay was longer for women who had a CS without labour compared with women who had a planned vaginal birth. However, it was not possible to determine if this result was significant. The evidence for this outcome was of low and very low quality.

Assisted ventilation or intubation

One study found that more women who had a CS without labour required assisted ventilation or intubation compared with women who had a planned vaginal birth. This finding was statistically significant. The evidence for this outcome was of very low quality.

Acute renal failure

One study did not find a statistically significant difference in the rate of acute renal failure for women with a planned CS compared with women with a planned vaginal birth. The evidence for this outcome was of very low quality.

Cardiac arrest

One study found that more women who had a CS without labour experienced cardiac arrest compared with women who had a planned vaginal birth. This finding was statistically significant. The evidence for this outcome was of very low quality.

Obstetric shock

One study found that fewer women who had a CS without labour experienced obstetric shock compared with women who had a planned vaginal birth. This finding was statistically significant. The evidence for this outcome was of very low quality.

Maternal outcomes: women with BMI ≥ 50

Infection (wound and postpartum)

One study did not find a statistically significant difference in the rate of wound and postpartum infection for women with a planned CS compared with women with a planned vaginal birth. The evidence for this outcome was of low quality.

Anaesthetic complication

One study did not find a statistically significant difference in the rate of anaesthetic complications for women with a planned CS compared with women with a planned vaginal birth. The evidence for this outcome was of low quality.
Major maternal morbidity
One study did not find a statistically significant difference in the rate of major maternal morbidity for women with a planned CS compared with women with a planned vaginal birth. The evidence for this outcome was of low quality.

Neonatal outcomes: born to women with an uncomplicated pregnancy

Neonatal mortality
One study found that the neonatal mortality rate was higher in neonates born by planned CS compared with neonates born by planned vaginal birth. This finding was statistically significant. One other study did not find a statistically significant difference in the neonatal mortality rate between the two groups. The evidence for this outcome was of very low quality.

Hypoxic-ischemic encephalopathy
One study did not find a statistically significant difference in the rate of hypoxic-ischemic encephalopathy for neonates born by planned CS compared with neonates born by planned vaginal birth. The evidence for this outcome was of very low quality.

Intracranial haemorrhage
One study did not find a statistically significant difference in the rate of intracranial haemorrhage for neonates born by planned CS compared with neonates born by planned vaginal birth. The evidence for this outcome was of very low quality.

Neonatal respiratory morbidity (composite outcome of intermittent positive pressure ventilation, transient tachypnoea, endotracheal tube insertion and pneumonia)
One study did not find a statistically significant difference in the rate of neonatal respiratory morbidity for neonates born by planned CS compared with neonates born by planned vaginal birth. The evidence for this outcome was of very low quality.

Composite of respiratory morbidity
One study did not find a statistically significant difference in the composite respiratory morbidity for neonates born by planned CS compared with neonates born by planned vaginal birth. The evidence for this outcome was of very low quality.

NICU admission
One study found that the rate of NICU admission was higher in neonates born by planned vaginal birth. This finding was statistically significant. The evidence for this outcome was of low quality.

Apgar score < 7 at 5 minutes
One study found that fewer neonates born by planned CS had an Apgar score of < 7 at five minutes compared with neonates born by planned vaginal birth. This finding was statistically significant. Another study found no statistically significant difference between the two groups. The evidence for this outcome was of very low quality.

Composite of neurological morbidity
One study did not find a statistically significant difference in the composite neurological morbidity for neonates born by planned CS compared with neonates born by planned vaginal birth. The evidence for this outcome was of very low quality. Neonatal outcomes: born to women with BMI ≥ 50

Neonatal mortality
One study did not find a statistically significant difference in the neonatal mortality rate for neonates born by planned CS compared with neonates born by planned vaginal birth. The evidence for this outcome was of low quality.

NICU admission
One study found that the rate of NICU admission was higher in neonates born by planned vaginal birth. This finding was statistically significant. However, when the data were adjusted for gestation < 37 weeks, this finding was not statistically significant. The evidence for this outcome was of low quality.
Evidence to recommendations

Relative value placed on outcomes considered

For this review, all reported outcomes have been presented as the point of the review was to inform women’s decision making. By presenting these outcomes it is hoped that women will have the relevant information to assess the risks and benefits of each mode of birth. The relative value of outcomes will vary from woman to woman.

The group noted that deep vein thrombosis and pulmonary embolism were not particularly useful outcomes for individual decision making as they are very rare and are likely to be even more rare in the future as there are recent guidelines from the RCOG on these topics (Greentop guidelines 37a and b, 2009) and from NICE on prevention of venous thromboembolism (NICE, 2009) which should help to standardise and improve prevention and care of these complications.

Trade-off between clinical benefits and harms

The data have been presented in their entirety in order to support women’s decision making. The group agreed that it is important that women are presented with evidence based information in order that they are able to make an informed decision. The reported benefits and harms can then be discussed with each individual woman to help her make decisions based on the relative trade off between the two modes of birth interpreted in light of her own circumstances.

Trade-off between net health benefits and resource use

The outcomes reported in this review were used to inform the economic modelling for cost-effectiveness of CS on maternal request (see section 5.9 for further details and conclusions).

Quality of the evidence

The evidence was based on observational studies, the findings from which were all of low or very low quality. However, the group noted that the sole inclusion of low obstetric risk populations meant that the evidence was more relevant compared with the data presented in the original guideline for women requesting a CS with no medical or obstetric indication.

The group agreed that as the purpose of the table was to inform women’s decision making about their chosen mode of birth, it was most appropriate to include the findings for women who underwent an unplanned CS with those women who had a vaginal birth. This is because the majority of unplanned or intrapartum CS will be required by women who had originally planned for a vaginal birth and not those who had planned a CS.

Other considerations

The group noted that the findings from the studies were relevant to the current UK population which has a CS rate of approximately 25% of women. Were the rate of unplanned CSs to reduce, it is anticipated that the risks associated with planned vaginal birth would also reduce.

The group agreed that when discussing the risks and benefits outlined in the table, the healthcare professional and woman also need to consider the woman’s individual circumstances which affect the risks associated with vaginal birth and CS such as previous abdominal or pelvic surgery, impaired mobility from pelvic girdle pain, or care of other children. It is also important to discuss the number of future babies that the woman and her partner are planning as some risks such as placenta praevia increase with an increasing number of CS.

The GDG noted that the evidence from the one UK study involving extremely obese women suggested that outcomes were similar for women planning a vaginal birth and those planning a CS. They felt this was contrary to what many obstetricians currently believe to be the case and that it was important to make a recommendation to underline this finding.

Recommendations

<table>
<thead>
<tr>
<th>Number</th>
<th>Recommendation</th>
</tr>
</thead>
<tbody>
<tr>
<td>4</td>
<td>Discuss the risks and benefits of CS and vaginal birth with the woman (see box A and recommendation 118), taking into account her circumstances, concerns, priorities and plans for future pregnancies (including the risks of placental problems with multiple CS). [new 2011]</td>
</tr>
</tbody>
</table>
Box A Planned caesarean section compared with planned vaginal birth for women with an uncomplicated pregnancy and no previous caesarean section

Planned caesarean section may reduce the risk of the following in women:
- perineal and abdominal pain during birth and 3 days postpartum
- injury to vagina
- early postpartum haemorrhage
- obstetric shock.

Planned caesarean section may reduce the risk of the following in babies:
- neonatal intensive care unit admission.

Planned caesarean section may increase the risk of the following in women:
- longer hospital stay
- hysterectomy caused by postpartum haemorrhage
- cardiac arrest.

Please refer to tables 4.5 and 4.6 for full details, including the absolute and relative risks for each effect.

Do not use a body mass index (BMI) of over 50 alone as an indication for planned CS. [new 2011]

<table>
<thead>
<tr>
<th>Number</th>
<th>Research Recommendation</th>
</tr>
</thead>
<tbody>
<tr>
<td>RR 1</td>
<td>What are the medium- to long-term risks and benefits to women and their babies of planned CS compared with planned vaginal birth?</td>
</tr>
</tbody>
</table>

The main focus would be the outcomes in women which could be measured at 1 year (medium term) and 5-10 years (long term). These outcomes could include:

- urinary dysfunction
- gastrointestinal dysfunction
- dyspareunia
- breastfeeding
- psychological health.

Infant outcomes could include medical problems, especially ongoing respiratory and neurological problems.

Why this is important

Morbidities arising intraoperatively or in the days after a caesarean section have been reasonably well described in the literature. Much less is known, however, about physical and emotional outcome measures in the longer term.

The Confidential Enquiries into Maternal Death in the UK, most recently published as ‘Saving mothers’ lives 2006-2008’ (Cantwell R. et al., 2011), devote a significant proportion of their work to investigating ‘late’ causes of maternal death. These include events arising in the medium term, namely, up to 1 year after a woman has given birth, many of which originate from the preceding pregnancy. The infectious, psychiatric and other conditions arising in or related to pregnancy do not always cause death but are responsible for arguably a greater burden of morbidity in the medium and long term, long after the pregnancy is over.

To provide more meaningful information to women when they are choosing their mode of birth, there is a pressing need to document medium- to long-term outcomes
in women and their babies after a planned CS or a planned vaginal birth. First, it should be possible to gather data using standardised questions (traditional paper-based questionnaires and face-to-face interviews) about maternal septic morbidities and emotional wellbeing up to 1 year after a planned CS in a population of women who have consented for follow-up. Internet-based questionnaires could also be devised, to achieve the high response rates required for a full interpretation of the data. Similarly, it would be important to collect high-quality data on infant morbidities after a planned CS compared with a planned vaginal birth. A long term morbidity evaluation (between 5 and 10 years after the CS) would use similar methodology but assess symptoms related to urinary and gastrointestinal function.

Further evaluation is needed to determine the impact of demographic and clinical factors (such as ethnic group, increase in body mass index) and attitudinal factors on CS rates.
### Table 4.5 Summary effect on women’s health of planned CS compared with planned vaginal birth for women with an uncomplicated pregnancy and no previous CS

<table>
<thead>
<tr>
<th>Effects around the time of birth</th>
<th>Finding for planned CS</th>
<th>Finding for planned vaginal birth</th>
<th>Absolute effect</th>
<th>Relative effect (95% confidence interval)</th>
<th>Evidence quality</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Absolute effect (95% confidence interval)</td>
<td>Evidence quality</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Studies suggest may be reduced after a planned CS</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Perineal and abdominal pain during birth (score/10 – higher scores indicate higher pain levels)</td>
<td>Median score 1.0</td>
<td>Median score 7.3</td>
<td>6.3 lower</td>
<td>NC</td>
<td>Very low</td>
</tr>
<tr>
<td>(including % unplanned CS in planned vaginal birth group)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Perineal and abdominal pain 3 days postpartum (score/10 – higher scores indicate higher pain levels)</td>
<td>Median score 4.5</td>
<td>Median score 5.2</td>
<td>0.7 lower</td>
<td>NC</td>
<td>Very low</td>
</tr>
<tr>
<td>Injury to vagina</td>
<td>0.0%</td>
<td>0.56%</td>
<td>6 fewer per 1000 (from 6 fewer to 2 fewer)</td>
<td>NC</td>
<td>Very low</td>
</tr>
<tr>
<td>Early postpartum haemorrhage</td>
<td>1.1%</td>
<td>6.0%</td>
<td>49 per 1000 (from 4 fewer to 56 fewer)</td>
<td>OR 0.23 (0.06 to 0.94)</td>
<td>Low</td>
</tr>
<tr>
<td></td>
<td>3.9%</td>
<td>6.2%</td>
<td>23 fewer per 1000 (from 35 fewer to 6 fewer)</td>
<td>RR 0.06 (0.4 to 0.9)</td>
<td>Very low</td>
</tr>
<tr>
<td>Obstetric shock</td>
<td>0.006%</td>
<td>0.018%</td>
<td>12 fewer per 100,000 (from 17 fewer to 0.1 fewer)</td>
<td>RR 0.33 (0.11 to 0.99)</td>
<td>Very low</td>
</tr>
<tr>
<td><strong>Studies suggest may be reduced after planned vaginal birth</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Length of hospital stay</td>
<td>3.2 days</td>
<td>2.6 days</td>
<td>0.6 days longer</td>
<td>Mean difference 1.58 (1.27 to 2.17)</td>
<td>Low</td>
</tr>
<tr>
<td></td>
<td>3.96 days</td>
<td>2.56 days</td>
<td>1.4 days longer</td>
<td>Adjusted mean difference 1.47 (1.46 to 1.49)</td>
<td>Very low</td>
</tr>
<tr>
<td>Event</td>
<td>Risk before</td>
<td>Risk after</td>
<td>Additional Risk</td>
<td>RR</td>
<td>Confidence Interval</td>
</tr>
<tr>
<td>--------------------------------------------</td>
<td>-------------</td>
<td>------------</td>
<td>-----------------</td>
<td>------</td>
<td>---------------------</td>
</tr>
<tr>
<td>Hysterectomy due to postpartum haemorrhage</td>
<td>0.03%</td>
<td>0.01%</td>
<td>14 more per 100,000 (from 3 more to 33 more)</td>
<td>RR 2.31</td>
<td>(1.30 to 4.09)</td>
</tr>
<tr>
<td>Cardiac arrest</td>
<td>0.19%</td>
<td>0.03%</td>
<td>15 more per 10,000 (from 11.5 more to 19.5 more)</td>
<td>RR 4.91</td>
<td>(3.95 to 6.11)</td>
</tr>
<tr>
<td>No difference found in studies</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Perineal and abdominal pain 4 months postpartum</td>
<td>Median score 0.0</td>
<td>Median score 0.17</td>
<td>0.17 lower</td>
<td>NC</td>
<td></td>
</tr>
<tr>
<td>Injury to bladder/ureter</td>
<td>0.0%</td>
<td>0.14%</td>
<td>1 fewer per 1000 (from 2 fewer to 2 more)</td>
<td>NC</td>
<td></td>
</tr>
<tr>
<td>Injury to cervix</td>
<td>0.0%</td>
<td>0.28%</td>
<td>3 fewer per 1000 (from 3 fewer to 1 more)</td>
<td>NC</td>
<td></td>
</tr>
<tr>
<td>Iatrogenic surgical injury</td>
<td>0.00%</td>
<td>0.07%</td>
<td>7 fewer per 10,000</td>
<td>NC</td>
<td></td>
</tr>
<tr>
<td>Pulmonary embolism</td>
<td>0.00%</td>
<td>0.003%</td>
<td>2 fewer per 10,000 (from 2 fewer to 40 more)</td>
<td>NC</td>
<td></td>
</tr>
<tr>
<td>Wound infection</td>
<td>0.01%</td>
<td>0.00%</td>
<td>1 more per 10,000</td>
<td>p = 1.0</td>
<td></td>
</tr>
<tr>
<td></td>
<td>1.5%</td>
<td>0.9%</td>
<td>6 fewer per 1000 (from 1 fewer to 19 more)</td>
<td>RR 1.7</td>
<td>(0.9 to 3.2)</td>
</tr>
<tr>
<td>Intraoperative trauma</td>
<td>0.1%</td>
<td>0.3%</td>
<td>1 fewer per 1000 (from 3 fewer to 7 more)</td>
<td>RR 0.5</td>
<td>(0.1 to 3.5)</td>
</tr>
<tr>
<td>Uterine rupture</td>
<td>0.02%</td>
<td>0.03%</td>
<td>13 fewer per 100,000 (from 22 fewer to 2.2 more)</td>
<td>RR 0.51</td>
<td>(0.25 to 1.07)</td>
</tr>
<tr>
<td>Event</td>
<td>Expected Rate</td>
<td>Actual Rate</td>
<td>Difference (per 100,000)</td>
<td>RR (95% CI)</td>
<td>Evidence Quality</td>
</tr>
<tr>
<td>--------------------------------------</td>
<td>---------------</td>
<td>-------------</td>
<td>--------------------------</td>
<td>-------------</td>
<td>------------------</td>
</tr>
<tr>
<td>Assisted ventilation or intubation</td>
<td>0.01%</td>
<td>0.005%</td>
<td>7 more (from 0 fewer to 22 more)</td>
<td>RR 2.21 (0.99 to 4.90)</td>
<td>Very low</td>
</tr>
<tr>
<td>Acute renal failure</td>
<td>0.004%</td>
<td>0.001%</td>
<td>2 more (from 9 fewer to 13 more)</td>
<td>RR 2.17 (0.58 to 8.14)</td>
<td>Very low</td>
</tr>
<tr>
<td>Maternal death</td>
<td>9/737</td>
<td>49/9133</td>
<td>No difference (no events)</td>
<td>NC</td>
<td>Very low</td>
</tr>
<tr>
<td>Deep vein thrombosis</td>
<td>0.00%</td>
<td>0.03%</td>
<td>0.7 fewer (from 0.2 fewer to 4 more)</td>
<td>NC</td>
<td>Very low</td>
</tr>
<tr>
<td>Blood transfusion</td>
<td>1.7%</td>
<td>1.9%</td>
<td>2 fewer (from 14 fewer to 34 more)</td>
<td>OR 0.87 (0.27 to 2.78)</td>
<td>Low</td>
</tr>
<tr>
<td></td>
<td>0.3%</td>
<td>0.3%</td>
<td>0 fewer (from 2 fewer to 5 more)</td>
<td>RR 0.89 (0.20 to 3.99)</td>
<td>Very low</td>
</tr>
<tr>
<td></td>
<td>0.3%</td>
<td>0.4%</td>
<td>1 fewer (from 2 fewer to 5 more)</td>
<td>RR 0.7 (0.2 to 2.7)</td>
<td>Very low</td>
</tr>
<tr>
<td></td>
<td>0.02%</td>
<td>0.07%</td>
<td>41 fewer (from 53 fewer to 86 more)</td>
<td>RR 0.20 (0.20 to 0.64)</td>
<td>Very low</td>
</tr>
</tbody>
</table>

**Conflicting findings from studies**

Maternal death

Of maternal deaths occurring in the planned vaginal birth group 13/49 (26.5%) were women who gave birth by unplanned CS

<table>
<thead>
<tr>
<th>Event</th>
<th>Expected Rate</th>
<th>Actual Rate</th>
<th>Difference (per 100,000)</th>
<th>RR (95% CI)</th>
<th>Evidence Quality</th>
</tr>
</thead>
<tbody>
<tr>
<td>Maternal death</td>
<td>0.00%</td>
<td>0.00%</td>
<td>No difference (no events)</td>
<td>NC</td>
<td>Very low</td>
</tr>
<tr>
<td>Deep vein thrombosis</td>
<td>0.00%</td>
<td>0.02%</td>
<td>10 fewer (from 0 fewer to 22 more)</td>
<td>RR 2.21 (0.99 to 4.90)</td>
<td>Very low</td>
</tr>
<tr>
<td>Blood transfusion</td>
<td>0%</td>
<td>0%</td>
<td>No difference (no events)</td>
<td>NC</td>
<td>Very low</td>
</tr>
</tbody>
</table>

Caesarean section: full guideline DRAFT (September 2011)
<table>
<thead>
<tr>
<th>Condition</th>
<th>Rate 1</th>
<th>Rate 2</th>
<th>Difference</th>
<th>RR (CI)</th>
<th>Evidence Level</th>
</tr>
</thead>
<tbody>
<tr>
<td>Infection – wound and postpartum</td>
<td>1.1%</td>
<td>0.8%</td>
<td>3 more per 1000 (from 2 fewer to 11 more)</td>
<td>RR 1.36 (0.75 to 2.4)</td>
<td>Very low</td>
</tr>
<tr>
<td></td>
<td>(14.7%)</td>
<td>(14.7%)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>0.6%</td>
<td>0.21%</td>
<td>390 more per 100,000 (from 323 more to 464 more)</td>
<td>RR 2.85 (2.52 to 3.21)</td>
<td>Very low</td>
</tr>
<tr>
<td></td>
<td>(8.2%)</td>
<td>(8.2%)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hysterectomy</td>
<td>0.6%</td>
<td>0.1%</td>
<td>5 more per 1000</td>
<td>p = 0.13</td>
<td>Low</td>
</tr>
<tr>
<td></td>
<td>(35%)</td>
<td>(35%)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>0.1%</td>
<td>0.01%</td>
<td>1 more per 1000 (from 0 more to 5 more)</td>
<td>RR 9.09 (1.36 to 60.33)</td>
<td>Very low</td>
</tr>
<tr>
<td></td>
<td>(14.7%)</td>
<td>(14.7%)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>0.06%</td>
<td>0.02%</td>
<td>41 more per 100,000 (from 23.6 more to 68 more)</td>
<td>RR 3.60 (2.44 to 5.31)</td>
<td>Very low</td>
</tr>
<tr>
<td></td>
<td>(8.2%)</td>
<td>(8.2%)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Anaesthetic complications</td>
<td>0.4%</td>
<td>0.3%</td>
<td>1 more per 1000 (from 2 fewer to 11 more)</td>
<td>RR 1.24 (0.34 to 4.59)</td>
<td>Very low</td>
</tr>
<tr>
<td></td>
<td>(14.7%)</td>
<td>(14.7%)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>0.53%</td>
<td>0.21%</td>
<td>319 more per 100,000 (from 257 more to 389 more)</td>
<td>RR 2.5 (2.22 to 2.86)</td>
<td>Very low</td>
</tr>
<tr>
<td></td>
<td>(8.2%)</td>
<td>(8.2%)</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
## Table 4.6 Summary effect on babies’ health of planned CS compared with planned vaginal birth for women with an uncomplicated pregnancy and no previous CS

<table>
<thead>
<tr>
<th>Effects around the time of birth</th>
<th>Finding for CS</th>
<th>Finding for vaginal birth</th>
<th>Absolute effect</th>
<th>Relative effect (95% confidence interval)</th>
<th>Evidence quality</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Studies suggest may be reduced after planned vaginal birth</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>NICU admission</td>
<td>13.9%</td>
<td>6.3%</td>
<td>76 more per 1000 (from 31 more to 134 more)</td>
<td>RR 2.20 (1.4 to 3.18)</td>
<td>Low</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>(35%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>No difference found in studies</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hypoxic-Ischemic Encephalopathy (CNS depression, seizures, pH &lt; 7)</td>
<td>0.2%</td>
<td>0.2%</td>
<td>0 fewer per 1000 (from 2 fewer to 5 more)</td>
<td>RR 0.81 (0.22 to 3.00)</td>
<td>Very low</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>(14.7%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Intracranial haemorrhage</td>
<td>0.00%</td>
<td>0.01%</td>
<td>0.2 fewer per 1000 (from 0.4 fewer to 3 more)</td>
<td>NC</td>
<td>Very low</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>(14.7%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Neonatal respiratory morbidity (intermittent positive pressure ventilation, transient tachypnoea, endotracheal tube insertion, pneumonia)</td>
<td>12.0%</td>
<td>11.5%</td>
<td>5 more per 1000 (from 14 fewer to 27 more)</td>
<td>RR 1.04 (0.88 to 1.23)</td>
<td>Very low</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>(14.7%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Conflicting findings from studies</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Neonatal mortality</td>
<td>0.0%</td>
<td>0.1%</td>
<td>1 fewer per 1000 live births (from 1 fewer to 2 more)</td>
<td>NC</td>
<td>Very low</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>(14.7%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>0.17%</td>
<td>0.07%</td>
<td>1 more per 1000 live births (from 1 more to 2 more)</td>
<td>RR 2.4 (2.20 to 2.65)</td>
<td>Very low</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>(7.9%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Apgar score at 5 mins &lt; 7</td>
<td>0.0%</td>
<td>0.5%</td>
<td>5 fewer per 1000 (from 5 fewer to 1 fewer)</td>
<td>NC</td>
<td>Very low</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>(14.7%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>0.6%</td>
<td>1.2%</td>
<td>6 fewer per 1000 (from 9 fewer to 157 more)</td>
<td>RR 0.44 (0.07 to 2.51)</td>
<td>Very low</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>(35%)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Consent for CS

Provision of information is central to the consent process, this should include information about the patient’s condition, possible investigations and treatment options; the risks or benefits of these options, including the risk of doing nothing.\textsuperscript{30–32} [evidence level 4] Information should be given in a way that patients can understand.\textsuperscript{32,33} [evidence level 4] The amount of information provided will vary between patients according to the nature of the condition, the complexity of the treatment, the associated risk of the procedure, the patient’s own wishes and individual needs. For the process of seeking consent to be meaningful, refusal of treatment needs to be one of the patient’s options. Competent adults are entitled to refuse treatment even when the treatment would clearly benefit their health. Therefore a competent pregnant woman may refuse CS, even if this would be detrimental to herself or the fetus.\textsuperscript{30} [evidence level 4] Ethical guidance for obtaining consent, points of law and model documentation are available in the above guidance.\textsuperscript{30–32,34} [evidence level 4]

Tubal ligation at CS

It is estimated tubal ligation overall has a failure rate of 1 in 200 lifetime risk.\textsuperscript{49} We did not identify any studies that describes the failure rate of tubal ligation at CS. Other guidelines recommend that tubal ligation should have been requested before or during pregnancy and agreed at least one week prior to the procedure. This advice is based on expert opinion.\textsuperscript{49} [evidence level 4]

Recommendations

<table>
<thead>
<tr>
<th>Number</th>
<th>Recommendation</th>
</tr>
</thead>
<tbody>
<tr>
<td>6</td>
<td>Consent for CS should be requested after providing pregnant women with evidence-based information and in a manner that respects the woman’s dignity, privacy, views and culture, while taking into consideration the clinical situation. [C] [2004]</td>
</tr>
<tr>
<td>7</td>
<td>A pregnant woman is entitled to decline the offer of treatment such as CS, even when the treatment would clearly benefit her or her baby’s health. Refusal of treatment needs to be one of the woman’s options. [D] [2004]</td>
</tr>
<tr>
<td>8</td>
<td>When a decision is made to perform a CS, a record should be made of all the factors that influence the decision, and which of these is the most influential. [GPP] [2004]</td>
</tr>
</tbody>
</table>
5 Planned CS

This chapter considers the evidence related to decisions about planned mode of birth. Other aspects of management of specific conditions or complications of pregnancy are not included because they are outside the scope of the guideline.

5.1 Breech presentation

About 4% of all singleton pregnancies are breech presentation. The proportion of breech presentation fetuses decreases with increasing gestation: 3% of term infants, 9% for those born at 33–36 weeks of gestation, 18% of those born at 28–32 weeks and 30% of those born at less than 28 weeks. Breech presentation, is associated with cerebral palsy and handicap, due principally to the association with preterm birth and congenital malformations. Breech presentation is the primary indication for 10% of all CS. Overall 88% of pregnancies with breech presentation in England and Wales are delivered by CS (56% planned and 44% unplanned CS). However CS rates vary with gestational age, at term 91% women with a breech presentation had a CS, while at less than 28 weeks the CS rate was less than 40%.

External cephalic version

Interventions to promote cephalic version of babies in the breech position include external cephalic version (ECV), moxibustion and postural management. The research basis for these interventions is included in the guideline on antenatal care of healthy pregnant women. External cephalic version involves applying pressure to the mother's abdomen to turn the fetus in either a forward or backward somersault to achieve a vertex presentation. Recognised complications of ECV attributable to the procedure (and incidence) include:

- fetal heart rate abnormalities: the commonest is transient bradycardia (1.1% to 16%)60–62
- placental abruption (0.4% to 1%)59,61
- painless vaginal bleeding (1.1%)61
- admission for induction of labour (3%).62

Two systematic reviews examined the effect of ECV at term and before term. Performing ECV at term reduced the number of non-cephalic births by 60% when compared with no ECV (6 RCTs, n = 612 women, RR 0.42, 95% CI 0.35 to 0.50). A reduction in caesarean section is also observed in the ECV group when compared with no ECV (6 RCTs, n = 612, RR 0.52, 95% CI 0.39 to 0.71). ECV before 37 weeks gestation does not reduce non-vertex births at term (RR 1.02, 95% CI 0.89 to 1.17). Success rates following ECV in primiparous women range from 35% to 57% and from 52% to 84% in multiparous women.59–62,65 Interventions to improve the success rates of ECV include the routine or selective use of tocolysis, the use of regional analgesia and the use of vibroacoustic stimulation. None of the RCTs has used newer tocolytics and the effectiveness of these is uncertain. Further guidance on ECV may be found in the RCOG green top guideline on the management of breech presentation.

In the NSCSA external cephalic version was offered to 33% of women having a CS for breech presentation at term, this was the same irrespective of the woman's parity. ECV was provided by consultants, specialist registrars or staff grade obstetricians. If ECV was offered to
all women at term, assuming a 50% success rate then it is likely this would reduce the overall CS rate by 1%.

Cost effectiveness of ECV
Six cost-effectiveness studies were identified that considered the role of ECV in decreasing the rate of CS, two in the United Kingdom and four in the USA.

The UK studies reported the cost of ECV. The first was a cost study that reported an expected cost of £1,452 for ECV versus £1,828 for not having ECV, an expected saving of around £380.53. The results were insensitive (i.e. did not alter the result) to changes in the cost of an ECV. The cost of CS would need to fall by £8,576 (a fall of 56%, again, a highly unlikely scenario) for the non-ECV option to be the less costly option. However, the sensitivity analysis showed that the ECV success rate would only have to fall by around 5% for the ECV option to be the less favourable option. Therefore the cost analysis cannot categorically determine which option is least costly overall. The second UK study (much smaller) found that the cost of birth with a successful ECV was £2,230 and £2,595 for an unsuccessful ECV, a cost saving of around £360 per birth.

Four American studies have been published. One used a decision analytic modelling technique to determine the overall costs of four management options for breeches at term: ECV with planned vaginal birth, ECV with CS, selected vaginal birth and planned CS. The decision model used hospital charges (not costs) for vaginal birth of US$6000 and US$10,000 for CS (a wider ratio than the reported UK cost data). The expected CS rate was 25.4% (± 5.4) for ECV plus planned vaginal birth; 31.9% (± 6.6) for ECV plus planned vaginal birth; 62.6% (± 5.9) for selected vaginal birth and 88.6% (± 3.4) for planned CS. The model estimated the expected cost for each pathway (the cost of vaginal birth and CS for each option arm) and found that ECV with planned vaginal birth was the least costly option, due to the lower proportion of CS for this group (US$8071) and planned CS to be the most costly (US$9544). Whether these reported costs were statistically different is not reported. The validity of the range of probabilities used in the decision analysis were subsequently questioned.

A study in the same year considered the costs of failed and successful ECV separately and reported a cost of US$8042 for women with failed ECV and US$5059 for women with successful ECV. However, the effectiveness data on which this study was based was a cohort study and not an RCT.

An American study also presented data to show that successful ECV would yield savings over unsuccessful ECV. The most recent US study was a much larger study of 695 women. This was a decision-analytic model to calculate the potential cost savings from ECV (in terms of reduced CS rates). The authors assumed that ECV would be successful in 44% of cases, of which 67% would proceed to vaginal birth and 33% to a CS. They further assumed that ECV would be unsuccessful in 56% of cases, of which only 7% would proceed to a successful vaginal birth. Given these assumptions, the model calculated a savings (in US hospital charges) of around $650 per birth. Savings from every ECV attempted (even if not successful) versus ECV not attempted were around US$3000 per birth (these are greater due to higher reported rates of CS for women not attempting ECV).

Therefore in conclusion ECV yields cost savings in comparison with CS. There is no UK-based economic evaluation comparing ECV with vaginal breech birth.

Term breech pregnancy and CS
A systematic review identified 3 RCTs (n = 2396) that evaluated the effect of mode of birth for term breech pregnancies. The majority of the information about the effect of planned CS in the review comes from one international multi-centre RCT which is of good methodological quality (n = 2088 women, 121 centres in 26 countries). Offering planned CS reduced perinatal or neonatal death (excluding fatal anomalies) or serious neonatal morbidity (RR 0.33, 95% CI 0.19 to 0.56). The risk of perinatal/neonatal mortality or serious morbidity was 1.6% in the planned CS group and 5.0% in the planned vaginal birth group. The absolute risk reduction in perinatal/neonatal mortality or serious neonatal morbidity was 3.4%, therefore for every 29 CS for term breech pregnancy one baby will avoid death or serious morbidity.
The findings of the RCT and the systematic review are the subject of continued debate. Therefore more details about this RCT are outlined here. The RCT included a number of maternity units in the UK. About 40% of women recruited to the trial were in labour at time of randomisation. The women in labour were not further divided into stages of labour so there is no information on how many were in the second stage of labour. However advanced labour was not listed as an exclusion criterion. The RCT protocol provided guidance on management of labour. This included intermittent fetal heart monitoring (every 15 minutes in the first stage and every 5 minutes in the second stage), adequate progress in labour was defined as 0.5 cm dilatation per hour and descent of the breech to the pelvic floor within 2 hours of full cervical dilatation. Delivery of the breech could be spontaneous or assisted; the after coming head could be controlled using the Mauriceau–Smellie–Veit manoeuvre or forceps. The position of the woman for the second stage of labour was not stipulated by the protocol nor was this information collected during the trial.

Sub group analysis within this RCT has been undertaken to evaluate if the effect on perinatal mortality or morbidity could be explained by specific factors. These effects remain consistent and are therefore not explained by differences in:

- operator experience
- prolonged labour
- induction of labour with oxytocin or prostaglandins
- augmentation of labour
- type of breech presentation (footling or uncertain)
- the use of epidural analgesia.

Women who were in labour were included in the RCT (therefore the findings of the trial are generalisable to women in labour); however the effect of CS on neonatal outcomes is not reported separately for this group. It is possible that the benefits and risks of caesarean section particularly during the second stage are different. Therefore further research that specifically addressed this issue was sought; however no studies evaluating the effect of CS for undiagnosed breech compared to expectant management were identified. An RCT to address this issue would require randomisation of at least 4230 women with undiagnosed breech pregnancy to either CS or vaginal birth in order to detect at least a 40% difference in neonatal morbidity.

The effects of planned CS for term breech on maternal health are less clear. The RCTs included in the systematic review assessed the impact of CS on maternal health using a variety of measures and combining the results across studies is not always possible. Where the estimates could be combined, no difference is detected in the measures of maternal morbidity (such as blood loss, blood transfusion, infection) between planned CS and planned vaginal birth. Estimates of composite measures of morbidity have previously been reported however these pooled estimates are not included in the guideline because it is unclear whether these estimates are based on person or event data. It is possible that the same woman may have more than one morbidity (for example a woman who needs additional surgery is more likely to need a blood transfusion or admission to ITU) so that composite morbidity measures based on summation of event rates rather than number of women affected can lead to spurious results. [evidence level 1b] Data for individual women was reported in one RCT, it did not detect any difference in composite maternal morbidity between women in the planned CS group or women in the planned vaginal birth group (RR1.24, 95% CI 0.79 to 1.95). [evidence level 1b] The specific estimates of the effect of planned CS on maternal health are outlined in Table 4.5

**Preterm breech**

Breech presentation, is associated with cerebral palsy and handicap, due principally to the association with preterm birth and congenital malformations. The proportion of breech presentation fetuses decreases with increasing gestation: 9% for those born at 33–36 weeks of gestation, 18% of those born at 28–32 weeks and 30% of those born at less than 28 weeks.
Overall 88% of pregnancies with breech presentation were delivered by CS. However CS rates varied by gestational age, 87% for babies born at 33–36 weeks, 81% of those born at 28–32 weeks, and 39% for babies born at less than 28 weeks.4 [evidence level 3]

The results of the term breech trial RCT are relevant for term breech pregnancies, extrapolation to preterm breech babies is inappropriate. In the CESDI Project 27/28 report, survival rates were lower for babies who were breech (84.5%) when compared to babies who were cephalic presentation (89.4%). Survival for breech presentation was significantly greater in those delivered by CS (86.5%) than those delivered vaginally (77.4%).76 [evidence level 3]

### Recommendations

<table>
<thead>
<tr>
<th>Number</th>
<th>Recommendation</th>
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<tbody>
<tr>
<td>9</td>
<td>Women who have an uncomplicated singleton breech pregnancy at 36 weeks’ gestation should be offered external cephalic version. Exceptions include women in labour and women with a uterine scar or abnormality, fetal compromise, ruptured membranes, vaginal bleeding or medical conditions. [A] [2004]</td>
</tr>
<tr>
<td>10</td>
<td>Pregnant women with a singleton breech presentation at term, for whom external cephalic version is contraindicated or has been unsuccessful, should be offered CS because it reduces perinatal mortality and neonatal morbidity. [A] [2004]</td>
</tr>
</tbody>
</table>

### Research Recommendation

<table>
<thead>
<tr>
<th>RR 3</th>
<th>Further research is needed to determine the effect of caesarean section compared with vaginal birth for women with:</th>
</tr>
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<tbody>
<tr>
<td></td>
<td>- preterm breech</td>
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<tr>
<td></td>
<td>- a breech presentation that is diagnosed in the second stage of labour</td>
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</table>

## 5.2 Multiple pregnancy

About 15 per 1000 pregnancies are multiple gestations; the majority of these are twin pregnancies (twins 14.4 per 1000, triplets 4 per 10,000).4 There have been increases in the rates of multiple pregnancy in the last ten years that are attributed to the use of ovulation induction in fertility treatments.77,78 Perinatal mortality and morbidity such as cerebral palsy are higher among multiple births than singleton births (stillbirths: multiple: 2%; singleton: 0.5%; neonatal deaths multiple: 2.5. singleton 0.3%); RR cerebral palsy twins 4.63 (3.32–6.46.)79,80 [evidence level 3] Some of the observed increase is explained by the association of multiple pregnancy with preterm birth.4,80 Other factors which have been associated with poorer outcome in twin pregnancy include low birth weight, discordant growth between twins, monochorionic twins and being a second born twin.81–85 The management of complications (such as discordant growth, monochorionic twins) and other obstetric complications in pregnancy (such as pre-eclampsia) will influence the mode of delivery decisions, however these are outside the scope of this guideline and are therefore not discussed further in this section.

Multiple pregnancy is the primary indication for 1% of caesarean sections.4 Overall 59% of twin pregnancies were delivered by CS. (37% planned and 63% unplanned CS). CS for delivery of the second twin following vaginal birth of the first baby was carried out in 3.5% of twins (n = 75). CS rates vary by gestational age, at term 60% women with a twin pregnancy had a CS, while at less than 28 weeks the CS rate was less than 29%.4 [evidence level 3] Where CS was planned for multiple pregnancy, breech presentation of the first twin was the most commonly reported indication (14%), together with previous CS (7%) and maternal request (9%). Of the unplanned caesarean sections,
fetal distress was the most influential factor in 29% and “failure to progress” in 12%. Almost all triplet pregnancies (92%) were delivered by CS. [evidence level 3]

A systematic review that included 1 RCT (n = 60) compared CS for a second twin with a non vertex presentation to vaginal birth. [evidence level 1b] The methodological quality of this trial is uncertain because ‘randomisation was according to a protocol that was changed randomly by a non-involved person, without prior notice, on a time basis’. No difference was detected in any of the baby outcome measures, however the study is too small to accurately estimate the effect on outcomes such as neonatal birth trauma and perinatal death. The study reported no difference in the average length of hospital stay (8 days compared to 5 days) and no difference in need for blood transfusion (RR 1.5 95% CI 0.27 to 8.28). Women in the planned CS group had increased risk of puerperal pyrexia compared to women in the planned vaginal birth group (RR 3.67 95% CI 1.15 to 11.69). [evidence level 1b]

A large number of observational studies using population based registers have been published. However the majority of these studies are analysed by actual mode of delivery rather than intended mode of delivery, the reports provide insufficient data on neonatal outcome for women who had planned CS and in the analysis paired tests have not been used to take into account that the outcome within twin pairs maybe related. One systematic review included only studies where the intended mode of delivery could be identified. The review included 3 retrospective cohort studies and the RCT discussed above. The results from these studies were consistent and did not detect differences in neonatal morbidity such as low 5-minute Apgar score, birth trauma, neurological complications, hyperbilirubinaemia, hypoglycaemia, transient tachypnoea or secondary apnoea. The studies are too small to evaluate perinatal mortality.

Triplet and higher order multiple births are rare. They most frequently are the result of ovulation induction for treatment of fertility problems. Tripletts are almost always born preterm and some of the poorer outcomes such as cerebral palsy seen in these infants are due to preterm birth. These and other complicating factors may influence the mode of delivery decisions. Almost all triplet pregnancies (92%) were delivered by CS. [evidence level 3] We identified 3 small retrospective case control studies which compared baby outcomes according to mode of birth for triplet pregnancies (119 sets of triplets in total). The babies born vaginally tended to have better outcomes such as higher Apgar scores than those delivered by CS. However these studies are analysed by actual mode of delivery rather than intended mode of delivery and do not use analysis to take into account that the outcome within triplets will be related.

Women who have multiple pregnancies have an increased risk of maternal mortality and morbidity. CEMD estimates maternal mortality is increased with multiple pregnancy (20.3 per 100 000 twin pregnancies; 215 per 100 000 triplet pregnancies, compared with 11.2 per 100 000 for singleton pregnancies). [evidence level 3] The effect of mode of delivery on this outcome is uncertain.

Timing of planned CS for twin pregnancy

Planned CS of twins between 36–37 weeks and 6 days is associated with increased risk of respiratory disorders (TTN or RDS) in one or both of the twins compared to CS between 38 and 40 weeks (RR 5.94, 95% CI 0.78 to 45.01). Multiple pregnancy is an established risk factor for preterm birth. About 29% of twin pregnancies are likely go into spontaneous labour before 37 weeks however CS in labour is associated with a reduced risk of respiratory disorders. We did not identify any studies that had evaluated the optimal timing for CS in higher order multiple births.

Recommendations

<table>
<thead>
<tr>
<th>Number</th>
<th>Recommendation</th>
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<tbody>
<tr>
<td>11</td>
<td>In otherwise uncomplicated twin pregnancies at term where the presentation of the first twin is cephalic, perinatal morbidity and mortality is increased for the second twin. However, the effect of planned CS in improving outcome for the second twin remains uncertain and therefore CS should not routinely be offered outside a research context.</td>
</tr>
</tbody>
</table>
In twin pregnancies where the first twin is not cephalic the effect of CS in improving outcome is uncertain, but current practice is to offer a planned CS. [GPP 2004]

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<tr>
<th>Number</th>
<th>Research Recommendation</th>
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<tbody>
<tr>
<td>RR 4</td>
<td>RCTs are needed to evaluate the benefits and risks to mothers and babies of CS for delivery of twin and triplet pregnancies.</td>
</tr>
</tbody>
</table>

### 5.3 Preterm birth and CS

Preterm birth is the most common cause of neonatal mortality (47% of neonatal deaths are due to immaturity). Babies born preterm are also at increased risk of morbidity (such as cerebral palsy) however the impact of mode of delivery on outcomes is uncertain. Preterm birth may result from spontaneous preterm labour or because delivery is thought to be beneficial to the mother’s (such as severe pre-eclampsia or HELLP) or baby’s health (for example presumed fetal compromise). Other obstetric complications (such as multiple pregnancies and breech presentation) are associated with preterm birth and will influence the mode of delivery decisions, however detailed discussion of the appropriate management of all these situations is outside the scope of this guideline. Changing the mode of birth for preterm infants to CS has been proposed as a means of reducing the morbidity and mortality however when the infant is very small delivery can be difficult at CS. In addition upper segment caesarean section (classical) may be needed in about 10% of babies born at 27–28 weeks which may have a significant impact on future pregnancies of these women.

A systematic review of planned CS versus expectant management for birth of the small baby identified six RCTs (n = 122). Three RCTs included only breech presentation and three included only cephalic presentations. All trials were discontinued before reaching their projected sample size because of difficulties in recruitment or difficulties in weight estimation where trial entry criteria were based on birthweight. About 1 in 6 of the babies allocated to CS were born vaginally, and vice versa. The findings of the review are inconclusive because there were too few events to give sufficiently precise estimates of effect that would be clinically useful.

A large number of observational studies evaluating mode of birth of preterm infants on mortality and morbidity (such as cerebral palsy) have been published. However the impact of mode delivery on neonatal outcome remains uncertain.

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<th>Recommendation</th>
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<tbody>
<tr>
<td>13</td>
<td>Preterm birth is associated with higher neonatal morbidity and mortality. However, the effect of planned CS in improving these outcomes remains uncertain and therefore CS should not routinely be offered outside a research context. [C 2004]</td>
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<th>Number</th>
<th>Research Recommendation</th>
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</thead>
<tbody>
<tr>
<td>RR 5</td>
<td>RCTs are needed to evaluate the impact of CS on the benefits and risks to mothers and babies born preterm.</td>
</tr>
</tbody>
</table>
5.4 Small for gestational age

Small for gestational age (SGA) refers to a fetus that has failed to achieve a specific biometric measurement (for example abdominal circumference) or estimated weight threshold by a specific gestational age. The commonly used threshold is the tenth centile. About half of these babies are constitutionally small, others are fetuses that are not achieving their growth potential (fetal growth restriction, FGR). SGA fetuses are at greater risk of stillbirth, birth hypoxia, neonatal complications and impaired neurodevelopment. However, most term SGA infants do not have significant morbidity or mortality.\textsuperscript{103} It is beyond the scope of this guideline to consider the investigation and management of small for gestational age infants other than the effect of CS on neonatal outcome, however this topic is covered by another guideline.\textsuperscript{103}

No RCTs were identified that directly reported on baby outcomes for planned CS versus planned vaginal birth for SGA babies. One RCT has compared delayed versus immediate delivery after diagnosis of fetal growth restriction. This trial reported that delayed delivery resulted in fewer CS (OR 2.7, 95% CI 1.6 to 4.5).\textsuperscript{104} [evidence level 1b] Observational data has suggested that SGA babies exposed to labour are more at risk of neonatal death than those not exposed to labour (RR 1.79, 95% CI 1.54 to 1.86).\textsuperscript{105} [evidence level 3] CS may reduce the need for neonatal resuscitation (OR 0.2, 95% CI 0.08 to 0.66).\textsuperscript{106} [evidence level 3]

The effect of CS on cerebral palsy in low birth weight babies is not certain. CS is not associated with a difference in rates of cerebral palsy.\textsuperscript{107,108} [evidence level 3] Currently available guidelines do not recommend a mode of birth for SGA babies.\textsuperscript{103} [evidence level 4]

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<thead>
<tr>
<th>Number</th>
<th>Recommendation</th>
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<tbody>
<tr>
<td>14</td>
<td>The risk of neonatal morbidity and mortality is higher with ‘small for gestational age’ babies. However, the effect of planned CS in improving these outcomes remains uncertain and therefore CS should not routinely be offered outside a research context. [C] [2004]</td>
</tr>
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<tr>
<th>Number</th>
<th>Research Recommendation</th>
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<tbody>
<tr>
<td>RR 6</td>
<td>RCT evidence is needed to determine the effect of planned CS on neonatal mortality and morbidity for ‘small for gestational age’ babies.</td>
</tr>
</tbody>
</table>

5.5 Placenta praevia

Placenta praevia is the primary indication for about 3% of all CS (2.2% not actively bleeding and 0.9% actively bleeding).\textsuperscript{4} The majority of low lying placenta detected at 20 weeks will resolve. If the placenta extends over the os a repeat US should offered at 32 weeks.\textsuperscript{8} (NCC-WCH, 2008) Placenta praevia may also present with painless bleeding. CS is usually necessary when the placenta covers the internal os at 36 weeks (minor or major placenta praevia). Women having a CS for placenta praevia are at increased risk of blood loss of greater than 1000 ml compared to CS for other indications (RR 3.97, 95% CI 3.24 to 4.85).\textsuperscript{4} In the last triennial report from the Confidential Enquiry into Maternal Deaths in the UK, four deaths occurred in women with placenta praevia, three as a result of haemorrhage.\textsuperscript{95} Hence, they should have the CS carried out by an experienced operator with a consultant readily available and at a maternity unit with on-site blood transfusion services.

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\textsuperscript{4} The original guideline indicated that the repeat scan should be offered at 36 weeks. This sentence has now been updated in line with the updated Antenatal Care Guideline (RCOG, 2008).
15. **Women with a placenta that partly or completely covers the internal cervical os (minor or major placenta praevia) should be offered CS. [D][2004]**

### 5.6 Morbidly adherent placenta

#### Introduction

Women who become pregnant after a single previous CS have a 0.6 to 1.3% risk of developing placenta praevia and of these, between 11 and 14% will have a morbidly adherent placenta (Guise et al. 2010). With two previous CS, there is a 1.1 to 2.3% risk of placenta praevia in a subsequent pregnancy and of these; between 23 and 40% will have a morbidly adherent placenta (Guise et al. 2010). With three or more previous caesarean sections, there is a 1.8 to 3.7% risk of placenta praevia in a subsequent pregnancy and of these; between 35 and 67% will have a morbidly adherent placenta (Guise et al. 2010).

Against this backdrop of incremental risk, CS is becoming an increasingly common mode of delivery in the UK, both as a primary and as a repeat procedure. Thus, clinicians can expect to see a gradual increase in the number of women presenting in pregnancy with a morbidly adherent placenta.

Morbidly adherent placenta is associated with serious maternal morbidity including major obstetric haemorrhage, transfusion of large quantities of blood products, hysterectomy and admission to an intensive care unit. However, exsanguination and maternal death from morbidly adherent placenta is now rare in the UK (Cantwell R. et al., 2011). It is hoped that improved prenatal identification of such cases has contributed to this.

This section will review the evidence for the accuracy of imaging techniques in diagnosing morbidly adherent placenta in a pregnant woman with a previous CS who present with placenta praevia. It also reviews the evidence relating to the optimum management once the diagnosis has been made.

#### Accuracy of diagnostic tests

**Review question**

What is the accuracy of imaging techniques (colour-flow ultrasound [US] and magnetic resonance imaging [MRI]) for diagnosis of a morbidly adherent placenta in pregnant women who have had a previous CS and are currently diagnosed with placenta praevia?

**Overview of evidence**

Five studies were included in this review (Warshak et al., 2006; Twickler et al., 2000; Masselli et al., 2008; Shih et al., 2009).

Three studies were conducted in the USA (Warshak et al., 2006; Twickler et al., 2000; Comstock et al., 2009), one in Italy (Masselli et al., 2008), and one in Taiwan (Shih et al., 2009). One retrospective study (Comstock et al., 2009) examined the diagnostic accuracy of transvaginal ultrasound for diagnosis of placenta accreta in pregnant women with antenatal diagnosis of placenta praevia who had prior CS. One retrospective study (Warshak et al., 2006) reported on the diagnostic accuracy of ultrasound (grey scale or colour Doppler) and MRI for diagnosis of placenta accreta in pregnant women with antenatal diagnosis of low anterior placenta and placenta praevia who had had at least one prior CS. One prospective study (Masselli et al., 2008) compared the value of pelvic ultrasound with colour Doppler and MRI for diagnosis of placenta accreta, increta and percreta. One prospective study (Shih et al., 2009) introduced additional criteria for diagnosis of placenta accreta using 3D power Doppler complementary to grey scale and colour Doppler technique. One prospective study (Twickler et al., 2000) evaluated the diagnostic accuracy of Doppler colour flow mapping for diagnosis of placenta accreta in pregnant women with prior caesarean section and diagnosis of anterior low lying placenta and placenta praevia.
<table>
<thead>
<tr>
<th>Table 5.1</th>
<th>GRADE summary of findings for diagnostic accuracy of tests for placenta accreta, increta and percreta</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of studies</td>
<td>Number of women</td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td>Grey scale transabdominal ultrasound (mean gestational age at diagnosis = 30 ± 2.2 weeks)</td>
<td>1 study (Shih et al., 2009)</td>
</tr>
<tr>
<td>Grey scale transvaginal ultrasound (gestational age at diagnosis = 15 to 20 weeks)</td>
<td>1 study (Comstock et al., 2009)</td>
</tr>
<tr>
<td>Grey scale transvaginal ultrasound (gestational age at diagnosis = 15 to 40 weeks)</td>
<td>1 study (Comstock et al., 2009)</td>
</tr>
<tr>
<td>Grey scale or colour Doppler ultrasound (mean gestational age at diagnosis = 25 weeks, range 11 to 37 weeks)</td>
<td>1 study (Warshak et al., 2006)</td>
</tr>
<tr>
<td>US colour Doppler (Masselli et al., 2008; mean gestational age at diagnosis = 30 weeks, range 20 - 37 weeks) (Twickler et al., 2000; gestational age at diagnosis not reported) (Shih et al., 2009; mean gestational age at diagnosis 30 ± 2.2 weeks)</td>
<td>1 study (Masselli et al., 2008)</td>
</tr>
<tr>
<td></td>
<td>1 study (Twickler et al., 2000)</td>
</tr>
<tr>
<td></td>
<td>1 study (Shih et al., 2009)</td>
</tr>
<tr>
<td>MRI (Masselli et al., 2008; mean gestational age at the diagnosis = 30 weeks, range 20 - 37 weeks) (Warshak et al., 2006; mean gestational age at diagnosis = 28 weeks, range 18 to 37 weeks)</td>
<td>1 study (Masselli)</td>
</tr>
</tbody>
</table>
Evidence statements

In the following statements the following definitions have been used when summarising the levels of sensitivity, specificity, positive predictive value (PPV) & negative predictive value (NPV):

- High – 90% and above
- Moderate – 75% to 89%

Evidence was identified using variety of ultrasounds to determine diagnostic accuracy for placenta accreta in women diagnosed with placenta praevia who had at least one prior CS. The quality of the evidence ranged from moderate to low for the included studies.

Grey scale transabdominal ultrasound

One study evaluated the diagnostic accuracy of grey scale ultrasound for placenta accreta in women diagnosed with placenta praevia who had a prior CS. The study reported a high sensitivity, a moderate specificity, a moderate PPV and a high NPV. The evidence for this test was of low quality.

Grey scale transvaginal ultrasound

One study evaluated the diagnostic accuracy of grey scale transvaginal ultrasound for placenta accreta in women at 15 to 20 weeks gestation who were diagnosed with placenta praevia and had a prior CS. The study reported a moderate sensitivity and a low PPV. Specificity and NPV were not reported. The evidence for this test was of low quality.

One study evaluated the diagnostic accuracy of grey scale transvaginal ultrasound for placenta accreta in women at 15 to 40 weeks gestation who were diagnosed with placenta praevia and had a prior CS. The study reported a moderate sensitivity and a low PPV. Specificity and NPV were not reported. The evidence for this test was of low quality.

Grey scale or colour Doppler ultrasound

One study evaluated the diagnostic accuracy of grey scale or colour Doppler ultrasound to diagnose placenta accreta in women diagnosed with placenta praevia who had a prior CS. The study reported moderate sensitivity, high specificity, a low PPV and a high NPV. The evidence for this test was of low quality.

US colour Doppler

Three studies evaluated the diagnostic accuracy of ultrasound colour Doppler to diagnose placenta accreta in women diagnosed with placenta praevia who had a prior CS. One study reported a high sensitivity and specificity with a high PPV and a high NPV. A second study reported high sensitivity, moderate specificity, a moderate PPV and a high NPV. A third study reported a high sensitivity, a low specificity, a moderate PPV and a moderate NPV. The evidence for this test was of moderate quality in the first study, and low quality in the other two studies.

MRI

Two studies evaluated the diagnostic accuracy of MRI to diagnose placenta accreta in women diagnosed with placenta praevia who had a prior CS. The first moderate quality study reported a high
sensitivity, specificity, PPV and NPV, whilst the second reported a moderate sensitivity with a high specificity, high PPV and moderate NPV. The evidence for this test was of moderate quality.

3D power colour sonography

One study evaluated the diagnostic accuracy of 3D power colour sonography to diagnose placenta accreta in women diagnosed with placenta praevia who had a prior CS. The study reported high sensitivity, moderate specificity, a moderate PPV and a high NPV. The evidence for this test was of low quality.

Evidence to recommendations

Relative value placed on the tests considered

The GDG felt that in current practice grey scale ultrasound would not generally be used to make a decision about placenta accreta. Instead the healthcare professional would use colour flow ultrasound in order to highlight movement.

Trade-off between clinical benefits and harms

The group noted evidence from one moderate quality study and one large study of low quality that colour ultrasound is moderately accurate at ruling out morbidly adherent placenta. MRI scan is better for a more complete diagnosis (i.e. considering both accurately ruling in and ruling out morbidly adherent placenta).

The group noted that evidence from one moderate quality prospective study showed that the diagnostic accuracy of MRI was 100% without the use of contrast dye.

However, the GDG also understood that women may not wish to undergo an MRI scan for a number of reasons (such as discomfort at being enclosed in a small space, the risk of supine hypotension, the noise of the machine and the length of the procedure).

The group therefore recognised the importance of discussing the procedure with the woman beforehand, explaining both the potential benefits and risks. This discussion should include an explanation of the degree of accuracy that can be expected, and information that the use of an MRI should enable better accuracy determining the degree of adherence.

The group noted that MRI is more accurate at identifying women who have a morbidly adherent placenta and therefore better able to help decision making around the choice of hospital advised for giving birth (local hospital vs. tertiary centre).

The group agreed with the generally held opinion that both MRI and ultrasound are safe for use in pregnancy.

Trade off between net health benefits and resources

Conclusions relating to cost-effectiveness of diagnosing morbidly adherent placenta are presented below where evidence relating to diagnostic accuracy of ultrasound and MRI and evidence relating to the effectiveness of antenatal diagnosis are considered together to inform the health economic modelling.

Quality of evidence

The group noted that in the majority of the studies, the imaging techniques were carried out prior to 32 weeks gestation, whereas in clinical practice, these scans would be more likely to be carried out after 32 weeks gestation (since the low lying placenta scan won’t generally occur until after a repeat scan for low-lying placenta which is usually carried out at 32-34 weeks).

The group noted that in the Warshak (2006) study looking at MRI, the person interpreting the MRI scans weren’t blinded to the results of the earlier colour ultrasound, thus potentially enhancing the diagnostic accuracy findings in favour of MRI.

The GDG noted that in all of the studies, whilst the high level of suspicion about morbidly adherent placenta in these women might be thought to inflate the figures for diagnostic accuracy, since this is the clinically relevant population, the figures reported are credible when generalised to clinical practice.
Other considerations

The GDG felt that as there was only one study investigating the use of 3D ultrasound, and given that it is not widely available throughout the UK, it was not appropriate to recommend its use.

The group also considered the relevance of evidence reviewed and recommendations to woman with other uterine scars (e.g. myomectomy) but in the absence of evidence pertaining specifically to this group, they did not feel this was possible.

<table>
<thead>
<tr>
<th>Number</th>
<th>Recommendation</th>
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<tbody>
<tr>
<td>16</td>
<td>If low-lying placenta is confirmed at 32-34 weeks in women who have had a previous CS, offer colour-flow doppler ultrasound as the first diagnostic test for morbidly adherent placenta. <strong>[new 2011]</strong></td>
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<tr>
<td>17</td>
<td>If a colour-flow doppler ultrasound scan result suggests morbidly adherent placenta:</td>
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<tr>
<td></td>
<td>- discuss with the woman the improved accuracy of magnetic resonance imaging (MRI) in addition to ultrasound to help diagnose morbidly adherent placenta and clarify the degree of invasion</td>
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<td></td>
<td>- explain what to expect during an MRI procedure</td>
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<td></td>
<td>- inform the woman that current experience suggests that MRI is safe, but that there is a lack of evidence about any long-term risks to the baby</td>
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<tr>
<td></td>
<td>- offer MRI if acceptable to the woman. <strong>[new 2011]</strong></td>
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<tr>
<td>18</td>
<td>Discuss the interventions available for delivery with women suspected to have morbidly adherent placenta, including cross matching of blood and planned CS with a consultant obstetrician present. <strong>[new 2011]</strong></td>
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<table>
<thead>
<tr>
<th>Number</th>
<th>Research Recommendation</th>
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<tbody>
<tr>
<td>RR 7</td>
<td>How accurate is 3D ultrasound compared with 2D ultrasound or MRI scanning for diagnosing morbidly adherent placenta?</td>
<td></td>
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</tbody>
</table>

Effect of diagnosis on outcomes

Review question

Does a diagnosis of morbidity adherent placenta using imaging techniques lead to improved outcomes in pregnant women with a previous caesarean section currently diagnosed with placenta praevia?

Overview of evidence

Two studies were included in this review (Warshak et al., 2009; Wong et al., 2008). One study was conducted in the USA (Warshak et al., 2009) and one in New Zealand (Wong et al., 2008). One observational study (Wong et al., 2008) examined the effects of antenatal diagnosis of placenta accreta on maternal outcomes and the other observational study (Warshak et al., 2009) compared maternal and neonatal outcomes in women with an antenatal diagnosis of placenta accreta (managed by planned caesarean hysterectomy) and those in whom an antenatal diagnosis was not made.

In the US study all women diagnosed with placenta accreta were offered a planned CS with hysterectomy (without attempted removal of the placenta). A caesarean hysterectomy was scheduled for 34 to 35 weeks gestation after a 48 hour course of betamethasone to enhance fetal lung maturity. A multidisciplinary team involving specialists from perinatology, anaesthetics, gynaecological oncology, interventional radiology and neonatology were involved in women’s care. Hysterectomies were performed under general anaesthesia. Internal iliac balloon catheters were passed pre-operatively and inflated during surgery only if significant bleeding was encountered. Most women spent the first day postoperatively in the intensive care unit and stayed longer if clinically indicated. The diagnosis of placenta accreta in all women was confirmed post delivery with a histological test. In the New Zealand study women diagnosed with placenta accreta were offered a planned CS. Five
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women had a hysterectomy and the uterus was conserved in two. No further details are reported regarding the package of care offered.

Maternal outcomes

Table 5.2 GRADE summary of findings for antenatal diagnosis of placenta accreta compared with no antenatal diagnosis (maternal outcomes)

<table>
<thead>
<tr>
<th>Number of studies</th>
<th>Number of women</th>
<th>Effect</th>
<th>Quality</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Antenatal diagnosis of placenta accreta</td>
<td>No antenatal diagnosis of placenta accreta</td>
<td>Relative (95% CI)</td>
</tr>
<tr>
<td>Estimated blood loss</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 study (Warshak et al., 2009)</td>
<td>Mean (litre) ± SD 2.3 ± 1.7 n = 62</td>
<td>Mean (litre) ± SD 2.9 ± 1.8 n = 37</td>
<td>Not calculable (NC)</td>
</tr>
<tr>
<td>1 study (Wong et al., 2008)</td>
<td>Mean (litre) ± SD 1.4 ± 1.0 n = 7</td>
<td>Mean (litre) ± SD 3.6 ± 1.3 n = 9</td>
<td>Not calculable (NC)</td>
</tr>
<tr>
<td>Number of units of blood transfused</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 study (Warshak et al., 2009)</td>
<td>Mean ± SD 4.7 ± 2.2 n = 62</td>
<td>Mean ± SD 6.9 ± 1.8 n = 37</td>
<td>NC</td>
</tr>
<tr>
<td>1 study (Wong et al., 2008)</td>
<td>2.3 ± 2.9 n = 7</td>
<td>5.1 ± 2.9 n = 9</td>
<td>NC</td>
</tr>
<tr>
<td>Emergency hysterectomy</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 study (Wong et al., 2008)</td>
<td>1/7 (14%) n = 7</td>
<td>9/9 (100%) n = 9</td>
<td>RR 0.14 (0.02 to 0.55)</td>
</tr>
<tr>
<td>Intensive care unit [ICU] admission</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 study (Warshak et al., 2009)</td>
<td>43/62 (69%)</td>
<td>22/37 (59%)</td>
<td>RR 1.16 (0.86 to 1.64)*</td>
</tr>
<tr>
<td>1 study (Wong et al., 2008)</td>
<td>1/7 (14%)</td>
<td>1/9 (11%)</td>
<td>RR 1.28 (0.14 to 11)*</td>
</tr>
</tbody>
</table>
### Neonatal outcomes

#### Table 5.3 GRADE summary of findings for antenatal diagnosis of placenta accreta compared with no antenatal diagnosis (neonatal outcomes)

<table>
<thead>
<tr>
<th>Number of studies</th>
<th>Number of neonates</th>
<th>Effect</th>
<th>Quality</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Antenatal diagnosis of placenta accreta</td>
<td>Relative (95% CI)</td>
<td>Absolute (95% CI)</td>
</tr>
<tr>
<td></td>
<td>No antenatal diagnosis of placenta accreta</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Neonatal intensive care unit [NICU] admission</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 study</td>
<td>(Warshak et al., 2009)</td>
<td>50/62 (80%)</td>
<td>19/37 (51%)</td>
</tr>
<tr>
<td>NICU length of stay</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 study</td>
<td>(Warshak et al., 2009)</td>
<td>9.8 days (SD 2.5) n = 62</td>
<td>6.3 days (SD 3.5) n = 37</td>
</tr>
</tbody>
</table>

*Calculated by NCC-WCH technical team
Evidence statements

Maternal outcomes

Estimated blood loss

One study found that the mean blood loss in women with an antenatal diagnosis of placenta accreta was lower than in women with no antenatal diagnosis of placenta accreta. This finding was statistically significant. A second study did not find a statistically significant difference for this outcome. The evidence for this outcome was of very low quality.

Number of units of blood transfused

One study found that women with an antenatal diagnosis of placenta accreta had a lower number of packed red blood cell transfusions compared with the women who had no antenatal diagnosis of placenta accreta. This finding was statistically significant. A second study did not find a statistically significant difference for this outcome. The evidence for this outcome was of low quality.

Emergency hysterectomy

One study found that the incidence of emergency hysterectomy was lower among women with an antenatal diagnosis of placenta accreta compared with those who had no antenatal diagnosis of placenta accreta. This finding was statistically significant. The evidence for this outcome was of very low quality.

Intensive care (ICU) admission

Two studies did not find a statistically significant difference in the rate of ICU admission for women with an antenatal diagnosis of placenta accreta compared with women who had no antenatal diagnosis of placenta accreta. The evidence for this outcome was of very low quality.

Length of hospital stay

Two studies did not find a statistically significant difference in length of hospital stay for women with an antenatal diagnosis of placenta accreta compared with women who had no antenatal diagnosis of placenta accreta. The evidence for this outcome was of very low quality.

Bladder injuries

Two studies did not find a statistically significant difference in the rate of bladder injuries for women with an antenatal diagnosis of placenta accreta compared with women who had no antenatal diagnosis of placenta accreta. The evidence for this outcome was of very low quality.

Neonatal outcomes

Neonatal intensive care unit (NICU) admission

One study found that the rate of NICU admission was higher in neonates born to women with an antenatal diagnosis of placenta accreta than in neonates born to women with no antenatal diagnosis of placenta accreta. This finding was statistically significant. The evidence for this outcome was of very low quality.

NICU length of stay

One study did not find a statistically significant difference in NICU length of stay for neonates born to women with an antenatal diagnosis of placenta accreta compared to neonates born to women with no antenatal diagnosis of placenta accreta. The evidence for this outcome was of very low quality.

Health economics

A de novo model to compare different diagnostic strategies for morbidly adherent placenta in praevia was developed for this guideline. The results of this analysis are summarised here; further details are provided in chapter 13.

The model compared the following diagnostic strategies:

i) None

ii) Ultrasound

iii) MRI

iv) Ultrasound followed by MRI in ultrasound test positives
There is an absence of evidence about how much a diagnosis of morbidly adherent placenta leads to improved outcomes. Even if it does, the “downstream” saving and QALY gain from averting “adverse outcomes” are further unknowns. Therefore, the model took a “what-if” approach to assess what would be considered cost-effective in different scenarios. There is also some uncertainty about the precise diagnostic accuracy of the different diagnostic tests although there is at least some evidence for these, which perhaps make this uncertainty of secondary importance in terms of making guideline recommendations.

The model suggested that a diagnostic strategy of ultrasound alone was dominated by other alternatives, which meant that other strategies were likely to be cheaper and more effective. Although it has the lowest diagnostic cost, the high cost of false positives in a low prevalence population makes it the most expensive strategy overall. Furthermore, the evidence suggests that such a strategy would miss more cases than a strategy of MRI alone. This finding did not depend on assumptions about improved outcomes arising from the detection of cases.

The “what-if” analysis started from the premise that identifying cases would lead to improved outcomes. Although there is an absence of evidence for this and an effect size can’t be estimated, the GDG were strongly of the opinion that “being prepared” offered some protection from risk. Under that premise there were scenarios where ‘do nothing’, ultrasound+MRI and MRI alone could be considered cost-effective. However, in general a much lower effect size, QALY gain and “downstream” cost saving from averting “adverse outcomes” was necessary for ultrasound+MRI to be cost-effective than for MRI alone.

Evidence to recommendations
Relative value placed on the outcomes considered
The GDG recognised that although blood loss is an important outcome, the way that it has been reported is not useful for their decision making. The group were particularly concerned about identifying the number of women where blood loss could be potentially life-threatening but this was not reported.

The group recognised that although the differences in the rates of hysterectomies appeared to be a significant finding, these results were due to the local protocol at the hospital (i.e. where a placenta accreta was first discovered when performing the CS, the clinicians would perform an elective hysterectomy). The group felt that this was not a common approach and so did not wish to place any value on the differences reported.

The group did not feel that the findings related to ICU admission and length of hospital stay, for both women and neonates, were particularly helpful as the decision about length of stay will often be determined by local protocols.

The group felt that the number of bladder injuries was an important outcome. However, they recognised that neither of the studies showed a statistically significant difference for this outcome.

Trade-off between clinical benefits and harms
The group believed from their experience that the main benefit of diagnosing a morbidly adherent placenta is that this allows clinicians to be prepared, and to ensure that appropriate measures are taken in cases of extreme blood loss. These include ensuring that there is sufficient cross-matched blood available and that experienced specialist clinicians are available to provide support when needed.

Trade-off between net health benefits and resource use
There is insufficient evidence to determine the cost-effectiveness of different diagnostic strategies for morbidly adherent placenta. The modelling undertaken for this guideline suggested that ultrasound alone was likely to be dominated because of a higher false positive rate. There is some evidence to support a view that ultrasound has a lower specificity than MRI although it is not conclusive. The model also suggested that a sequential strategy of ultrasound followed by a confirmatory MRI in ultrasound test positives is cheaper than a strategy of MRI alone. This is because the sequential MRI test removes the costs of false positives which more than offsets the costs associated with an additional test. The sequential strategy involves a much smaller number of MRI scans than with a strategy based on MRI alone and because of the substantial difference in costs between an
ultrasound and MRI this means that the sequential strategy has markedly lower diagnostic costs even if the absolute number of tests undertaken is higher.

The question then is whether the additional costs associated with ultrasound+MRI or MRI alone is worth the additional costs. The evidence doesn’t exist to answer this question and the model therefore took a “what-if” approach. The GDG were strongly of the opinion that identifying cases was likely to lead to better outcomes. The model suggested that much smaller gains were necessary for ultrasound+MRI to reach a cost-effectiveness threshold relative to “do-nothing” than for MRI alone to be considered “cost-effective” relative to ultrasound+MRI. Therefore, it would be difficult to justify a recommendation for a diagnostic strategy of MRI alone given existing evidence. Such a strategy is not common in current UK practice and there could be capacity issues which would hinder the implementation of such a recommendation. Although current UK practice varies, ultrasound+MRI is an approach used in some centres. Whilst further evidence is required, a recommendation of ultrasound+MRI seems to make pragmatic sense given current practice, GDG opinion and the insights available from the model produced for this guideline.

Quality of evidence

The group recognised that there were only two studies which provided evidence for this question, and that the quality of the evidence for the findings from these studies was very low. They noted that one of the studies only contained a small number of women in each arm and so was likely to be underpowered for rare outcomes such as bladder injury.

Given the poor quality of the evidence and lack of detail in one study about the specific management regimes used, the group did not feel able to make a strong recommendation for specific interventions. It was noted that in one study (Warshak, 2009) the management strategy of elective caesarean hysterectomy was used for all women, an approach which is not usual practice and thus findings from this study are not generalisable to situations where conservative management is undertaken.

In light of the large amount of blood loss associated with both arms of each study, the GDG agreed that there were steps that should be taken to minimise morbidity associated with this.

Other considerations

The group recognised that there are a number of interventions which are used in to reduce blood loss during surgery such as balloon catheters and interventional radiology. In addition, some trusts have cell salvage equipment available which can also be used to reduce the need for cross-matched blood. There is variation in practice concerning the use of these interventions in the management of morbidly adherent placenta and a lack of evidence to support their use. Consequently the GDG felt it important to recommend that further research is conducted in this area.

<table>
<thead>
<tr>
<th>Number</th>
<th>Recommendation</th>
</tr>
</thead>
</table>
| 19     | When performing a CS for women suspected to have morbidly adherent placenta, ensure that:  
  - a consultant obstetrician and a consultant anaesthetist are present  
  - an experienced paediatrician is present  
  - a senior haematologist is available for advice  
  - a critical care bed is available  
  - sufficient cross-matched blood and blood products are readily available. [new 2011] |
<p>| 20     | When performing a CS for women suspected to have morbidly adherent placenta the consultant obstetrician should decide which other healthcare professionals need to be consulted or present. [new 2011] |
| 21     | All hospitals should have a locally agreed protocol for managing morbidly adherent placenta that sets out how these elements of care should be provided. [new 2011] |</p>
<table>
<thead>
<tr>
<th>Number</th>
<th>Research Recommendation</th>
</tr>
</thead>
<tbody>
<tr>
<td>RR 8</td>
<td>What is the effectiveness of procoagulant agents (such as recombinant factor VIIa, beriplex, tranexamic acid, fibrinogen concentrate) in reducing blood loss in women with morbidly adherent placenta?</td>
</tr>
<tr>
<td>RR 9</td>
<td>What is the effectiveness of point of care testing for haematological indices in women with an established postpartum haemorrhage and in cases of morbidly adherent placenta in reducing maternal morbidity?</td>
</tr>
<tr>
<td>RR 10</td>
<td>What is the effectiveness of the components of the package of care for morbidly adherent placenta such as imaging techniques (e.g. interventional radiology including balloon catheters), stenting of ureters, removal of the placenta, and cell salvage in reducing morbidity associated with maternal blood loss?</td>
</tr>
<tr>
<td>RR 11</td>
<td>What is the appropriate gestational age of elective birth for babies of women with a morbidly adherent placenta</td>
</tr>
<tr>
<td>RR 12</td>
<td>What is the effectiveness of performing an elective hysterectomy to reduce morbidity associated with blood loss in women with morbidly adherent placenta?</td>
</tr>
</tbody>
</table>

5.7 Predicting CS for cephalopelvic disproportion in labour

Pelvimetry (clinical or X-ray) has been used to predict the need for CS in pregnant women. A systematic review of 4 RCTs (n = 895) assessed the effects of x ray pelvimetry on mode of birth. Two RCTs included women with a previous CS. The women on whom pelvimetry was performed were more likely to be delivered by CS (Peto OR 2.17, 95% CI 1.63 to 2.88); There were no differences in neonatal outcomes (asphyxia, admission to neonatal unit, scar dehiscence).\[109\] [evidence level 1a]
Guidelines have recommended that pelvimetry is not used except in rare circumstances such as if the woman has had a previous fracture of the pelvis.\[110\]
Other tests to predict failure to progress (FTP) have included shoe size, maternal height and size of fetus. Observational studies have not demonstrated their value in predicting FTP in labour.\[111,112\] [evidence level 3]

<table>
<thead>
<tr>
<th>Number</th>
<th>Recommendation</th>
</tr>
</thead>
<tbody>
<tr>
<td>22</td>
<td>Pelvimetry is not useful in predicting ‘failure to progress’ in labour and should not be used in decision making about mode of birth. [A] [2004]</td>
</tr>
<tr>
<td>23</td>
<td>Shoe size, maternal height and estimations of fetal size (ultrasound or clinical examination) do not accurately predict cephalopelvic disproportion and should not be used to predict ‘failure to progress’ during labour. [B] [2004]</td>
</tr>
</tbody>
</table>

5.8 Mother-to-child transmission of maternal infections

This section addresses CS as an intervention to reduce mother-to-child transmission (MTCT) of viral infections (such as human immunodeficiency virus [HIV]), other interventions also impact on the risk of the MTCT of viral infections (such as anti-retrovirals for HIV) but these topics are outside the scope of this guideline.
HIV

Introduction

Approximately 86,500 people in the UK are carriers of the Human Immunodeficiency Virus (Health Protection Agency, 2010). In the pregnant population about 1 in 450 pregnant women nationally are HIV positive and this rises to about 1 in 250 pregnant women in London. In general, around one quarter of all people carrying the virus is unaware of her or his HIV positive status.

Current NICE guidelines for routine antenatal care recommend that screening for HIV should be offered to all pregnant women in the UK because various interventions can decrease the maternal to fetal (vertical) transmission of the virus (NCC-WCH, 2008). The previous version of this guideline recommended that ‘HIV-positive women who are pregnant should be offered a planned CS because it reduces the risk of mother-to-child transmission of HIV’. However, since the publication of the original guideline there has been a growing body of evidence suggesting that for some women taking antiretroviral therapy (ART) or highly active antiretroviral therapy (HAART), the chance of vertical transmission is reduced so effectively that CS may no longer associated with reduced vertical transmission rates compared to vaginal birth, even in the presence of a detectable viral load.

This new evidence is reviewed in this section, together with an update of the recommendations for clinical practice.

Review question

What is the effectiveness of planned caesarean section compared with planned vaginal birth at decreasing the mother-to-child transmission of the virus in pregnant women with HIV, for both low and high viral load?

Overview of evidence

Four studies (Boer et al., 2010; Warszawski et al., 2008; Islam et al., 2010; Townsend et al., 2008) were included in this review which investigated the effectiveness of planned CS compared with vaginal birth at decreasing the MTCT of the virus in pregnant women with HIV for both low and high viral load.

One study (Boer et al., 2010) was a prospective observational study; three (Townsend et al., 2008; Warszawski et al., 2008; Islam et al., 2010) were retrospective observational studies. Two of the studies (Islam et al., 2010; Townsend et al., 2008) were conducted in the UK, one in France (Warszawski et al., 2008) and one (Boer et al., 2010) in eight Western European countries (Italy, Spain, Belgium, Netherlands, UK, Germany, Denmark and Sweden). All four studies reported mother to child transmission rate, viral load count and mode of birth. Two studies (Islam et al., 2010; Townsend et al., 2008) investigated planned CS versus planned vaginal birth and the other two (Boer et al., 2010; Warszawski et al., 2008) compared planned CS with vaginal birth (planned and unplanned).

Three studies (Boer et al., 2010; Islam et al., 2010; Townsend et al., 2008) reported on mother to child transmission rate and mode of birth in HIV infected pregnant women with low viral load defined as undetectable viral load (< 50 copies/ml). One study (Warszawski et al., 2008) reported on mother to child transmission rate and mode of birth in women with low viral load defined as < 400 copies/ml.

The table below is the summary of the evidence by mother to child transmission rate for all the published studies identified for this review question. The tables have been divided up by plasma viral load count.

<table>
<thead>
<tr>
<th>Number of studies</th>
<th>Number of women</th>
<th>Effect</th>
<th>Quality</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Planned CS</td>
<td>Vaginal birth</td>
<td>Relative (95% CI)</td>
</tr>
<tr>
<td>Mother to child transmission (MTCT) in women with viral load &lt; 50 copies/ml on highly active antiretroviral therapy (HAART)</td>
<td>1 study</td>
<td>1/238</td>
<td>1/321</td>
</tr>
<tr>
<td>Study</td>
<td>MTCT in women with viral load &lt; 50 copies/ml on HAART</td>
<td>MTCT in women with viral load ≥ 50 and &lt; 1000 copies/ml on HAART</td>
<td>MTCT in women with viral load &lt; 400 copies/ml with and without HAART</td>
</tr>
<tr>
<td>-------</td>
<td>----------------------------------------------------</td>
<td>---------------------------------------------------------------</td>
<td>---------------------------------------------------------------</td>
</tr>
<tr>
<td>1 study Countries Townsend et al. (2008)</td>
<td>2/1135 (0.2%)</td>
<td>4/417 (0.95%)</td>
<td>1/242 (4.5%)</td>
</tr>
<tr>
<td>1 study Countries Islam et al. (2010)</td>
<td>1/23 (0.08%)</td>
<td>2/81 (2.5%)</td>
<td>OR 0.39 (0.07 to 2.17)**</td>
</tr>
<tr>
<td>1 study Countries Boer et al. (2010)</td>
<td>OR 0.73 (0.06 to 8.12)**</td>
<td>NC</td>
<td>NC</td>
</tr>
<tr>
<td>1 study Countries Townsend et al. (2008)</td>
<td>0/23 (0%)</td>
<td>OR 0.39 (0.07 to 2.17)**</td>
<td>NC</td>
</tr>
<tr>
<td>1 study Countries Islam et al. (2010)</td>
<td>Not reported (NR)</td>
<td>Not reported (NR)</td>
<td>NC</td>
</tr>
</tbody>
</table>

1. *planned vaginal birth
2. **calculated by NCC technical team

3. **Evidence statements**
4. Mother to child transmission (MTCT) in women with viral load < 50 copies/ml on HAART
Two studies did not find a statistically significant difference in the rates of mother to child transmission in women on HAART having a planned CS compared with those having a vaginal birth (planned or unplanned; including intrapartum CS) with viral load < 50 copies/ml.

One study found no reported incidences of mother to child transmission of HIV for women having planned vaginal birth in women with viral load < 50 copies/ml, of whom 14/23 were on HAART. The evidence for this outcome was of very low quality.

**MTCT in women with viral load ≥ 50 and < 1000 copies/ml on HAART**

One study did not find a statistically significant difference between transmission rates for women on HAART with viral load ≥ 50 and < 1000 copies/ml having a planned CS compared with those having a planned vaginal birth. The evidence for this outcome was of very low quality.

**MTCT in women with viral load < 400 copies/ml with and without HAART**

One study found that planned CS reduces the risk of mother to child transmission compared with vaginal birth (planned or unplanned; including intrapartum CS) when maternal viral load is < 400 copies/ml (some women receiving HAART, some receiving ART, a few receiving no ART, numbers not reported). This finding was statistically significant. The evidence for this outcome was of very low quality.

**MTCT in women with viral load < 400 copies/ml on ART**

One study did not find a statistically significant difference in transmission rate for women on ART with a viral load < 400 copies/ml having a planned CS compared with those having a planned vaginal birth.

The evidence for this outcome was of very low quality.

**MTCT in women with viral load < 1000 copies/ml on HAART**

One study reported transmission rates in women on HAART with a viral load < 1000 copies/ml. This was slightly higher for women having a planned CS but the statistical significance was not calculable.

The evidence for this outcome was of very low quality.

**MTCT in women with viral load ≥ 1000 copies/ml on ART (including HAART)**

One study did not find a statistically significant difference between transmission rates for women on ART (including HAART) with a viral load ≥ 1000 copies/ml having a planned CS compared with those having a planned vaginal birth.

The evidence for this outcome was of very low quality.

**Evidence to recommendations**

Relative value placed on outcomes considered

The only relevant outcome for consideration in this question was the rate of mother to child transmission of HIV. This outcome was then split according to the viral load (number of copies per ml) reported in each study (and in some cases further subdivided according to the treatment used).

Trade-off between clinical benefits and harms

The evidence provided a number of results, both for different viral loads and different therapies. For women with a viral load of < 50 copies/ml on HAART, there was no significant difference in mother to child transmission rates between women who gave birth vaginally and those who gave birth by CS.

The group felt that this matched their clinical experience and so were confident in recommending that women on HAART with a viral load of < 50 copies/ml should not be offered a CS.

The group noted that one study (Warszawski et al., 2008) with the majority of women on ART did not show a statistically significant difference in the mother to child transmission rate between women giving birth vaginally and those giving birth by CS in women with a viral load of < 400 copies/ml. However, the group recognised that the study did include a number of women on HAART and that this might have affected the results. As a result the group agreed that either a vaginal birth or CS could be considered for women on ART with a viral load from 50-400 copies/ml.
The evidence also suggested that for women on HAART with a viral load between 50 copies/ml and 1000 copies/ml there was no significant difference in transmission rates between women who gave birth vaginally and those who gave birth by CS. Further discussion of all the evidence pertaining to viral loads of greater than 400 copies/ml led the group to decide that the evidence was of too low a quality to change current practice and they felt it important to remain cautious in these instances. They ultimately agreed that women on HAART with a viral load less than 400 copies/ml should not be offered a CS on the grounds of their HIV status but that all women with a viral load > 400 copies/ml should be offered a CS regardless of the therapy being received.

One study (Boer et al., 2010) which included women receiving different therapies showed a significant reduction in the risk of mother to child transmission with planned CS. The GDG felt that this difference was likely to be due to the fact that the study included women receiving no therapy. As a result, the group agreed that it was appropriate to adopt a cautious approach and recommended that women on no therapy should be offered a CS regardless of their viral load.

### Trade-off between health benefits and resources

The economic modelling for CS compared with vaginal birth for women with healthy, uncomplicated pregnancy suggests vaginal birth may be more cost effective, although the evidence for this is not strong and based upon incomplete outcome data. However, where a woman is HIV positive the balance may be tipped in favour of CS in order to reduce the risk of mother to child transmission of HIV and the subsequent long term health loss and treatment costs for the baby. The point at which the balance favours CS above vaginal birth depends upon the relative risk of HIV transmission. The GDG decided, based on the evidence reviewed plus their own experience, that this tipping point comes at viral loads of above 400 copies/ml regardless of therapy. The use of HAART reduces the risk of transmission and so where there is low viral load (< 400 copies/ml) it is appropriate to advise vaginal birth.

### Quality of evidence

All evidence reviewed for this question was of very low quality. Most often this was due to the studies being retrospective observational studies, underpowered and with flaws in reporting meaning it was not always possible to determine actual mode of birth for all women within each study group. The low quality of evidence meant the GDG remained cautious with their recommendations.

### Other considerations

The group agreed that further UK-based data about the diagnosis of HIV in pregnant women, its treatment, the mode of birth chosen in different circumstances, and mother-to-child transmission rates were all required. The group was aware that some of this information is currently collected by the RCOG and so included a recommendation that this data continue to be collected.

<table>
<thead>
<tr>
<th>Number</th>
<th>Recommendation</th>
</tr>
</thead>
<tbody>
<tr>
<td>24</td>
<td>As early as possible give women with HIV information about the risks and benefits for them and their child of the HIV treatment options and mode of birth so that they can make an informed decision. [new 2011]</td>
</tr>
</tbody>
</table>
| 25     | Do not offer a CS on the grounds of HIV status to prevent mother-to-child transmission of HIV to:  
  
  - women on highly active anti-retroviral therapy (HAART) with a viral load of less than 400 copies per ml or  
  - women on any anti-retroviral therapy with a viral load of less than 50 copies per ml.  
  
  Inform women that in these circumstances the risk of HIV transmission is the same for a CS and a vaginal birth. [new 2011] |
| 26     | Consider either a vaginal birth or a CS for women on anti-retroviral therapy (ART) with a viral load 50-400 copies per ml because there is insufficient evidence that a CS prevents mother-to-child transmission of HIV. [new 2011] |
Offer CS to women with HIV who:

- are not receiving any anti-retroviral therapy or
- are receiving any anti-retroviral therapy and have a viral load of 400 copies per ml or more. [new 2011]

Researchers and national bodies responsible for the collection of UK population data should continue to collect data about HIV diagnoses in pregnant women, including treatment, mode of birth, and mother-to-child transmission rates. [new 2011]

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**Hepatitis B virus**

Serological screening for hepatitis B should be offered to all pregnant women. The prevalence of hepatitis B surface antigen (HBsAg) in pregnant women in the UK has been found to range from 0.5 to 1% [evidence level 3]. The wide range in prevalence rates is most likely due to wide variation in prevalence among different ethnic groups. [evidence level 3]

Hepatitis B immunoglobulin and hepatitis B vaccine reduce mother-to-child transmission (MTCT). The vaccine and immunoglobulin are given to the infant at birth followed by either a one month and six month dose or at 5 weekly intervals [evidence level 1b]

Most MTCT occurs at birth or postnataally. Transmission at birth may be due to microperfusion of maternal blood into the infant's circulation during placental separation or by the infant swallowing maternal blood, amniotic fluid or vaginal secretions at vaginal birth. [evidence level 3]

It has been suggested that CS could further reduce MTCT however no RCTs have addressed this issue. One cohort study was identified (n = 447 infants). The methodology of this study is not clearly reported and the generalisability of the findings is not clear. [evidence level 2a]

<table>
<thead>
<tr>
<th>Number</th>
<th>Recommendation</th>
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<tbody>
<tr>
<td>29</td>
<td>Mother-to-child transmission of hepatitis B can be reduced if the baby receives immunoglobulin and vaccination. In these situations pregnant women with hepatitis B should not be offered a planned CS because there is insufficient evidence that this reduces mother-to-child transmission of hepatitis B virus. [B] [2004]</td>
</tr>
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**Hepatitis C virus**

Women are not routinely offered screening for hepatitis C infection in the UK. The prevalence of hepatitis C virus (HCV) in women of child-bearing age is not known as large scale serological studies have not been done. It is however estimated that 1–2% of women of child-bearing age in the US are positive for antibody to HCV. An estimate for EU countries is 0.9% (0.1–3%).

Mother-to-child transmission (MTCT) of HCV was first described in the early 1990s and is now widely recognized. The risk of MTCT of HCV is usually low at 3–5% but higher rates of 10–20% are observed among HIV co-infected women. [evidence level 3] A cohort study involving 441 mother-child pairs from the UK and Ireland of which 5% were known to be HIV-positive, estimated overall MTCT risk at 6.7% (95% CI 4.1 to 10.2). Women co-infected with HIV and HCV had a 3.8 times higher risk of transmitting HCV to their infants than HIV-negative women. [evidence level 2b]

The effect of mode of birth on the risk of MTCT of HCV has not been evaluated in RCTs. We identified a pooled retrospective analysis of prospectively collected data on 1474 HCV infected...
women from 36 centres in eight European countries.\textsuperscript{138} [evidence level 3] For women with hepatitis C infection, there was no difference in risk of vertical transmission by mode of birth (OR 1.19, 95% CI 0.64 to 2.20). This lack of association persisted with adjustment for breastfeeding status, geographic region and maternal age at birth (OR 1.26, 95% CI 0.68 to 2.34), (OR 1.29, 95% CI 0.69 to 2.42) and (OR 1.17, 95% CI 0.59 to 2.31).\textsuperscript{138}[evidence level 3]

Within this study subgroup analysis of women co-infected with HIV (n = 503, 35.4%), reported that the risk of vertical transmission for HCV was reduced by 60% with CS (OR 0.43, 95% CI 0.23 to 0.80). Of the HIV co-infected women, 14 (7.3%) were classified as clinical stage C, the remainder of the women are described as being asymptomatic. There is no mention of whether any of the women were on anti-retroviral therapy. Thirteen (2.6%) of the HIV co-infected women breastfed their infants.\textsuperscript{138} [evidence level 3]

<table>
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<tr>
<th>Number</th>
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<tbody>
<tr>
<td>30</td>
<td>Women who are infected with hepatitis C should not be offered a planned CS because this does not reduce mother-to-child transmission of the virus. [C] [2004]</td>
</tr>
<tr>
<td>31</td>
<td>Pregnant women who are co-infected with hepatitis C virus and HIV should be offered planned CS because it reduces mother-to-child transmission of both hepatitis C virus and HIV. [C] [2004]</td>
</tr>
</tbody>
</table>

**Genital herpes simplex virus**

Genital herpes simplex virus (HSV) infection is an ulcerative sexually transmitted infection which can recur and is associated with considerable physical and psychological morbidity. Genital ulcers may cause pain but can be asymptomatic (for example cervical lesions). Between 1972 and 2001, there was a 9–fold increase in the incidence of genital HSV diagnosed in women in the UK.\textsuperscript{139} [evidence level 3] Currently HSV-2 antibody prevalence in England and Wales is 3% in men and 5% in women.\textsuperscript{140} [evidence level 3]

Neonatal HSV can cause severe systemic disease and is associated with a high mortality rate. Active surveillance in the UK suggests that neonatal HSV infection occurs in 1.65 per 100,000 live births.\textsuperscript{141} [evidence level 3] Neonatal HSV may result from contact of the newborn with the birth canal of an infected mother.

**Primary HSV infection and MTCT of HSV**

The accepted practice of offering CS to women with HSV infection is based on three case series. The first study included 101 pregnant women with HSV (both primary and recurrent disease). This study found the risk of neonatal herpes to be highest for women who acquired primary infection during the third trimester (3 cases of neonatal infection out of 9 cases of exposure).\textsuperscript{142} [evidence level 3] Subsequently a study evaluating screening for HSV identified 94 women who acquired HSV during pregnancy but with no MTCT to the infants. There were an additional 9 women who acquired genital HSV near the onset of labour and in this group, 4 of the 9 infants developed neonatal HSV infection.\textsuperscript{143} [evidence level 3] A study of 15,923 asymptomatic women in early labour reported isolating HSV from 56 women of whom 18 (35%) had a primary infection. Neonatal HSV developed in 6 infants (33%).\textsuperscript{144} [evidence level 3] None of the studies are large enough to address the effect of mode of birth on MTCT.

Despite limited evidence the high mortality associated with neonatal herpes means there is consensus about current practice to offer CS for primary infection.\textsuperscript{145,146}

**Recurrent HSV infection and history of HSV infection and MTCT**

Observational data suggests that the risk of neonatal infection with recurrent HSV is lower than with primary HSV infection (8% with recurrent infection and 33% with primary HSV infection).\textsuperscript{147,148} [evidence level 3] In the Netherlands there has not been a policy of CS for women with recurrent HSV
149 Recurrent HSV may not cause symptomatic lesions, for example with cervical ulceration. A study of 15923 asymptomatic women in early labour reported isolating HSV from 34 women, neonatal HSV developed in 1 of the infants (3%).\footnote{144 [evidence level 3]} To prevent MTCT of HSV in asymptomatic women antenatal screening using HSV cultures was proposed, but this test also did not predict infants risk at birth.\footnote{150 [evidence level 3]}

Three RCTs evaluate using oral acyclovir from 36 weeks to prevent recurrence of HSV at the time of birth. These found a reduction in CS for HSV; however do not report the effect of acyclovir on MTCT.\footnote{151–153 [evidence level 1b]}

A survey of obstetricians in the UK found there was no consensus of opinion or practice for recurrent disease or a history of disease.\footnote{146 [evidence level 3]}

### Cost effectiveness of CS to prevent MTCT of HSV

Three American studies have considered the factors that promote or inhibit the cost-effectiveness of various strategies to prevent MTCT of HSV.\footnote{154–156 Two studies by the same author have examined the additional efficacy, risks, and costs of CS for three groups of women: those presenting with primary HSV; women with a history of HSV; and women with no clinical HSV or history of HSV. The first study was a decision analytic model using data from a review of 19 studies.\footnote{154 [evidence level 3]} Marginal (additional) costs and benefits over and above standard delivery were calculated.}

Adopting a programme of offering routine CS for women with a history of HSV, 9 neonatal cases would be averted per million births at an estimated cost of US$2.5 million per case of neonatal HSV averted. For women with primary HSV, 18 neonatal cases prevented per million with estimated cost saving of US$38,000 per case of neonatal HSV averted.\footnote{154 However more data on transmission rates and the efficacy of CS are required to make these estimates robust.}

A later study\footnote{155} modelled the cost-effectiveness of four strategies to prevent MTCT of HSV in women with at least one previous episode of HSV. CS only, acyclovir prophylaxis in late pregnancy with vaginal birth, acyclovir prophylaxis in late pregnancy with screening and follow-up, and a ‘do nothing’ option. The incremental cost per case prevented compared with ‘do nothing’ was highest for CS with 2.8 cases prevented at an additional cost of US$1.3 million, and lowest for acyclovir prophylaxis with screening and follow-up of neonates (an additional cost of US$400,300). This suggests that acyclovir therapy with follow-up was a more cost-effective strategy than CS alone.

The third paper examined whether acyclovir suppression was a more cost-effective option compared to offering CS only to women with a history of HSV.\footnote{156 The analysis showed that CS rate was the most sensitive variable (since it represents a high proportion of the total costs). The authors concluded that acyclovir suppression was a cost-effective alternative to CS for women with a history of genital herpes in agreement with analysis of the authors of the previous two papers. However, given the lack of data around the estimates of costs, the small sample size (46 women presenting with HSV or with a history of HSV) and the setting of the study, the findings are of limited value to this guideline.}

In conclusion CS is the preferred (the most cost-effective and cost-saving) option in women presenting with primary HSV late in pregnancy. Acyclovir prophylaxis may be a more cost-effective option for women with recurrent HSV.

<table>
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<tr>
<th>Number</th>
<th>Recommendation</th>
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<tbody>
<tr>
<td>32</td>
<td>Women with primary genital herpes simplex virus (HSV) infection occurring in the third trimester of pregnancy should be offered planned CS because it decreases the risk of neonatal HSV infection. [C] [2004]</td>
</tr>
<tr>
<td>33</td>
<td>Pregnant women with a recurrence of HSV at birth should be informed that there is uncertainty about the effect of planned CS in reducing the risk of neonatal HSV infection. Therefore, CS should not routinely be offered outside a research context.</td>
</tr>
</tbody>
</table>
Number Research Recommendation
RR 14 RCTs are needed to determine whether planned CS should be offered to prevent MTCT of HSV to women with recurrence of HSV at birth and in women in whom the primary HSV infection occurs in the first trimester of pregnancy.

5.9 Maternal request for CS

Introduction
In general, CS is a safe operation, especially when performed as planned procedure. CS rates are rising worldwide, with an increasing proportion being undertaken for maternal request, in contrast to those that are performed for obstetric indications. There are many reasons for such requests but these are not always revealed by the women or adequately explored and clearly documented by their carers. This chapter addresses the issue of CS requested by women who have no apparent clinical reason or who report fear of giving birth vaginally.

Rates of maternal request for CS
We identified 19 observational studies that report rates of maternal request for CS. Twelve of these are included in a systematic review (n = 13285) and 7 studies have been published since the review. The largest of these studies were a survey of women attending antenatal clinics in Sweden (n = 3061) and a survey of women’s views of childbirth carried out within the National Sentinel Caesarean Section Audit (NSCSA) (n = 2475).

The rates of preference for CS expressed by the women that were surveyed during pregnancy in UK, Australia and Sweden range from 6% - 8%. Within these studies there was a consistent relationship between women’s preference for CS and either previous CS, previous negative birth experience, a complication in the current pregnancy or a fear of giving birth. The main reason given for preference for CS was that it was perceived to be safest for the baby. The main reason given by those who expressed a preference for vaginal birth was the experience of a natural event. One study concluded that maternal request for CS seems to be a marker for previously negative birth experiences and should prompt enquiries to address any issues or concerns.

Fear of childbirth
It is estimated that about 6%--10% of pregnant women experience fear of childbirth. Fears concerning childbirth such as pain, obstetric injury, unplanned CS, health care staff and the effects on family life have been reported to be more common among primiparous compared to multiparous women, and among those who had not attended antenatal classes. Fear of health care workers was reported to be more common among women who either had problems in the current pregnancy or those who were intending a planned CS. Manifestations of this fear included stress symptoms influencing everyday life, nightmares, a wish to have CS and a wish to avoid the current pregnancy and childbirth.

Fear of childbirth has been measured using different scoring systems. One case–control study found that women who requested planned CS due to fear of child birth were more likely to have also experienced a spontaneous miscarriage (OR 1.73, 95% CI 1.05 to 2.85), a longer time between pregnancies (OR 1.44, 95% CI 1.19 to 1.75), a longer duration of second stage of labour and a previous assisted vaginal birth (OR 4.50, 95% CI 2.18 to 9.31) or “emergency” CS (OR 26.91, 95% CI 11.86 to 61.07).
duration or intervention in the third stage of labour in a previous pregnancy were not found to be associated with fear of childbirth in this study.\textsuperscript{166}

Another study reported that women who had “emergency” CS had higher scores for fear of childbirth during pregnancy compared to those who had vaginal births.\textsuperscript{167} However a prospective study carried out in the U.K. did not find an association between fear of childbirth and “emergency” CS (OR 1.00, 95% CI 0.98 to 1.01).\textsuperscript{26} [evidence level 3]

One RCT randomised women referred to an antenatal clinic for fear of childbirth to receive either cognitive behavioural therapy or usual care. No difference was detected in the proportion of women who chose to deliver by CS (OR 0.82, 95% CI 0.50 to 1.36), however fewer women in the intervention group who had vaginal births reported fear of pain in labour and had shorter labours.\textsuperscript{168} [evidence level 1b]

Responding to requests for CS

Obstetricians estimate that they agree to perform a CS for about half of the requests they receive.\textsuperscript{4} [evidence level 3] A woman’s request for CS is the ‘start of a continuing dialogue and process’ during which a negotiated plan of care can be developed which enables women to continue to feel in control with the support of her health care providers.\textsuperscript{169} [evidence level 4] When a woman requests a CS the first response should be to determine the reason for the request and the factors that are contributing to the request. This can then be followed by the provision of information comparing the risks and benefits of planned CS and vaginal birth (refer to Tables 4.5 & 4.6).

Review question

What is the appropriate care pathway for women who request a primary caesarean section where there is no obstetric or medical indication?

Overview of evidence

When addressing this question for the guideline, the GDG hoped to find evidence relating to the effectiveness of interventions for providing care for women requesting a CS. Unfortunately no such evidence was identified. One prospective cohort study (Wiklund et al., 2007) conducted in Sweden, investigated post partum outcomes in women having their first baby planning CS in the absence of medical indication compared to those planning a vaginal birth. Questionnaires were completed by the women on recruitment prior to giving birth, at two days and three months post partum. The outcomes recorded included their reason for the request, self-estimated health, expectations of birth and experience of delivery as well as duration of breastfeeding and time to re-establishment of sexual life. The study also had a secondary aim to study whether post partum depression was more common in the group planning CS.

Evidence profile

The table below is the summary of the evidence from this one study which included 91 women who planned to have a CS and 266 consecutively recruited controls who were planning a vaginal birth.

Maternal outcomes

| Table 5.5 GRADE summary of findings for comparison of planned CS vs. vaginal birth (maternal outcomes) |
|-------------------------------------------------|-------------------------------------------------|-------------------------------------------------|-------------------------------------------------|-------------------------------------------------|
| Number of studies | Results | Effect | Quality |
| Maternal request CS | Planned vaginal birth | Comparative t test/chi\(^2\) (p value) | Absolute | Maternal hospital stay (mean days) |
| 1 study (Wiklund et al.,) | 3.6 | 2.8 | 34.40 (0.001) | 0.8 | Very low |
### Table 5.6 GRADE summary of findings for comparison of planned CS vs. vaginal birth (neonatal outcomes)

<table>
<thead>
<tr>
<th>Number of studies</th>
<th>Number of neonates (%)</th>
<th>Effect</th>
<th>Quality</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Maternal request CS</td>
<td>Planned vaginal birth</td>
</tr>
<tr>
<td><strong>Neonatal intensive care unit (NICU) care</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 study (Wiklund et al., 2007)</td>
<td></td>
<td>5/99 (5%)</td>
<td>12/237 (5%)</td>
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</table>

* Total number in case and/or control group not provided. Percentage reported by authors presented here.
Evidence statements

Maternal outcomes

Maternal hospital stay
One study found that women who had a planned CS remained in hospital for a longer time than women having a planned vaginal birth. This finding was statistically significant. The evidence for this outcome was of very low quality.

Birth experience
One study found that women with a planned CS reported a higher satisfaction score regarding their birth experience 2 days after birth compared to women having a planned vaginal birth and this effect remained at 3 months. These findings were both statistically significant. The evidence for these outcomes was of very low quality.

Uncomplicated breastfeeding
One study did not find a statistically significant difference in breastfeeding rates at 2 days postpartum between women who had a planned vaginal birth compared with women who had a planned CS. However, the same study found that more women who had a planned vaginal birth were breastfeeding at three months postpartum compared with women who had a planned CS. This finding was statistically significant. The evidence for these outcomes was of very low quality.

Coitus
One study did not find a statistically significant difference in the numbers of women resuming coitus at 3 months following a planned CS compared with those who had a planned vaginal birth. The evidence for this outcome was of very low quality.

Family planning
One study found that more women who had a planned vaginal birth had plans for a second child at 3 months postpartum compared to women having a planned CS. This finding was statistically significant. The evidence for this outcome was of very low quality.

Depression
One study did not find a statistically significant difference in signs of postnatal depression comparing women who had given birth by planned CS compared with those who had had a planned vaginal birth. The evidence for this outcome was of very low quality.

Neonatal outcomes

Neonatal intensive care unit care
One study found did not find a statistically significant difference in the number of neonates who received neonatal intensive care following planned CS compared with planned vaginal birth. The evidence for this outcome was of very low quality.

Health economics
A model to compare the cost-effectiveness of maternal request CS versus planned vaginal birth in primiparous women without any medical or obstetric indication for CS was developed. Full details of this model are presented in chapter 13 but a summary is provided below.

Risks for the two modes of birth were taken from a clinical review undertaken for this guideline update comparing outcomes by planned mode of birth, rather than actual mode, of birth (see section 4.2). The analysis considered both the costs of birth and “downstream” costs associated with the outcomes reported in the clinical review and found that a planned vaginal birth was approximately £800 cheaper than a maternal request caesarean section. This implies that the NHS could save £5.6 million for every percentage point reduction in caesarean section if the characteristics of the population were similar to those of women included within the guideline model. A cost utility analysis found that planned vaginal birth dominated maternal request caesarean section.

However, there may be other outcomes, such as urinary incontinence, which were not reported in the studies that were included in the clinical review which make the findings reported above more uncertain. Sensitivity analysis suggested that this could, under certain assumptions, produce a different cost-effectiveness result.
Evidence to recommendations

Relative value placed on outcomes considered

The group agreed that the most important outcomes to consider were women’s birth experience along
with women’s satisfaction and experiences of care. They noted that these are difficult outcomes to
measure given the disparate reasons that women request a CS.

The group also felt that women’s mental health was an important outcome to consider. It was
acknowledged that not agreeing to a request for a CS could have a negative impact on a woman’s
mental health and potentially lead to a long term need for psychological support postnatally.

The group noted that the length of hospital stay will not always be an important consideration for
women (as they accept that a surgical procedure will be associated with inpatient stay). However,
they agreed that it was important to recognise the increased cost and resources required associated
with CS.

Trade-off between clinical benefits and harms

From the evidence reviewed for maternal request the GDG noted that CS is associated with a longer
hospital stay and a higher rate of women not breastfeeding at three months. However, they weighed
this against the finding that women who had a CS described a significantly better birthing experience,
both immediately postnatally and three months after birth.

The group noted that the findings for breastfeeding might have been influenced by the different
demographic profile of the 2 groups of women. Women in the planned CS group were significantly
older than those in the planned vaginal birth group, were more likely to have come from abroad, less
likely to have received parenthood education and less likely to report their perceived health as good
compared with women in the planned vaginal birth group.

The GDG noted that the findings for depression were poorly reported and they did not feel that they
were helpful. From their own experience, the group was aware that for some women, not to receive a
requested CS can lead to poorer mental health outcomes such as anxiety both during and after the
pregnancy, and difficulty bonding with their baby.

Trade-off between net health benefits and resource use

The group noted that there was likely to be an increased resource use with CS due to the increased
length of hospital stay.

An economic model developed for this guideline suggested that planned vaginal birth was cost-
effective compared to a maternal request CS. However, this finding was limited to outcomes that were
reported in the included studies for the clinical review undertaken for this guideline (see section 4.2).
A sensitivity analysis suggested that the inclusion of adverse outcomes not reported, such as urinary
incontinence, could make the cost-effectiveness conclusion less certain. On balance this model does
not provide strong evidence to refuse a woman’s request for CS on cost-effectiveness grounds.

The group agreed that there was likely to be an associated cost with providing psychological support
to those women who experience mental health problems as a result of not receiving a CS on request.
However, they noted that this was only likely to be the case for a small proportion of women. The
GDG’s experience of caring for women requesting a CS was that anxiety about giving birth vaginally
was often at the root of the request, for example as a result of a previous poor birth experience. The
GDG believed that when women are given the opportunity to discuss these anxieties in a supportive
environment, the anxieties can often be reduced to the point where the woman is able to choose a
planned vaginal birth. The group agreed this was the preferred approach. It was not felt to be
necessary for the person providing this psychological support to be a mental health expert unless
clinically indicated, but rather that it could be provided by a member of the maternity team e.g. a
midwife or obstetrician. It was felt that the extra resource required to provide this support would be
offset by resources saved where a request for planned CS was appropriately changed to a planned
vaginal birth where a woman’s anxieties or concerns have been able to be addressed antenatally.
However, in situations where women persist in their request for a CS, following provision of the
opportunity to discuss and explore their reasons for the request, the GDG believed that the potential
for psychological harm caused by denying this request was sufficient to warrant this unacceptable in
terms of the woman’s health and has the potential to be costly in terms of long-term need for
psychological support. It was concluded, therefore, that if a vaginal birth is not an acceptable option to
the woman after discussion and the offer of support, she should be supported in her choice of a
planned CS.

The GDG were aware of instances where women had been offered referral to a perinatal mental
health expert and that this expert had not been granted access to the planned place of birth. The
group recognised that having this access is important in order to provide appropriate support and
adequately address any anxieties regarding the birth setting. As a result, the group agreed that it was
appropriate to recommend that the healthcare professional providing this care be given access.

**Quality of the evidence**

The GDG were hoping to find evidence of the effectiveness of antenatal interventions aimed at
supporting women who request a CS in the absence of medical indications. Unfortunately no such
evidence was identified. The one included study compared outcomes for women requesting and
receiving a CS with those who had a planned vaginal birth. This information was only marginally
useful in helping the GDG to decide its recommendations.

The women in the 2 groups were significantly different in a number of characteristics at baseline i.e.
compared to the planned vaginal birth group, the women who had a planned CS were older, more
had come from abroad and more had had IVF, although fewer reported their pregnancy was planned,
or that they had received parenthood education or perceived their health as good. The groups were
only similar in terms of having a university education and in number of smokers.

Analysis was not performed to assess the effects of these differences on the results obtained from the
questionnaires. In addition, a sub-group/per protocol analysis was not performed to estimate
outcomes separately for women who planned a vaginal birth but subsequently had an unplanned CS
(n = 29, 11%) or an instrumental delivery (n = 36, 13%).

The study was conducted in a middle-high income urban area in Sweden where women were highly
educated. The GDG considered the results to be relevant to a UK population but noted that the study
was not representative of women from a low socio-economic background. They agreed that the
quality of the study means that it is of limited relevance.

The group noted that there was no evidence comparing women who requested a CS and received
one with those who wanted a CS and didn’t receive one. This would have been a useful comparison.

It was also noted that there appeared to be incomplete reporting of some of the findings, such as
postnatal depression, which undermined the validity of those findings.

**Other considerations**

The group was aware that some groups of women such as those women who don’t speak English as
a first language can find it more difficult to express their concerns. They recognised the importance of
ensuring that all women are encouraged to discuss their concerns with a healthcare professional at
an early stage in pregnancy. Discussions with women requesting a CS need to sensitively explore
reasons behind the request including previous birth trauma, women’s perceptions of the risks of both
vaginal birth and CS, women’s perceptions of vaginal birth including misconceptions and lack of
knowledge surrounding birth, as well as issues surrounding planning a date for giving birth and
convenience.

The GDG also believed it was important for an individual obstetrician to be able to exercise his/her
own beliefs about what is the best course of action in any given situation. Thus, if an obstetrician feels
a woman’s request for CS is not appropriate after the woman has received appropriate counselling
and support, then s/he should be able to decline to support the woman’s request. This does not over-
rule the woman’s rights to express a preference for a CS however, and in this instance the
obstetrician should transfer care of the woman to an NHS obstetrician within the same unit who is
happy to support her choice.
### Recommendation

34 When a woman requests a CS explore, discuss and record the specific reasons for the request. [new 2011]

35 If a woman requests a CS when there is no other indication, discuss the overall risks and benefits of CS compared with vaginal birth (see box A on page 64) and record that this discussion has taken place. Include a discussion with other members of the obstetric team (including the obstetrician, midwife and anaesthetist) if necessary to explore the reasons for the request, and to ensure the woman has accurate information. [new 2011]

36 When a woman requests a CS because she has anxiety about childbirth, offer referral to a healthcare professional with expertise in providing perinatal mental health support to help her address her anxiety in a supportive manner. [new 2011]

37 Ensure the healthcare professional providing perinatal mental health support has access to the planned place of birth during the antenatal period in order to provide care. [new 2011]

38 For all women requesting a CS, if after discussion and offer of support (including perinatal mental health support for women with anxiety about childbirth), a vaginal birth is still not an acceptable option, offer a planned CS. [new 2011]

39 An obstetrician has the right to decline a woman's request for a CS. If this happens, they should refer the woman to an NHS obstetrician in the same unit who will carry out the CS. [new 2011]

### Research Recommendation

RR 15 What support or psychological interventions would be appropriate for women who have a fear of vaginal childbirth and request a CS?

Interventions for evaluation could include:

- support from a named member of the maternity team
- continuity of carer
- formal counselling
- cognitive behavioural therapy.

Outcomes could include:

- mode of birth planned at term
- psychological outcomes (postnatal depression, post-traumatic stress disorder, self-esteem, mother-infant bonding)
- breastfeeding.

**Why this is important**

Fear of vaginal childbirth may stem from:

- fear of damage to the maternal pelvic floor
- damage to the baby during childbirth
- self-doubt on the ability to physically achieve vaginal birth
- previous childbirth experience
- unresolved issues related to the genital area.

Currently there is a wide variation in practice and limited resources lead to limited availability of effective interventions. Interventions that may be appropriate include:

- antenatal clinics dedicated to providing care for women with no obstetric indications who request a CS
• referral to a psychologist or a mental health professional
• referral to an obstetric anaesthetist
• intensive midwifery support.

Continuity of healthcare professional support from the antenatal to the intrapartum periods and ‘one to one’ midwifery care during labour are also often lacking and may make a difference to women who are anxious or afraid.

All of these interventions have different resource implications and there is no clear evidence to suggest that any are of benefit. The proposed research would compare in a randomised controlled trial two or more of these interventions in women requesting a CS. In the absence of any evidence, there is a case for comparing these interventions with routine antenatal care (that is, no special intervention).

This research is relevant because it would help to guide the optimal use of these limited resources and future guideline recommendations.

RR 16 Medium to long term quality of life study comparing psychological and physical outcomes in women who have had a requested and given birth by CS compared with women who plan a vaginal birth.

RR 17 Qualitative and quantitative research should be carried out to look at the reasons that lead to pregnant women’s request for CS

RR 18 The effect of counselling and other interventions such as second opinion and provision of support on the likelihood of CS for women who express a preference for CS need further evaluation.
6 Factors affecting likelihood of CS during intrapartum care

6.1 Place of birth

Planned home birth

One systematic review that includes one small RCT comparing planned home birth to planned hospital birth was identified (n = 11). The RCT included operative delivery but not specifically CS. No difference was reported for any of the outcomes measured however this was a small RCT and has limited power to detect a difference.\(^{171}\) [evidence level 1b]

A systematic review of observational studies evaluating the safety planned home births (in countries with good health resources) versus planned hospital births identified six cohort studies (n = 24,092)\(^{172}\) [evidence level 2b] Outcome measures included perinatal and maternal mortality, Apgar scores and incidence of maternal lacerations. The review also reported other outcomes including CS rates. No difference was detected in perinatal mortality in any of the individual studies, nor in the pooled data. In the home birth group, both low 5 minute Apgar and maternal lacerations were less frequent in all studies. The odds of CS were lower in the planned home birth group in five studies (reported crude OR of CS in studies: 0.04; 0.09; 0.31; 0.05; 0.27). No maternal deaths occurred but the studies are underpowered to evaluate this outcome.\(^{172}\) [evidence level 2a]

A subsequent cohort study in Canada (n = 2176) reported on CS rates and maternal and perinatal morbidity between 3 groups, women who had a planned home birth, women who were attended by a physician in hospital and women who were attended by a midwife in hospital. They reported that less women in the home birth group had a CS, compared to women in the physician-attended hospital group (adjusted OR 0.3, 0.22 to 0.43) and compared to the midwife attended hospital group (adjusted OR 0.66, 0.44 to 0.99). Odds ratios were adjusted for maternal age, lone parent status, income quintile, substance use and parity. No difference was detected between the groups for maternal or perinatal morbidity.\(^{172}\) [evidence level 2a]

A large prospective case controlled UK study of 5971 planned home births and 4724 planned hospital births reported that planning a home birth halved the chance of having a CS (unadjusted OR 0.49, 95% CI 0.39 to 0.62).\(^{174}\) [evidence level 2b]

<table>
<thead>
<tr>
<th>Number</th>
<th>Recommendation</th>
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</thead>
<tbody>
<tr>
<td>40</td>
<td>During their discussions about options for birth, healthy pregnant women with anticipated uncomplicated pregnancies should be informed that planning a home birth reduces the likelihood of CS. [B] [2004]</td>
</tr>
</tbody>
</table>

‘Midwifery-led unit’ or ‘birthing centre’

Current convention in the UK is that the term “midwifery-led units” refers to units that are near to or adjacent to a hospital maternity facility and that “birthing centres” are stand alone units. However this convention is not standardised in the literature. The centres are intended for “low risk” women. The care is midwife led with minimal medical intervention, sometimes described in the literature as...
‘home like’. Case series have reported reduced CS or operative delivery in ‘midwifery-led units’ or ‘birthing centres’.¹⁷⁵–¹⁸⁰ [evidence level 4]

A systematic review that included six RCTs (n = 8677) compared clinical outcomes between women delivering in a midwife led unit or in a hospital.¹⁸¹ [evidence level 1a] The RCTs were conducted in Stockholm¹⁸², Australia¹⁸², United Kingdom¹⁸⁴–¹⁸⁶ and Canada.¹⁸⁷ The centre in each of the RCTs was situated close to the conventional labour ward within the same hospital setting. The RCTs all describe the environment as ‘home like’ and that the care was aimed at women retaining control and choice with minimal medical intervention. Three of the studies do not describe the study environment any further.¹⁸²,¹⁸³,¹⁸⁵ Three of the studies describe the furnishings in detail (for example “furnished to appear like a normal household bedroom”)¹⁸⁴,¹⁸⁶,¹⁸⁷ and one RCT also mentions specifically interventions that were avoided such as enemas, perineal shaving, intravenous infusion and electronic fetal monitoring.¹⁸⁷ [evidence level 1b]

All RCTs (n = 8646) reported on CS rates, a further 39 outcomes are also reported. No difference was detected in CS rates between ‘midwifery-led unit’ and conventional birth settings (RR 0.85, 0.72 to 1.00). The review has a 90% power to detect a difference of at least 2% in CS rates if such a difference exists. No difference in instrumental vaginal deliveries was detected (OR 0.87, 0.74 to 1.01). Birth in a ‘midwife-led unit’/‘birth centre’ was associated with lower rates of intrapartum analgesia (OR 0.82, 0.72 to 0.93); less augmented labour (OR 0.72, 0.64 to 0.81); and fewer women ‘less than completely satisfied with care’ (OR 0.62, 0.55 to 0.70).¹⁸¹ [evidence level 1a]

A further UK RCT (n = 2578) comparing care ‘midwifery-led unit’ or in a conventional labour ward did not evaluate mode of delivery but assessed maternal satisfaction using a postal questionnaire. No difference was detected in rates of satisfaction between the groups. Women who had their babies in the ‘midwifery-led unit’/‘birthing centre saw fewer medical staff, were more likely to report having had a choice as to moving around during childbirth and alternative positions for birth and more likely to have made their own decisions regarding analgesia.¹⁸⁸ [evidence level 1b]

We did not identify any RCTs that compared birthing centres which are stand alone to conventional maternity facilities. However we did identify a case series following women admitted for labour and delivery at 84 ‘free standing’ birthing centres in the United States (n = 11,814). The overall rate of CS was 4.4%. The rate of transfer to other maternity facilities before birth was 11.9%. Other morbidity outcomes reported include 5-minute Apgar of less than 7 occurred in less than 0.5% of births.¹⁷⁸ [evidence level 4]

An Australian postnatal survey of women’s views about their birth experience (n = 395) reports that women who had given birth at home or at a ‘midwifery-led unit’ were more likely to feel that the birth place affected the bonding process and less likely to see birth as a medical condition compared to women who gave birth in a conventional labour ward. Women who gave birth at home were older, more educated, more likely to be multiparous and better informed about childbirth compared to the women who gave birth in the ‘midwife-led unit’ or in the conventional labour ward. Adjusting for these differences, place of birth correlated with women’s satisfaction with health care providers.¹⁸⁹ [evidence level 3]

<table>
<thead>
<tr>
<th>Number</th>
<th>Recommendation</th>
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<tbody>
<tr>
<td>41</td>
<td>During their discussions about options for birth, healthy pregnant women with anticipated uncomplicated pregnancies should be informed that planned childbirth in a ‘midwifery-led unit’ does not reduce the likelihood of CS. [A] [2004]</td>
</tr>
</tbody>
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<tr>
<th>Number</th>
<th>Research Recommendation</th>
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<tbody>
<tr>
<td>RR 19</td>
<td>RCTs comparing planned birth in a stand-alone birthing centre to birth in conventional maternity facilities or midwifery led units.</td>
</tr>
<tr>
<td>RR 20</td>
<td>Qualitative research is needed to explore women’s opinions on place of birth and the impact of place of birth on their birth experiences.</td>
</tr>
</tbody>
</table>
**Delayed admission to labour ward**

A systematic review included one RCT (n = 209) compared a labour assessment program in a separate unit within the hospital and delayed admission to labour ward until labour is in the active phase, with direct admission to the labour ward.\(^{190,191}\) The RCT did not detect a difference in CS rates between the two groups (OR 0.70, 95% CI 0.27 to 1.79). At least two thousand women would be needed in each group to detect a 3% difference in CS therefore this RCT is underpowered to detect this difference in CS rates. There were differences in other outcomes such as length of time spent in the labour ward, analgesia requirements, oxytocic use and maternal satisfaction, measured using sense of control (see evidence table). [evidence level 1b]

An observational study (n = 3220) reported a reduced likelihood of CS with increased cervical dilatation at the time of presentation in labour. The CS rates for nulliparous women presenting at 0–3 cm was 10% compared with 4% for those presenting at 4–10 cm (p = 0.001). This was consistent for nulliparous and parous women.\(^{192}\) [evidence level 2b]

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<th>Research Recommendation</th>
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<tr>
<td>RR 21</td>
<td>Further RCTs are needed to determine the effect of ‘delayed admission in labour’ on the likelihood of CS.</td>
</tr>
</tbody>
</table>

### 6.2 Factors reducing the likelihood of CS

#### One-to-one support

One-to-one support in labour had been evaluated in recently published systematic review;\(^{193}\) this current review replaces the previous review on this subject by the same authors.\(^{194}\) [evidence level 1a]. The first review included 14 RCTs (n = 5000), the new review includes 15 RCTs (n = 12,791) the newly included study is a multi centre RCT (n = 6915 women) conducted in Canada and the US (13 centres). The trial evaluated the effectiveness of continuous labour support by a specially trained nurse/midwives to usual care. Each hospital in the RCT had a CS rate of at least 15%. The main outcome measure was CS rate. The study did not detect a difference in CS rate between the two groups. The use of continuous electronic fetal monitoring higher in the usual care group (79%) compared to those in the continuous support group (75%, p < 0.001). All comparisons of women’s likes and dislikes, and their future preference for amount of nursing support, favoured the continuous labour support group.\(^{195}\) [evidence level 1a]

The new systematic review (15 RCTs, n = 12,791 women) evaluates the effects of one-to-one support on women and their babies. In addition the new review also considers whether the effects of continuous support are influenced by routine practices and policies in the birth environment that may affect a woman’s autonomy, freedom of movement and ability to cope with labour; whether the caregiver is a member of staff and whether the continuous support begins early or late in labour.\(^{195}\) [evidence level 1a] The RCTs in the review included support persons that varied in terms of their experience, qualifications and relationship to the women in childbirth. In eight RCTs the support was provided by a member of hospital staff. The remaining 7 RCTs included women from the community (“doulas”), with or without prior training, a childbirth educator, or a close female relative. Half of the RCTs were conducted in developed countries, where hospital policy permitted women to be accompanied by their husband/partners or other family members during labour. The remaining RCTs were conducted in developing countries in settings in which only the support person allocated by the study was allowed to accompany the woman during labour. No RCT evaluated the effects of husbands or partners as providers of support.

The results of the review reported that women who had continuous one-to-one support during labour were less likely to have a CS (15 trials, n = 12,791, RR 0.90, 95% CI 0.82 to 0.99). The effects of continuous support on CS appeared to be stronger in settings which did not permit the presence of
additional support people (chi squared = 4.46, p < 0.05) and when epidural was not routinely available (chi squared 4.97, p < 0.05). The routine use of EFM did not affect the impact of one-to-one support on CS rates. The reduction in CS was influenced by who was giving the support and the reduction was only seen in the RCTs where the support was not provided by members of staff (RR 0.74, 95% CI 0.61 to 0.9). The difference between different sub-groups of non medical providers of support was not statistically significant. The impact of timing of onset of continuous support was of borderline statistical significance (chi squared = 5.93, p = 0.05) favouring support that began before active labour. Thirty other outcomes were considered in the review, but are not reported here.\textsuperscript{193} [evidence level 1a]

<table>
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<th>Number</th>
<th>Recommendation</th>
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<tbody>
<tr>
<td>42</td>
<td>Women should be informed that continuous support during labour from women with or without prior training reduces the likelihood of CS. [A] [2004]</td>
</tr>
</tbody>
</table>

Pregnancy after 41 weeks

A systematic review of 26 RCTs compared induction of labour with expectant management after 41 weeks. Offering routine induction after 41 weeks reduced perinatal death (19 RCTs, n = 7925. Peto OR 0.20, 95% CI 0.06 to 0.70) and the rate of CS (9 RCTs, n = 5954 Peto OR 0.87, 95% CI 0.77 to 0.99).\textsuperscript{196} [evidence level 1a]

It is estimated that by 41 weeks 74% of women have given birth, this increases to 82% by 42 weeks. The risk of stillbirth increases from 1 per 3000 ongoing pregnancies at 37 weeks to 3 per 3000 ongoing pregnancies at 42 weeks to 6 per 30000 with ongoing pregnancies at 43 weeks. A similar increase in neonatal mortality is also reported.\textsuperscript{197} [evidence level 2a]

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<th>Number</th>
<th>Recommendation</th>
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<tbody>
<tr>
<td>43</td>
<td>Women with an uncomplicated pregnancy should be offered induction of labour beyond 41 weeks because this reduces the risk of perinatal mortality and the likelihood of CS. [A] [2004]</td>
</tr>
</tbody>
</table>

Partogram

Progress in labour can be assessed using the clinical parameters of descent of the presenting part and dilatation of the cervix. No study has evaluated tests based on maternal and fetal outcomes. The partogram is derived from a curve describing normal labour (Friedman’s curve). The original Freidman’s curve was developed using observational data from 100 American primigivad women at term in spontaneous labour (included 98 singleton cephalic, 1 breech presentation and 1 multiple pregnancy). Twenty two percent of the women received caudal anaesthesia and 10 percent received oxytocin augmentation. Cervical dilatation was determined using rectal examination predominantly at 10, 30 or 60 minute intervals. Curves of dilatation versus time were produced and resulted in a sigmoid curve of progress of labour with average progress during the active phase of 1.1cm per hour and average length of labour of 12 hours for nulliparous women and 6 hours for multiparous women.\textsuperscript{196} [evidence level 3] More recent observational studies from the USA (n = 2511) measured the length of labour in women who had not received oxytocin or epidurals and report average length of labour for nulliparous women to be 19.4 hours and 13.7 hours for multiparous women. This is longer than the originally described normal labour curve.\textsuperscript{199} [evidence level 3]

On a partogram cervical dilatation and descent of the presenting part are plotted graphically against time. The partogram was initially proposed as a screening tool for use in poorly resourced countries to identify women who needed referral to hospital. The partogram includes two lines, an alert line and an action line. The alert line is set at a rate of 1cm per hour (derived from Friedman’s curve). The action line is drawn 4 hours to the right of the alert line. If the progress of labour crossed the action line women were referred to hospital for either augmentation of labour or CS.\textsuperscript{200,201} [evidence level 3]
Three RCTs have evaluated the use of partograms in the management of labour. The first RCT compared using a partogram with a four hour action line to not using a partogram in the management of labour. This was a cluster randomised trial where the unit of randomisation was a maternity hospital. Four pairs of hospitals participated. Each hospital had a practice of active management of labour including oxytocin use. The effect of the partogram was analysed in a before and after design which compared labour outcome data on 10,049 women who delivered before implementation of the partogram (4 hour action line) with data on 9130 women who delivered after implementation. This RCT did not report CS rates but did report rates of spontaneous vaginal birth. The number of spontaneous cephalic births was increased after implementation of the partogram (83% vs. 86.3%, \( p < 0.001 \)). There was a decrease in the proportion of women with labours of more than 18 hours (551 versus 249, \( p < 0.001 \)), labours augmented by oxytocin (\( p = 0.041 \)) and the number of intrapartum stillbirths (0.5% vs. 0.31%, \( p = 0.024 \)). There was no change in the overall duration of labour or other neonatal indices. Similar patterns were noted for multiparous and primiparous women.\(^{202} \) [evidence level 1b]

The second RCT (n = 928 women) compared partograms with different action lines (either 2, 3 or 4 hours to the right of the alert line set at 1 cm per hour). The primary outcomes were CS rate and maternal satisfaction. CS rate was lowest when labour was managed using a partogram with a 4 hour action line. Women in the 2 hour arm were most satisfied with their labour experience. No difference was found in the secondary outcomes of neonatal and maternal morbidity.\(^{203} \) [evidence level 1b]

The third RCT conducted in South Africa (n = 694) compared management using a single alert line partogram offering oxytocin if the alert line was crossed (with 2 hour vaginal examinations) to management using a 4 hour action line. CS was a primary outcome. Women in the intervention group were less likely to have a CS (RR 0.68, 95% CL 0.50 to 0.93).\(^{204} \) [evidence level 1b]

Meta-analysis of the 2 RCTs that included comparison of the two hour action line with a four hour action line partogram showed no difference in CS rate between the use of 2 or 4 hour action lines (RR 0.93, 95% CL 0.48 to 1.78).\(^{203,204} \) [evidence level 1b] The use of a 4 hour partogram reduces the number of vaginal examinations that women would undergo during labour.

<table>
<thead>
<tr>
<th>Number</th>
<th>Recommendation</th>
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<tbody>
<tr>
<td>44</td>
<td>A partogram with a 4-hour action line should be used to monitor progress of labour of women in spontaneous labour with an uncomplicated singleton pregnancy at term, because it reduces the likelihood of CS. [A] [2004]</td>
</tr>
</tbody>
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<th>Number</th>
<th>Research Recommendation</th>
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</thead>
<tbody>
<tr>
<td>RR 22</td>
<td>RCT evidence is needed to determine the impact of partograms based on different curves of labour on CS rates and morbidity outcomes.</td>
</tr>
</tbody>
</table>

**Decision making for unplanned CS**

Second opinion has been proposed as an intervention to decrease CS rates. Second opinion refers to a doctor needing the agreement of another usually more senior second opinion before a decision for CS can be made. A large multi centred RCT in five South American countries has recently been completed however the results have not been reported.

Using the NSCSA data the proportion of CS cases with consultant involvement varied between maternity units, although in the majority of CS, the consultant was the most senior obstetrician involved in the decision (see table).

In maternity units where consultant obstetricians were frequently involved either in the decision for CS or present in theatre for “emergency” CS the crude and adjusted CS rates (having taken into account case mix differences) were lower (see Tables 6.1 and 6.2).
Table 6.1 Proportion of CS with consultant involvement in maternity units

<table>
<thead>
<tr>
<th>Consultant present in theatre</th>
<th>Median (%)</th>
<th>IQR (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>All CS</td>
<td>12.6</td>
<td>7.6 – 18.5</td>
</tr>
<tr>
<td>“Emergency” CS</td>
<td>8.7</td>
<td>5.8 – 13.3</td>
</tr>
<tr>
<td>“Emergency” CS out of hours (1800 – 0700)</td>
<td>4.8</td>
<td>2.1 – 8.8</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Consultant involved on decision making to perform CS</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>All CS</td>
<td>76.4</td>
</tr>
<tr>
<td>“Emergency” CS</td>
<td>75.0</td>
</tr>
<tr>
<td>“Emergency” CS out of hours (1800 – 0700)</td>
<td>72.4</td>
</tr>
</tbody>
</table>

Table 6.2 Relationship between proportion of CS where there was consultant involvement and CS rates

<table>
<thead>
<tr>
<th>Consultant present in theatre</th>
<th>Crude CS rate</th>
<th>Adjusted CS rate</th>
<th>Spearman's rank correlation coefficient</th>
<th>P value</th>
<th>Spearman's rank correlation coefficient</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>All CS</td>
<td>-0.01</td>
<td>0.85</td>
<td>-0.05</td>
<td>0.48</td>
<td></td>
<td></td>
</tr>
<tr>
<td>“Emergency” CS</td>
<td>-0.12</td>
<td>0.06</td>
<td>-0.14</td>
<td>0.04</td>
<td></td>
<td></td>
</tr>
<tr>
<td>“Emergency” CS out of hours (1800 – 0700)</td>
<td>-0.12</td>
<td>0.07</td>
<td>-0.14</td>
<td>0.04</td>
<td></td>
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</tbody>
</table>

Consultant involved in decision making to perform CS

<table>
<thead>
<tr>
<th>Consultant involved in decision making to perform CS</th>
<th>Crude CS rate</th>
<th>Adjusted CS rate</th>
<th>Spearman's rank correlation coefficient</th>
<th>P value</th>
<th>Spearman's rank correlation coefficient</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>All CS</td>
<td>-0.19</td>
<td>&lt; 0.01</td>
<td>-0.19</td>
<td>&lt; 0.01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>“Emergency” CS</td>
<td>-0.18</td>
<td>&lt; 0.01</td>
<td>-0.17</td>
<td>0.01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>“Emergency” CS out of hours (1800 – 0700)</td>
<td>-0.19</td>
<td>&lt; 0.01</td>
<td>-0.17</td>
<td>0.01</td>
<td></td>
<td></td>
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</table>

Number Recommendation

45 Consultant obstetricians should be involved in the decision making for CS, because this reduces the likelihood of CS. [C] [2004]

Electronic fetal monitoring and fetal blood sampling

Systematic reviews of 9 RCTs (conducted between 1976–1993, n = 18,561 women) have compared the use of electronic fetal monitoring (EFM) during labour to intermittent auscultation. No difference is detected in perinatal mortality (RR 0.89, 95% CI 0.60 to 1.33). The use of EFM during intrapartum care results in increased CS rates (RR 1.4, 95% CI 1.23 to 1.61) This increase is less marked if fetal blood sampling (FBS) is used (RR 1.27, 95% CI 1.08 to 1.51 for EFM with FBS, compared with RR 1.41, 95% CI 1.23 to 1.61 for EFM without FBS). It is therefore recommended that where delivery is contemplated because of an abnormal fetal heart rate pattern, in cases of suspected fetal acidosis, FBS should be undertaken in the absence of technical difficulties or any contraindications. Contraindications to FBS include maternal infection (such as HIV, hepatitis viruses or herpes simplex
virus); fetal bleeding disorders such as haemophilia and prematurity (less than 34 weeks). Where there is clear evidence of acute fetal compromise, e.g. prolonged decelerations (longer than 3 minutes), FBS should not be undertaken and the baby should be delivered urgently.\textsuperscript{2}

The NSCSA measured practice against this audit standard for CS.\textsuperscript{4} Overall an abnormal CTG was noted in 69% of singleton cephalic pregnancies delivered by CS for presumed fetal compromise. If the CTG was noted to be severely abnormal or cervical dilatation was less than 4cm these cases were not included (50%). Overall practice concorded with the audit standard in 44% of cases. However, there was marked variation in practice. Five percent of maternity units met the standard in all cases (100%), in 9% the standard was not reached for any case. Units and regions which used FBS more frequently before CS had lower CS rates. Overall, cases in which this recommendation was not met contributed 4.6% to the overall CS rate or about 1% of all births.\textsuperscript{4}

<table>
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<th>Number</th>
<th>Recommendation</th>
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<tbody>
<tr>
<td>46</td>
<td>Electronic fetal monitoring is associated with an increased likelihood of CS. When CS is contemplated because of an abnormal fetal heart rate pattern, in cases of suspected fetal acidosis, fetal blood sampling should be offered if it is technically possible and there are no contraindications. [B] [2004]</td>
</tr>
</tbody>
</table>

### 6.3 No influence on likelihood of CS

The following interventions during intrapartum care have not been shown to influence the likelihood of CS. These interventions may have other effects (beneficial or harmful) which are outside the scope of this guideline and are not considered here.

### Walking in labour

Two RCTs have evaluated the effect of walking in labour to usual care, one conducted in the UK (n = 68)\textsuperscript{207} [evidence level 1b] and the other conducted in the USA (n = 1067)\textsuperscript{208} [evidence level 1b]. No difference was detected in the CS rates between women who walked around during labour and those who did not (RR 0.71, 95% CI 0.43 to 1.20). Most of the weight of the pooled RR in the meta-analysis comes from the larger RCT. Therefore it is not surprising that the US RCT did not detect a difference in CS rates between groups (RR 0.73, 95% CI 0.43 to 1.24). The study has 80% power to detect a difference of at least 4% in CS rate, therefore if walking in labour has an impact on CS rates it is likely to be less than 4%. The RCT did not detect a difference in other outcomes including length of the first stage of labour and need for analgesia. The results were similar for parous and multiparous women.\textsuperscript{208} [evidence level 1b]

### Position in the second stage of labour

A systematic review\textsuperscript{209} of 18 RCTs evaluated the effect of different positions for the second stage of labour. No difference was detected between any upright or lateral position during second stage on CS rates compared to supine or lithotomy positions (12 RCTs; n = 2250; RR 0.87, 95% CI 0.52 to 1.45). Use of any upright or lateral position, compared with supine or lithotomy positions, was associated with the following: reduced duration of second stage of labour (12 RCTs. Weighted mean difference: 5.4 minutes, 95% CI 3.9, 6.9 minutes); a reduction in assisted deliveries (17 RCTs. OR 0.82, 95% CI 0.69 to 0.98); a reduction in episiotomies (11 RCTs: OR 0.73, 95% CI 0.64 to 0.84); an increase in second degree perineal tears (10 RCTs: OR 1.30, 95% CI 1.09 to 1.54); increased estimated risk of blood loss greater than 500 ml (10 RCTs: OR 1.76, 95% CI 1.34 to 3.32); reduced reporting of severe pain during second stage of labour (1 RCT: OR 0.59, 95% CI 0.41 to 0.83) and fewer abnormal fetal heart rate patterns (1 RCT: OR 0.31, 95% CI 0.11 to 0.91).\textsuperscript{209} [evidence level 1a]

### Immersion in water in labour

Water births and the use of immersion in water during labour comprise 0.6% of births in the UK.\textsuperscript{210} [evidence level 3] A systematic review\textsuperscript{211} [evidence level 1a] that included three RCTs (n = 988)
compared immersion in water during labour (no births occurred in the water) to conventional care. Another RCT (n = 1237) on this topic has been published since this review.\textsuperscript{212} [evidence level 1b] The CS rate in the intervention arm of these RCTs ranged from 1.8% to 8.9%, in the control group it ranged from 0% to 7.9%. A new meta-analysis of the findings from these 4 RCTs (n = 2225) did not detect a difference in CS rates between the two groups (RR 1.31, 95% CI 0.89 to 1.93) [evidence level 1a]. Overall these studies have a 90% power to detect a difference of at least 4% in CS rates between the two groups therefore if water birth has an effect on CS rate it is likely to be less than 4%.

One of the above RCTs interviewed a subset of women about their use and satisfaction with care in labour. Women most liked the presence of a support person and immersion in water.\textsuperscript{213}

A national cohort study using regional UK survey data compared the perinatal mortality and morbidity of 4032 births either in water (or following labours in water) to births not in water. They report no difference in perinatal mortality (RR 0.9, 99% CI 1.2 to 3.6). There were two cases of water aspiration which required admission to NICU.\textsuperscript{214} [evidence level 3] A prospective observational study in Switzerland of 7508 births of which 2014 were water births showed no increased risk for women or their babies. The study reported: lower episiotomy rates, higher rates of intact perineum, lower blood loss and lower use of pain killers in women who had a waterbirth.\textsuperscript{215} [evidence level 3]

A number of position papers have provided guidelines for water births in the absence of adequate evidence, and have suggested the continued reporting of adverse events.\textsuperscript{216,217} [evidence level 4]

### Analgesia during labour

There has been an increase in the use of epidural analgesia in labour and there has been concern that this may have contributed to an increase in CS. Observational data provides conflicting results.\textsuperscript{218–227} [evidence level 3]

Two systematic reviews have included RCTs of women in spontaneous labour who requested analgesia and were randomised to receive either epidural analgesia or usual analgesia (such as intravenous or intramuscular pethidine). The first review of 10 RCTs (n = 2369) did not detect a difference in CS rates between the two groups (OR 1.5, 95% CI 0.81 to 2.76).\textsuperscript{228} [evidence level 1a]. A subsequent review includes 11 RCTs (n = 3157, it includes 6 RCTs from the previous review, 2 new RCTs\textsuperscript{229,230} and 2 RCTs not included in the first review\textsuperscript{231–234}). It also did not detect a difference in CS rates (OR 1.30, 95% CI 0.93 to 1.83).\textsuperscript{235} [evidence level 1a]

We did not identify any RCTs that had compared parenteral analgesia (intravenous or intramuscular opiate derived analgesia) to placebo or complementary therapies on mode of birth and risk of CS.

### Raspberry leaf during labour

An RCT (n = 192) was conducted that looked at the use of raspberry leaf, given in tablet form during labour. No difference was detected in length of labour or mode of birth, including “emergency” CS.\textsuperscript{236} [evidence level 1b]. Earlier descriptive studies of raspberry leaf used in labour excluded women who had a CS from their analysis.\textsuperscript{237} [evidence level 3]

<table>
<thead>
<tr>
<th>Number</th>
<th>Recommendation</th>
</tr>
</thead>
<tbody>
<tr>
<td>47</td>
<td>Women should be informed that the following interventions during intrapartum care have not been shown to influence the likelihood of CS, although they may affect other outcomes that are outside the scope of this guideline:</td>
</tr>
<tr>
<td></td>
<td>- walking in labour</td>
</tr>
<tr>
<td></td>
<td>- non-supine position during the second stage of labour</td>
</tr>
<tr>
<td></td>
<td>- immersion in water during labour</td>
</tr>
<tr>
<td></td>
<td>- epidural analgesia during labour</td>
</tr>
<tr>
<td></td>
<td>- the use of raspberry leaves. [A] [2004]</td>
</tr>
</tbody>
</table>

Caesarean section: full guideline DRAFT (September 2011)
**Number** | **Research Recommendation**
---|---
RR 23 | RCT evidence is required to evaluate the effect of parenteral analgesia (intramuscular and intravenous morphine based analgesia) used during childbirth on the likelihood of CS.

### Complementary therapies during labour

Complementary therapies used during pregnancy include acupuncture, aromatherapy, hypnosis, Chinese medicines, herbal products and nutritional supplements, homeopathic medicines and raspberry leaf (discussed previously). We have only considered their use during labour in this guideline. The antenatal use of complementary therapies is included in the NICE Antenatal Care Guideline.¹

We identified a systematic review of complementary therapies for pain management in labour which includes seven RCTs (n = 366) using different modalities of pain management²³⁸ [evidence level 1a]. CS rates were considered as secondary outcomes in two of the included studies: one RCT using acupuncture (n = 90), one aromatherapy RCT (n = 22), neither showed any difference in CS rates however the trials were underpowered to evaluate this outcome. Two RCTs (n = 125) have compared the use of hypnosis to usual analgesia. CS was not reported. However women in the hypnosis group were more likely to have a spontaneous vaginal birth (RR 1.38, 95% CI 1.13 to 2.47).²³⁸ [evidence level 1b]

A large survey (n = 8058) of women views on the effect of using aromatherapy during labour. Effect was measured using a Likert scale. About half of the women reported aromatherapy was helpful, a minority (14%) found it unhelpful.²³⁹ [evidence level 4]

The suggested benefits of Chinese medicines in labour include prevention of nausea and vomiting, heartburn and fatigue. We did not identify any RCTs on their use in labour. We identified a cohort study on the use of Chinese medicines during pregnancy which reported no effect on mode of birth.²⁴⁰ [evidence level 2b]

Surveys from the USA and Australia suggest that there is widespread use of herbal products and nutritional supplements during pregnancy, 12% of women in Australia²⁴¹ [evidence level 4] and 7% in the USA.²⁴² [evidence level 3] A UK survey of midwives estimated that 34% of midwives offer some form of complementary medicine to women during pregnancy or childbirth.²⁴³ [evidence level 4] The majority of this use is antenatal with only certain herbal products used during labour or to induce labour. We did not identify any RCTs on the use of herbs during labour but a number of expert opinion papers offer advice and suggested guidelines for their use. Using information from midwives surveys they recommend caution with the use of blue cohosh (due to reports of dizziness, fainting, nausea and meconium stained liquor as well as case reports of neonatal heart failure); black cohosh and castor oil to induce labour.²⁴⁴ [evidence level 4] There have not been reported complications with either evening primrose oil or raspberry leaf.²⁴⁵,²⁴⁶ [evidence level 4]

### Number Recommendation

| 48 | Women should be informed that the effects on the likelihood of CS of complementary therapies used during labour (such as acupuncture, aromatherapy, hypnosis, herbal products, nutritional supplements, homeopathic medicines, and Chinese medicines) have not been properly evaluated and further research is needed before such interventions can be recommended. [D] [2004] |

### Number | Research Recommendation
---|---
RR 24 | RCTs are needed to establish the safety and efficacy of complementary therapies
used during labour.

6.4 ‘Failure to progress’ in labour and CS

In the NSCSA, “failure to progress” in labour (FTP) was the primary indication for CS in 35% (n = 4,896) of women with term cephalic pregnancies and no uterine scar. For 17% (n = 811) of these women cervical dilatation at the time of CS was less than 4 cm. While 74% of these women had their labour augmented (65% were given oxytocin and amniotomy, 7% amniotomy only, 2% oxytocin only), 24% (n = 193) had no augmentation of labour before CS. The majority (98%) of women with cervical dilatation of at least 4 cm at the time of CS had either amniotomy or oxytocin or both. Twenty-five percent (n = 1231) of CS for FTP were done at a cervical dilatation of 10 cm, 28% (n = 345) of these women did not have oxytocin before CS. These cases in which labour augmentation with oxytocin was not used contributed 3.2% to the overall CS rate.5 [evidence level 3]

We searched for research that evaluated the impact of packages of interventions such as active management of labour and interventions such as routine amniotomy or oxytocin infusion used together or alone are included.

Active management of labour

Active management of labour refers to a labour ward protocol that includes routine amniotomy and early augmentation with oxytocin as well as strict criteria for the diagnosis of labour, abnormal progress in labour and fetal compromise. It also includes the continual presence of a midwife or support person during labour and peer review of assisted deliveries. Observational studies by the initiators of active management reported lower CS rates, reduction in the number of women having prolonged labour, better neonatal outcomes and improved maternal satisfaction.247 Subsequent observational studies did not replicate these findings.248,249 It has remained an area of controversy.250 [evidence level 3]

A systematic review of 10 RCTs (n = 5111) evaluated the effects of a package intervention of early augmentation of labour with amniotomy and oxytocin in nulliparous women compared to usual care (“care at the discretion of the individual doctor/midwife attending the woman in labour”). Overall there was no reduction in the likelihood of CS with early amniotomy and early oxytocin infusion (OR 0.9, 95% CI 0.7 to 1.1). Subgroup analysis of the therapy RCTs (recruited women in whom a delay in progress was diagnosed) (3 RCTs, n = 109) and prevention RCTs (7 RCTs, n = 5002) were undertaken. No difference in CS rate was apparent in these subgroups. However the therapy subgroup is too small and is therefore underpowered to evaluate this outcome.251 [evidence level 1a]

None of the RCTs had maternal satisfaction as an outcome measure.251 [evidence level 1a]

A recently published RCT from South Africa (n = 694) compared using a single line partogram, two-hourly vaginal examinations and use of oxytocin if the partogram line was crossed in nulliparous women to usual management (4 hour vaginal examinations). CS rates in the intervention group were lower (RR 0.68 95% CI 0.50 to 0.93) Analysis is by intention to treat but it was noted that there was a high proportion of protocol violations in both groups (about 30%)204 [evidence level 1b]. It was not possible to include this RCT with the earlier RCTs as the descriptions of management of labour were not consistent.

Oxytocin

Most RCTs identified incorporate the use of oxytocin into active management of labour. However we identified one RCT (n = 60) that looked at the effect of oxytocin without other components of active management of labour in women in whom there was a delay in labour progress. Women whose cervical dilatation was less than 0.5 cm per hour were randomised to one of three groups: group one – oxytocin was deferred for 8 hours; group two – low-dose oxytocin infusion (2mu/minute) or group three – high-dose oxytocin (7mu/minute). The CS rates between the three groups were not statistically different (45%, 35% and 26% respectively $\chi^2 = 1.6346$ 2df). There were no differences between the groups in terms of neonatal outcomes.252 [evidence level 1b] This RCT is underpowered to assess these outcomes.
Observational data from the original active birth management study suggested benefit of the early use of high dose oxytocin infusions. Subsequent observational studies that looked at the use of oxytocin alone in labour suggested that it decreased the CS rates and did not result in increased neonatal morbidity.

**Amniotomy**

A systematic review of nine RCTs looked at the impact of early routine amniotomy. CS rate was reported in 8 of the included RCTs (n = 4008). No difference in CS rates was found between early routine amniotomy and no routine amniotomy (OR 1.26, 95% CI 0.96 to 1.66). Amniotomy was associated with a reduction in labour duration of between 60 and 120 minutes, reduction in the likelihood of 5 minute Apgar of less than 7 (OR 0.54, 95% CI 0.30 to 0.96) and a decrease in the use of oxytocin (OR 0.79, 95% CI 0.67 to 0.92). Groups were similar with respect to other neonatal indicators.

**Operative delivery in the second stage**

Four percent (n = 1203) of all CS were performed for failure to progress in the second stage of labour (in women without a previous CS who had a term singleton cephalic infant). In the majority 55% (n = 661) no other method of delivery had been attempted before CS. In 35% (n = 427) of these occurrences, CS followed a failed attempt at ventouse, in 7% (n = 81) both ventouse and forceps had been attempted prior to CS and in 2% (n = 27) CS followed a failed attempt at forceps delivery. Overall in the UK while CS rates have increased, operative vaginal delivery rates have remained relatively constant (about 10–11%). However there has been a marked reduction in the use of forceps and an increase in the use of ventouse since the early nineties. A cohort study has compared the maternal and neonatal outcomes following either instrumental vaginal delivery or CS in the second stage of labour (n = 393 women, 184 had a vaginal delivery, 209 CS). Major haemorrhage (blood loss > 1000 ml) was more common after CS than vaginal delivery (adjusted OR 2.92, 95% CI 1.1 to 7.62). Length of hospital stay was increased with CS. No difference was detected in wound infection, blood transfusion, need for opiate analgesia or rates of breastfeeding. Odds ratios were adjusted for maternal body mass index, pre-eclampsia, maternal diabetes, duration of second stage, station and position of the presenting part, demographic differences and experience of the operator. A further study following up the same women after 3 years reported half had achieved a further pregnancy after 3 years. There was no difference the proportion of women who had difficulty conceiving but there was an increase in involuntary infertility of more than 1 year. Of women who choose not to have more children there was no difference in the proportion that stated they “could not go through childbirth again”. Of women who had a further pregnancy those who had had a previous instrumental vaginal birth were more likely to aim for and achieve a vaginal birth again (adjusted OR 15.55, 95% CI 5.25 to 46.04; adjusted OR 9.50, 95% CI 3.48 to 25.97). Qualitative research of women views on the impact of operative delivery in the second stage of labour (n = 27) described that women felt unprepared for operative delivery and that antenatal education had not adequately prepared them for this event.

<table>
<thead>
<tr>
<th>Number</th>
<th>Recommendation</th>
</tr>
</thead>
<tbody>
<tr>
<td>49</td>
<td>The following aspects of intrapartum care have not been shown to influence the likelihood of CS for ‘failure to progress’ and should not be offered for this reason, although they may affect other outcomes which are outside the scope of this guideline:</td>
</tr>
<tr>
<td></td>
<td>• active management of labour</td>
</tr>
<tr>
<td></td>
<td>• early amniotomy. [A] [2004]</td>
</tr>
</tbody>
</table>
**Number** | **Research Recommendation**
--- | ---
RR 25 | More RCTs are required to determine the effect of oxytocin augmentation as single interventions or as part of a package of interventions (such as “active management of labour”) on the likelihood of CS and other outcomes including women’s satisfaction with care.
RR 26 | Further research on the short and longer term health impacts of CS during the second stage compared to operative vaginal delivery are needed.

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**Female genital mutilation**

Female genital mutilation is defined by the World Health Organization (WHO) as, ‘all procedures that involve partial or total removal of the female external genitalia or other injury to the female genital organs whether for cultural, religious or other non-therapeutic reasons’. An estimated 10,000 to 20,000 girls in the UK are thought to have undergone genital mutilation. [evidence level 3]

The association between female genital mutilation and intrapartum complications has been systematically reviewed by the WHO. [evidence level 3] Possible complications include obstructed labour, fetal distress and increased perinatal mortality however the evidence for these are contradictory. [evidence level 3] No RCTs or observational studies have addressed the effect on health outcomes of CS in the management of female genital mutilation. It is outside the scope of this guideline to address the antenatal or intrapartum management of female genital mutilation.

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**6.5 Eating during labour**

The practice of encouraging women to eat and drink during labour in order to maintain their strength for the second stage changed following publication of a case series (n = 66) of aspiration pneumonitis. In this paper Mendelson suggested that mortality due to aspiration pneumonitis could be reduced if women did not eat and drink during labour. [evidence level 3] This work continues to influence practice both in the UK and elsewhere. In the UK less than 5% (12/268) maternity hospitals have a policy of unrestricted intake during labour, [evidence level 3] this is also usual practice in many other countries. [evidence level 3]

An exception to this is the Netherlands where a survey reported that the majority of obstetricians and midwives had an unrestrictive policy on fluid and food intake. The Netherlands do not have a higher mortality rate due to aspiration pneumonitis than other countries. [evidence level 3] A UK survey of women’s views about eating in labour reported that 31% of women said that would have liked to have eaten during labour. [evidence level 3] Many historical overviews, comments, surveys or non-systematic literature reviews have been written discussing the benefits and harms of eating during labour. [evidence level 3]

One RCT (n = 94) compared offering a low residue diet of toast cereal, crackers and low fat cheese during labour to offering a range of drinks to women during labour (water, tea, coffee, cocoa). Women included in the trial were in spontaneous labour, at term with singleton cephalic presentation and who did not request parenteral opioids (because opioids can delay gastric emptying). Outcome measures used were women’s metabolic profile, volume of gastric contents as well as labour outcomes such as length of labour, use of oxytocin and mode of birth. [evidence level 1b] Women who had a low residue diet were less likely to have ketosis and had higher plasma glucose at the end of labour than women in the drinks only group. Gastric contents were significantly higher in those eating a low residue diet and these women were more likely to vomit at birth, vomit higher volumes and vomit more solid material. Higher gastric volumes could be of importance if unexpected general anaesthesia was needed. No differences were detected in labour outcomes between the two groups but the study is underpowered to evaluate these outcomes. [evidence level 1b] This issue is currently being evaluated in another RCT.

A further RCT (n = 60) compared drinking an isotonic drink to drinking water only during labour. Metabolic indices and gastric volumes were measured. Isotonic drinks reduced ketosis but did not
increase gastric volume. There was no change in labour outcomes but the study was underpowered to assess these outcomes.\textsuperscript{280} [evidence level 1b]

<table>
<thead>
<tr>
<th>Number</th>
<th>Recommendation</th>
</tr>
</thead>
<tbody>
<tr>
<td>50</td>
<td>Women should be informed that eating a low-residue diet during labour (toast, crackers, low-fat cheese) results in larger gastric volumes, but the effect on the risk of aspiration if anaesthesia is required is uncertain. [A] [2004]</td>
</tr>
<tr>
<td>51</td>
<td>Women should be informed that having isotonic drinks during labour prevents ketosis without a concomitant increase in gastric volume. [A] [2004]</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Number</th>
<th>Research Recommendation</th>
</tr>
</thead>
<tbody>
<tr>
<td>27</td>
<td>RCTs that evaluate the effects of eating during labour compared with restricting intake on labour outcomes are needed. Cohort or case control studies on the risk factors for aspiration and other morbidities for women having CS are needed.</td>
</tr>
</tbody>
</table>
7 Procedural aspects of CS

7.1 Timing of planned CS

Babies born preterm are at increased risk of respiratory distress syndrome. One UK survey (n = 179,701) of babies born at 34 weeks gestation or more reported 0.08% (149 babies) had respiratory distress requiring surfactant therapy. Of these babies, 24% (n = 36) were born at or after 37 weeks but 88% (n = 32) of these babies were born by planned CS.\(^\text{281}\) [evidence level 3]

Babies born by planned CS at term (37–42 weeks of gestation) are at risk of respiratory distress syndrome and this decreases with increasing gestational age.\(^\text{282}\) A large prospective UK survey looked at all cases of respiratory distress syndrome (RDS) or transient tachypnoea of the newborn (TTN) at term requiring neonatal intensive care unit (NICU). This study found a decrease in respiratory morbidity from 39 weeks onwards (from 42.3 per 1000 at 38 weeks to 17.8 per 1000 at 39 weeks – OR 8.2 and 3.5 respectively). Respiratory morbidity among neonates born by CS before the onset of labour across the different gestational ages was increased.\(^\text{282}\) [evidence level 3] Figure 7.1 shows respiratory morbidity per 1000 for CS before labour.\(^\text{282}\) [evidence level 3]

From the National Sentinel Caesarean Section Audit [NSCSA] data it is estimated that about 18% of women went into spontaneous labour between 37–39 weeks (see figure 7.2). The average planned CS rate is about 10%. Therefore between 1-2% of women booked for a planned CS after 39 weeks would be expected to go into labour before this time. For an average hospital with 3000 births this would prevent 1 case of TTN or RDS per year and would increase unscheduled CS rate by 18% (assuming timing of planned CS goes from 37 to 39 weeks).

Figure 7.1 Respiratory morbidity per 1000 for CS before labour\(^\text{282}\) [evidence level 3].
The risk of respiratory morbidity is increased in babies born by CS before labour, but this risk decreases significantly after 39 weeks. Therefore planned CS should not routinely be carried out before 39 weeks.

[B] [2004]

### 7.2 Classification of urgency

CS has traditionally been divided into either elective or emergency procedures. More recently these terms have been replaced by “planned” and “unplanned” The unplanned category is broad, as it may include procedures done within minutes to save the life of a woman or baby as well as those in which woman and baby are well but where early delivery is needed, (for example, a woman with a planned CS who is admitted in labour). A clear classification of the perceived degree of urgency of the CS can facilitate communication and reduce misunderstanding between health care professionals. The NCEPOD classification system recommended the categorisation of operations into four grades of urgency. This categorisation scheme has been piloted and evaluated. Although new to most maternity units, there was consistent use of the new scheme when compared with the binary categories and indication for CS. The categorisation also independently predicted baby outcome.

The categories are:

1. Immediate threat to the life of the woman or fetus
2. Maternal or fetal compromise which was not immediately life-threatening
3. No maternal or fetal compromise but needs early delivery
4. Delivery timed to suit woman or staff

Grade 1 (immediate threat to the life of the woman or fetus) includes CS for acute severe bradycardia, cord prolapse, uterine rupture, fetal blood sampling pH less than 7.2. Grade 2 (maternal or fetal compromise which was not immediately life-threatening), there is ‘urgency’ to deliver the baby in order to prevent further deterioration of either the mother or baby’s condition (e.g. antepartum haemorrhage, ‘failure to progress’ in labour with maternal or fetal compromise). Grade 3 (no maternal or fetal compromise but needs early delivery) includes CS carried out where there is no maternal or fetal compromise but early delivery is necessary (e.g. a woman booked for planned CS who is admitted with pre-labour SROM or ‘failure to progress’ with no maternal or fetal compromise). Grade 4
(delivery timed to suit woman or staff) includes all CS carried out ‘electively’ at a planned time to suit the mother and clinicians.

**Recommendations**

<table>
<thead>
<tr>
<th>Number</th>
<th>Recommendation</th>
</tr>
</thead>
<tbody>
<tr>
<td>53</td>
<td>The urgency of CS should be documented using the following standardised scheme in order to aid clear communication between healthcare professionals about the urgency of a CS:</td>
</tr>
<tr>
<td></td>
<td>1 immediate threat to the life of the woman or fetus</td>
</tr>
<tr>
<td></td>
<td>2 maternal or fetal compromise which is not immediately life-threatening</td>
</tr>
<tr>
<td></td>
<td>3 no maternal or fetal compromise but needs early delivery</td>
</tr>
<tr>
<td></td>
<td>4 delivery timed to suit woman or staff. [C] [2004]</td>
</tr>
</tbody>
</table>

### 7.3 Decision-to-delivery interval for unplanned CS

#### Introduction

The appropriate decision-to-delivery interval in unplanned (category 1 and 2) CS remains a controversial issue. This is especially true of the 30 minute interval which has become a critical clinical threshold in clinical practice, despite the fact that in the original guideline, it was intended to be an audit standard and not a recommendation for practice. A further concern is the current lack of distinction between the category 1 and 2 CS in practice. The RCOG Good Practice Guideline no. 11, ‘Classification of urgency of caesarean section – a continuum of risk’ (2010) has advised and strongly recommended that the ‘continuum of risk’ be recognised in the unplanned situation. With these considerations it was felt timely to review the current NICE recommendations in this area.

#### Review question

What is the appropriate decision-to-delivery interval (DDI) for unplanned caesarean section?

#### Overview of evidence

Ten studies were included in this review (Holcroft et al., 2005; Nasrallah et al., 2004; Bloom et al., 2006; Roy et al., 2008; Thomas et al., 2004; Kolas et al., 2006; Hillemanns et al., 2005; Leung et al., 2009; Chauleur et al., 2009; Hillemanns et al., 2003).

Three studies were conducted in the USA (Holcroft et al., 2005; Nasrallah et al., 2004; Bloom et al., 2006), one in India (Roy et al., 2008), one in the UK (Thomas et al., 2004), one in Norway (Kolas et al., 2006), two in Germany (Hillemanns et al., 2003; 2005), one in China (Leung et al., 2009), and one in France (Chauleur et al., 2009).

Six observational studies (Holcroft, et al. 2005; Nasrallah et al., 2004; Bloom et al., 2006; Hillemanns et al., 2003; Roy et al., 2008; Chauleur et al., 2009) examined the effects of less and higher than 30 minutes DDI on neonatal and maternal outcomes. Two studies (Thomas et al., 2004; Kolas et al., 2006) examined the association between different DDI (ranking from less than 15 minutes to higher than 75 minutes) on neonatal and maternal outcomes. One study (Hillemanns et al., 2005) retrospectively examined the correlation between umbilical cord arterial blood pH and decision-to-delivery time. One retrospective study (Leung et al., 2009) investigated the relationship between bradycardia to delivery interval and adverse perinatal outcomes.

Maternal and neonatal outcomes were chosen by the GDG as being of priority to inform recommendations and the results for these are presented in the evidence profile below.
Maternal outcomes

Table 7.1 GRADE findings for comparison of a decision-to-delivery interval of < 30 minutes with a decision-to-delivery interval of > 30 minutes (maternal outcomes)

<table>
<thead>
<tr>
<th>Number of studies</th>
<th>Number of women</th>
<th>Effect</th>
<th>Quality</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>DDI &lt; 30 minutes</td>
<td>DDI &gt; 30 minutes</td>
<td>Relative (95% CI)</td>
</tr>
<tr>
<td>Blood transfusion</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 study (Nasrallah et al., 2004)</td>
<td>6/83 (7.2%)</td>
<td>0/28 (0%)</td>
<td>Not calculable (NC)</td>
</tr>
<tr>
<td>1 study (Hillemanns et al., 2003)</td>
<td>11/109 (10.1%)</td>
<td>1/109 (0.9%)</td>
<td>11 (1.8 to 68)*</td>
</tr>
<tr>
<td>Uterine/bladder rupture</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 study (Hillemanns et al., 2003)</td>
<td>7/109 (6.4%)</td>
<td>8/109 (7.3%)</td>
<td>0.87 (0.34 to 2.24)*</td>
</tr>
<tr>
<td>Ureteric injuries</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 study (Bloom et al., 2006)</td>
<td>2/1814 (0.1%)</td>
<td>1/994 (0.1%)</td>
<td>1.09 (0.14 to 8.35)*</td>
</tr>
<tr>
<td>Cystotomy</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 study (Bloom et al., 2006)</td>
<td>2/1814 (0.1%)</td>
<td>3/994 (0.3%)</td>
<td>0.36 (0.07 to 1.82)*</td>
</tr>
<tr>
<td>Wound complication</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 study (Bloom et al., 2006)</td>
<td>23/1814 (1.3%)</td>
<td>9/994 (0.9%)</td>
<td>1.40 (0.66 to 2.96)*</td>
</tr>
<tr>
<td>Urinary tract infection</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 study (Hillemanns et al., 2003)</td>
<td>3/109 (2.8%)</td>
<td>2/109 (1.8%)</td>
<td>1.5 (0.30 to 7.40)*</td>
</tr>
<tr>
<td>Wound infection</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 study (Hillemanns et al., 2003)</td>
<td>1/109 (0.9%)</td>
<td>5/109 (4.6%)</td>
<td>0.2 (0.03 to 1.26)*</td>
</tr>
</tbody>
</table>

Caesarean section: full guideline DRAFT (September 2011)
### Surgical injuries

<table>
<thead>
<tr>
<th>Study</th>
<th>N/S per 1000</th>
<th>NC per 1000</th>
<th><em>P value range</em></th>
<th><em>Number of additional cases</em></th>
<th>Grade</th>
</tr>
</thead>
<tbody>
<tr>
<td>Caesarean hysterectomy</td>
<td>2/83 (2.4%)</td>
<td>0/28 (0%)</td>
<td>NC</td>
<td>NC</td>
<td>Very low</td>
</tr>
<tr>
<td>Postpartum haemorrhage</td>
<td>2/109 (1.8%)</td>
<td>1/109 (0.9%)</td>
<td>2 (0.26 to 15.1)</td>
<td>9 more per 1000 (from 7 fewer to 129 more)</td>
<td>Very low</td>
</tr>
<tr>
<td>Bowel laceration</td>
<td>1/1814 (0.1%)</td>
<td>1/994 (0.1%)</td>
<td>0.54 (0.05 to 5.24)</td>
<td>0 fewer per 1000 (from 1 fewer to 4 more)</td>
<td>Very low</td>
</tr>
<tr>
<td>Intensive care unit</td>
<td>11/109 (10.1%)</td>
<td>5/109 (4.6%)</td>
<td>2.2 (0.82 to 5.90)</td>
<td>55 more per 1000 (from 8 fewer to 225 more)</td>
<td>Very low</td>
</tr>
<tr>
<td>Endometritis</td>
<td>3/109 (2.8%)</td>
<td>2/109 (1.8%)</td>
<td>1.5 (0.30 to 7.40)</td>
<td>9 more per 1000 (from 13 fewer to 117 more)</td>
<td>Very low</td>
</tr>
<tr>
<td>Special care requirements**</td>
<td>495/3958 (12.5%)</td>
<td>1587/12,606 (12.5%)</td>
<td>0.99 (0.90 to 1.09)</td>
<td>1 fewer per 1000 (from 13 fewer to 11 more)</td>
<td>Very low</td>
</tr>
</tbody>
</table>

1. *Calculated by NCC-WCH
2. **Defined as care following CS that was additional to ‘routine’ post-operative care
Maternal special care requirement is defined as any care above or under standard postnatal care.

A logistic regression analysis was performed adjusting data for primary indication for caesarean section, cardiotocography findings, grade of urgency, and type of anaesthesia. There was no statistically significant difference in the adjusted odds ratios for maternal requirement for special care in women with a DDI < 15 minutes compared with women with DDI > 16 to 75 minutes. However, there was a significantly increased risk of maternal requirement for special care in women with DDI > 75 minutes (OR = 1.5, 95% CI 1.2 to 1.8) compared with neonates born < 15 minutes (OR = 1).

Table 7.2 GRADE summary of findings for comparison of a decision-to-delivery interval of < 30 minutes with a decision-to-delivery interval of > 30 minutes (neonatal outcomes)

<table>
<thead>
<tr>
<th>Number of studies</th>
<th>Number of neonates</th>
<th>Effect</th>
<th>Quality</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>DDI &lt; 30 minutes</td>
<td>DDI &gt; 30 minutes</td>
<td>Relative (95% CI)</td>
</tr>
<tr>
<td><strong>Neonatal deaths</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 study (Holcroft et al., 2005)</td>
<td>1/34 (2.9%)</td>
<td>0/83 (0%)</td>
<td>Not calculable (NC)</td>
</tr>
<tr>
<td>1 study (Bloom et al., 2006)</td>
<td>7/1814 (0.4%)</td>
<td>1/994 (0.1%)</td>
<td>3.83 (0.61 to 23.8)*</td>
</tr>
<tr>
<td><strong>Stillbirth</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 study (Roy et al., 2008)</td>
<td>1/121 (0.8%)</td>
<td>0/96 (0%)</td>
<td>NC</td>
</tr>
<tr>
<td>1 study</td>
<td>27/3958</td>
<td>23/12,606</td>
<td>3.73</td>
</tr>
<tr>
<td>Event</td>
<td>Studies</td>
<td>Incidence</td>
<td>Confidence Interval</td>
</tr>
<tr>
<td>------------------------------------</td>
<td>-------------------------------------------------------------------------</td>
<td>---------------------</td>
<td>---------------------</td>
</tr>
<tr>
<td><strong>Fetal death in labour</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 study</td>
<td>(Bloom et al., 2006)</td>
<td>3/1814 (0.2%)</td>
<td>0/994 (0%)</td>
</tr>
<tr>
<td><strong>Perinatal mortality</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 study</td>
<td>(Hillemanns et al., 2003)</td>
<td>7/124 (5.6%)</td>
<td>3/124 (2.4%)</td>
</tr>
<tr>
<td><strong>5 mins Apgar &lt; 7</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 study</td>
<td>(Holcroft et al., 2005)</td>
<td>3/34 (8.8%)</td>
<td>8/83 (9.6%)</td>
</tr>
<tr>
<td>1 study</td>
<td>(Hillemanns et al., 2003)</td>
<td>21/124 (16.9%)</td>
<td>9/124 (7.3%)</td>
</tr>
<tr>
<td>1 study</td>
<td>(Nasrallah et al., 2004)</td>
<td>8/83 (9.5%)</td>
<td>1/28 (3.6%)</td>
</tr>
<tr>
<td>1 study</td>
<td>(Roy et al., 2008)</td>
<td>18/121 (14.9%)</td>
<td>15/96 (15.6%)</td>
</tr>
<tr>
<td>1 study</td>
<td>(Kolas. et al., 2006)</td>
<td>50/624 (8%)</td>
<td>8/576 (1.4%)</td>
</tr>
<tr>
<td>1 study</td>
<td>(Thomas. et al., 2004)</td>
<td>226/3958 (5.7%)</td>
<td>328/12606 (2.6%)</td>
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<tr>
<td><strong>5 minute Apgar ≤ 3</strong></td>
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<td></td>
</tr>
<tr>
<td>1 study</td>
<td>(Bloom et al., 2006)</td>
<td>18/1814 (1%)</td>
<td>9/994 (0.9%)</td>
</tr>
<tr>
<td><strong>Cord pH &lt; 7.0</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 study</td>
<td>(Hillemanns et al., 2003)</td>
<td>10/124 (8.1%)</td>
<td>0/124 (0%)</td>
</tr>
<tr>
<td>1 study</td>
<td>6/34 (17.6%)</td>
<td>2/83 (2.4%)</td>
<td>8.20 (1.97 to 34.2)*</td>
</tr>
<tr>
<td>----------------------------------------------</td>
<td>--------------</td>
<td>-------------</td>
<td>----------------------</td>
</tr>
<tr>
<td>(Holcroft et al., 2005)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 study</td>
<td>8/121 (6.6%)</td>
<td>5/96 (5.2%)</td>
<td>1.26 (0.45 to 3.59)*</td>
</tr>
<tr>
<td>(Roy et al., 2008)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 study</td>
<td>52/1814 (2.9%)</td>
<td>9/994 (0.9%)</td>
<td>3.16 (1.59 to 6.31)*</td>
</tr>
<tr>
<td>(Bloom et al., 2006)</td>
<td></td>
<td></td>
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<tr>
<td>1 study</td>
<td>5/83 (6%)</td>
<td>0/28 (0%)</td>
<td>NC</td>
</tr>
<tr>
<td>(Nasrallah et al., 2004)</td>
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</tbody>
</table>

**Admission to neonatal intensive care unit (NICU)**

<table>
<thead>
<tr>
<th>1 study</th>
<th>74/124 (59.7%)</th>
<th>65/124 (52.4%)</th>
<th>1.13 (0.91 to 1.42)*</th>
<th>68 more per 1000 (from 74 fewer to 220 more)*</th>
<th>Very low</th>
</tr>
</thead>
<tbody>
<tr>
<td>(Hillemanns et al., 2003)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 study</td>
<td>21/83 (25.3%)</td>
<td>6/28 (21.4%)</td>
<td>1.18 (0.56 to 2.67)*</td>
<td>39 more per 1000 (from 94 fewer to 358 more)*</td>
<td>Very low</td>
</tr>
<tr>
<td>(Nasrallah et al., 2004)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 study</td>
<td>26/121 (21.5%)</td>
<td>7/96 (7.3%)</td>
<td>2.94 (1.38 to 6.43)*</td>
<td>141 more per 1000 (from 28 more to 396 more)*</td>
<td>Very low</td>
</tr>
<tr>
<td>(Roy et al., 2008)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 study</td>
<td>147/624 (23.6%)</td>
<td>104/576 (18.1%)</td>
<td>1.30 (1.04 to 1.63)</td>
<td>54 more per 1000 (from 7 more to 114 more)</td>
<td>Very low</td>
</tr>
<tr>
<td>(Kolas. et al., 2006)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 study</td>
<td>24/25 (96%)</td>
<td>35/46 (76%)</td>
<td>1.26 (1.02 to 1.55)*</td>
<td>198 more per 1000 (from 15 more to 418 more)*</td>
<td>Very low</td>
</tr>
<tr>
<td>(Chauleur et al., 2009)</td>
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</table>

**Seizures**

<table>
<thead>
<tr>
<th>1 study</th>
<th>2/34 (5.9%)</th>
<th>5/83 (6%)</th>
<th>0.97 (0.22 to 4.08)*</th>
<th>2 fewer per 1000 (from 47 fewer to 176 more)*</th>
<th>Very low</th>
</tr>
</thead>
<tbody>
<tr>
<td>(Nasrallah et al., 2004)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Encephalopathy**

<table>
<thead>
<tr>
<th>1 study</th>
<th>5/83 (6%)</th>
<th>0/28 (0%)</th>
<th>NC</th>
<th>NC</th>
<th>Very low</th>
</tr>
</thead>
<tbody>
<tr>
<td>(Nasrallah et al., 2004)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
1 study (Bloom et al., 2006) 12/1814 (0.7%) 5/994 (0.5%) 1.31 (0.48 to 3.57)* 2 more per 1000 (from 3 fewer to 13 more)* Very low

Median NICU stay (days)

1 study (Nasrallah et al., 2004) 13 (range 1-40) 9 (range 3-35) NC 4 Very low

Neonate requiring immediate ventilation

1 study (Roy et al., 2008) 4/121 (3.3%) 2/96 (2.1%) 1.58 (0.34 to 7.31)* 12 more per 1000 (from 14 fewer to 121 more)* Very low

* Calculated by NCC-WCH technical team

Figure 7.4 Stillbirth and 5 minute Apgar < 7 findings from Thomas et al., 2004

A logistic regression analysis was performed adjusting data for primary indication for caesarean section, cardiotocography findings, grade of urgency, and type of anaesthesia.

There was no statistically significant difference in the adjusted odds ratios for 5 minutes Apgar score < 7 in neonates born < 15 minutes compared with the neonates born with DDI > 16 to 75 minutes. However there was a significantly higher odds ratio of a 5 minutes Apgar score < 7 in neonates born with DDI > 75 minutes (OR 1.7, 95% CI 1.2 to 2.4) compared with neonates born < 15 minutes (OR 1).

There was no statistically significant difference in the adjusted odds ratios for stillbirth in neonates born < 15 minutes compared with the neonates born with DDI > 16 to 75 minutes.
Figure 7.5 – NICU admission and 5 minute Apgar < 7 findings from Kolas et al., 2006

![Graph showing NICU admission and 5 minute Apgar < 7 findings from Kolas et al., 2006](image)

Series 1 = NICU admission; DDI < 15 min n = 242 (29.0%); DDI 16 to 30 min n = 382 (23.4%); DDI 31 to 60 min n = 394 (19.3%); DDI > 60 min n = 182 (15.5%) p < 0.01

Series 2 = 5 min Apgar < 7; DDI < 15 min n = 242 (11.0%); DDI 16 to 30 min Term n = 382 (5.9%); DDI 31 to 60 min n = 394 (1.0%); DDI > 60 min n = 182 (2.2%) p < 0.001

One retrospective cohort study (Hillemans et al., 2005) examined the effect of the < 30 minutes DDI of “crash emergency” CS on Apgar and umbilical artery pH. A very short DDI < 20 minutes was inversely correlated to fetal outcome. Babies born by “emergency” CS performed within 19 minutes presented with lower Apgar score after 5 and 10 minutes than those born after a 20 minutes or more DDI (p = 0.003 and p = 0.01 respectively). The umbilical cord pH was not significantly related to decision-to-delivery time (correlation coefficient r =0.36, p > 0.05).

One retrospective cohort study (Leung et al., 2009) examined the relation between fetal bradycardia and DDI, or DDI to cord arterial pH according to different causes of fetal distress. The causes of the bradycardia were reviewed and categorized into; ‘irreversible’ (median DDI = 10 min), ‘potentially reversible’ (median DDI = 11.5 min) and ‘unknown cause’ (median DDI = 11 min).

The median cord arterial pH was lower in babies born in the ‘irreversible’ group than in the ‘potentially reversible’ group or ‘unknown’ group (p < 0.001). No relationship was found between cord arterial pH and base excess with either bradycardia to delivery interval or DDI in ‘irreversible’, ‘potentially reversible’ and ‘unknown cause’. However, in subgroup analysis, the cord arterial pH was significantly inversely correlated with bradycardia to delivery interval in the ‘irreversible’ group (Spearman’s rho = -0.354; p = 0.027), but no significant inverse correlation was seen in the other two groups.

**Evidence statements**

**Maternal outcomes**

**Blood transfusion**

One study found that blood transfusion in women with a DDI < 30 minutes was higher than in women with DDI > 30 minutes. This finding was statistically significant. Another study investigated this outcome but the statistical significance was not calculable. The evidence for this outcome was of very low quality.

**Uterine/bladder rupture**

One study did not find a statistically significant difference in the rate of uterine/bladder rupture for women with a DDI < 30 minutes compared with women with DDI > 30 minutes. The evidence for this outcome was of very low quality.
Ureteric injuries
One study did not find a statistically significant difference in the rate of ureteric injuries for women with a DDI < 30 minutes compared with women with DDI > 30 minutes. The evidence for this outcome was of very low quality.

Cystotomy
One study did not find a statistically significant difference in the rate of inadvertent cystotomy for women with a DDI < 30 minutes compared with women with DDI > 30 minutes. The evidence for this outcome was of very low quality.

Wound complication
One study did not find a statistically significant difference in the rate of wound complication for women with a DDI < 30 minutes compared with women with DDI > 30 minutes. The evidence for this outcome was of very low quality.

Urinary tract infection
One study did not find a statistically significant difference in the rate of urinary tract infection for women with a DDI < 30 minutes compared with women with DDI > 30 minutes. The evidence for this outcome was of very low quality.

Wound infection
One study did not find a statistically significant difference in the rate of wound infection for women with a DDI < 30 minutes compared with women with DDI > 30 minutes. The evidence for this outcome was of very low quality.

Surgical injuries
One study did not find a statistically significant difference in surgical injuries for women with a DDI < 30 minutes compared with women with DDI > 30 minutes. The evidence for this outcome was of very low quality.

Caesarean hysterectomy
One study investigated the outcome of caesarean hysterectomy in women with a DDI < 30 minutes compared with women with DDI > 30 minutes. The statistical significance of this finding was not calculable in this study. The evidence for this outcome was of very low quality.

Postpartum haemorrhage
One study did not find a statistically significant difference in postpartum haemorrhage for women with a DDI < 30 minutes compared with women with DDI > 30 minutes. The evidence for this outcome was of very low quality.

Bowel laceration
One study did not find a statistically significant difference in the rate of bowel laceration for women with a DDI < 30 minutes compared with women with DDI > 30 minutes. The evidence for this outcome was of very low quality.

Intensive care unit admission
One study did not find a statistically significant difference in the rate of intensive care unit admission for women with a DDI < 30 minutes compared with women with DDI > 30 minutes. The evidence for this outcome was of very low quality.

Endometritis
One study did not find a statistically significant difference in the rate of endometritis for women with a DDI < 30 minutes compared with women with DDI > 30 minutes. The evidence for this outcome was of very low quality.

Special care requirements
One study did not find a statistically significant difference in maternal requirements for special care for women with a DDI < 30 minutes compared with women with DDI > 30 minutes. The evidence for this outcome was of very low quality.
Neonatal outcomes

Neonatal deaths
One study did not find a statistically significant difference in the number of neonatal deaths in neonates born with a DDI < 30 minutes compared with neonates born with a DDI > 30 minutes. Another study investigated the same outcome but the statistical significance was not calculable. The evidence for this outcome was of very low quality.

Stillbirth
One study found a higher number of stillbirths in neonates born with a DDI < 30 minutes compared with neonates born with a DDI > 30 minutes. This finding was statistically significant. A second study investigated the same outcome but the statistical significance was not calculable. The evidence for this outcome was of very low quality.

Stillbirth
One study did not find a significant difference in the number of stillbirths in neonates born with a DDI < 15 minutes compared with neonates born with a DDI > 16-75 minutes. The evidence for this outcome was of very low quality.

Fetal death in labour
One study investigated the outcome of fetal death in neonates born with a DDI < 30 minutes compared with neonates born with a DDI > 30 minutes. The statistical significance of this finding was not calculable. The evidence for this outcome was of very low quality.

Perinatal mortality
One study did not find a statistically significant difference in the perinatal mortality in neonates born with a DDI < 30 minutes compared with neonates born with a DDI > 30 minutes. The evidence for this outcome was of very low quality.

5 minutes Apgar < 7
Three studies found a higher rate of 5 minutes Apgar score < 7 in neonates born with a DDI < 30 minutes compared with neonates born with a DDI > 30 minutes. This finding was statistically significant. Three further studies did not find a statistically significant difference in this outcome between the two groups. The evidence for this outcome was of very low quality.

One study did not find a statistically significant difference in the adjusted odds ratio of 5 minutes Apgar score < 7 in neonates born with a DDI < 15 minutes compared with neonates born with a DDI > 16-75 minutes. However there were significantly higher odds of 5 minutes Apgar score < 7 in neonates born with DDI > 75 minutes. The evidence for this outcome was of very low quality.

One study found a higher rate of 5 minutes Apgar score < 7 in neonates born with a DDI < 15 minutes compared with neonates born with a DDI > 16-30 minutes, DDI > 31-60 minutes, and DDI > 60 min. This finding was statistically significant. The evidence for this outcome was of very low quality.

5 minutes Apgar ≤ 3
One study did not find a statistically significant difference in the number of neonates with 5 minutes Apgar score ≤ 3 in neonates born with DDI < 30 minutes compared with neonates born with a DDI > 30 minutes. The evidence for this outcome was of very low quality.

One study did not find a statistically significant difference in lower Apgar score after 5 and 10 min in neonates born within 19 minutes compared with neonates born ≥ 20 minutes. The evidence for this outcome was of very low quality.

Cord pH
Two studies found a higher rate of cord pH < 7.0 in neonates born with a DDI < 30 minutes compared with neonates born with a DDI > 30 minutes. This finding was statistically significant. A third study did not find a statistically significant difference for this outcome. Two further studies investigated this outcome but the statistical significance was not calculable. The evidence for this outcome was of very low quality.

One study did not find a statistically significant relationship between a low umbilical cord pH and the DDI interval. The evidence for this outcome was of very low quality.
One study did not find a statistically significant difference in median cord arterial pH in neonates born with median DDI = 10 minutes, median DDI = 11 minutes and median DDI = 11.5 minutes. The evidence for this outcome was of very low quality.

**NICU admissions**

Three studies found a higher number of NICU admission in neonates born with a DDI < 30 minutes compared with neonates born with a DDI > 30 minutes. This finding was statistically significant. Two further studies did not find a statistically significant difference for this outcome. The evidence for this outcome was of very low quality.

One study found a higher rate of NICU admissions in neonates born with a DDI < 15 minutes compared with neonates born with DDI > 16-30 minutes, DDI > 31-60 minutes and DDI > 60 minutes. This finding was statistically significant. The evidence for this outcome was of very low quality.

**Seizures**

One study investigated the number of seizures in neonates born with a DDI < 30 minutes compared with neonates born with a DDI > 30 minutes. The statistical significance of this finding was not calculable. The evidence for this outcome was of very low quality.

**Encephalopathy**

Two studies investigated the incidence of encephalopathy in neonates born with a DDI < 30 minutes compared with neonates born with a DDI > 30 minutes. One study did not find a statistically significant difference between the two groups, and the other study, the statistical significance of the finding was not calculable. The evidence for this outcome was of very low quality.

**NICU stay**

One study did not find a statistically significant difference in the length of NICU stay in neonates born with DDI < 30 minutes compared with neonates born with a DDI > 30 minutes. The evidence for this outcome was of very low quality.

**Neonate requiring immediate ventilation**

One study did not find a statistically significant difference in the length of NICU stay in neonates born with DDI < 30 minutes compared with neonates born with a DDI > 30 minutes. The evidence for this outcome was of very low quality.

**Evidence to recommendations**

**Relative value placed on outcomes considered**

The group was keen to see whether there was any evidence that a DDI of less than 30 minutes was related to poorer maternal outcomes as there is a concern that performing CS too quickly can lead to iatrogenic injury. Given this, the group rated all of the maternal outcomes as being important in determining whether or not this is the case.

In terms of neonatal outcomes, the group recognised that there is a treatment paradox: the babies who are delivered the quickest are likely to be the ones who are most compromised, and are therefore more likely to have poorer outcomes. As a result, samples of babies born within 30 minutes will consistently contain a higher proportion of babies in poorer condition. Thus, differences in findings between groups might reflect this disparity, rather than differences due to speed of delivery. Given this, the group did not feel able to attach as much weight to the neonatal findings.

**Trade off between clinical benefits and harms**

The GDG noted there is a trade-off between the baby being born as quickly as possible against the risk of injuring the mother or the baby.

The group went on to consider appropriate DDI in relation to varying degrees of urgency as classified in this guideline and repeated in the RCOG Good Practice Guideline no 11, ‘Classification of urgency of caesarean section – a continuum of risk’ (2010). There was no evidence that performing a CS within 30 minutes resulted in greater injury to the woman. However, whilst they agreed that in general the CS should be accomplished as quickly as reasonably possible, the GDG still felt there are occasions in which a very rapid delivery could do harm. The group did not feel therefore that it was appropriate to recommend a time within which a category 1 CS should be performed.
It was agreed that a 30 minute DDI is useful as an audit standard. However, it was felt important to highlight that this should only be used as a standard by which to measure the performance of an obstetric unit as a whole. It should not be used as a clinical standard and should not be used to judge the quality of care in individual cases.

The group recognised that there was evidence of adverse neonatal and maternal outcomes in category 2 CS which have a DDI longer than 75 minutes. Whilst the group recognised that there are particular instances where it would not be appropriate to perform a category 2 CS before 75 minutes (e.g. where maternal blood pressure needs stabilising or essential specialist health is being awaited), in the large majority of cases, clinicians should aim to perform category 2 CS within this time.

**Trade off between net health benefits and resource use**

The trade off being considered in the clinical situation where an unplanned CS is necessary is to ensure safe birth of the baby in as good a condition as possible without causing harm to the woman or her baby through iatrogenic injury or mistakes made due to carrying out the procedure with too much haste. Issues of resource use mirror these considerations in that iatrogenic injury has the potential to be costly as well as causing pain and distress to the woman. Similarly unwarranted delay also has the potential to be hugely damaging in terms of the baby's health. Thus optimal timing of birth will be both clinically effective in terms of the health outcomes for the woman and her baby and cost-effective in terms of resource use.

**Quality of evidence**

The group recognised that the quality of the evidence was low as all of the studies included in the review were retrospective observational studies.

Whilst there were statistically significant findings which indicated poorer outcomes for babies born before 30 minutes, the group agreed that this was due to the treatment paradox noted above i.e. that the most compromised babies are those who require the fastest intervention.

The group had anticipated that there might be evidence to show that there were more iatrogenic injuries to the woman where the CS had been performed with a DDI of less than 30 minutes. However, in the evidence reviewed, surgical injury was found not be significantly different between the two groups (under 30 minutes vs. over 30 minutes). The low quality of all the included studies meant the group were less certain of the reliability of this finding, although it was noted that it was a consistent finding across a number of studies reporting different types of injury.

There was evidence from one study of a significantly higher need for blood transfusion in women with a DDI of less than 30 minutes. However, it was not possible to determine the reasons for this. The group felt that it was not possible to determine whether this was a consequence of the rapid delivery, or a reason for it.

**Other considerations**

The group wished to distinguish between a clinical standard and an audit standard. Whilst a clinical standard indicates the care which should be provided in each individual case, an audit standard indicates the overall level of care which should be provided by a unit. In the case of the DDI of 30 minutes, the group recognised that this was inappropriate as a clinical standard – in a number of instances (e.g. complete cord-occlusion), a DDI of 30 minutes would be too long, whilst in others (e.g. when necessary to stabilise the woman’s clinical condition) some delay would be appropriate. However, the group agreed that 30 minutes was a useful standard by which to assess the performance of a unit – whilst in individual cases, a DDI of 30 minutes might not be appropriate, overall, there would be cause for concern if the vast majority of category 1 CS were not carried out within this time.

<table>
<thead>
<tr>
<th>Number</th>
<th>Recommendation</th>
</tr>
</thead>
<tbody>
<tr>
<td>54</td>
<td>Perform category 1 and 2 CS[9] as quickly as possible after making the decision, particularly for category 1. [new 2011]</td>
</tr>
</tbody>
</table>

[9] Category 1 CS is when there is immediate threat to the life of the woman or fetus, and category 2 CS is when there is maternal or fetal compromise which is not immediately life-threatening.
55 Perform category 2 CS\(^{10}\) in most situations within 75 minutes of making the decision. [new 2011]

56 Take into account the condition of the woman and the unborn baby when making decisions about rapid delivery. Remember that rapid delivery may be harmful in certain circumstances. [new 2011]

57 Use the following decision-to-delivery intervals to measure the overall performance of an obstetric unit:

- 30 minutes for category 1 CS\(^{10}\)
- both 30 and 75 minutes for category 2 CS.

Use these as audit standards only and not to judge multidisciplinary team performance for any individual CS. [new 2011]

<table>
<thead>
<tr>
<th>Number</th>
<th>Research Recommendation</th>
</tr>
</thead>
<tbody>
<tr>
<td>RR 28</td>
<td>What factors influence the decision-to-delivery interval when there is a category 1 level of urgency for CS?</td>
</tr>
</tbody>
</table>

Factors to be investigated could include:

- staff grade/level of experience
- skill mix within the multidisciplinary team
- task allocation
- methods of communication
- time of day
- availability of ongoing staff training about emergency procedures and levels of attendance.

The research could be conducted using simulation methods and video observation to determine what factors influence the decision-to-delivery interval for category 1 CS. The videos could also be used to train staff.

**Why this is important**

‘Crash’ CS is a psychologically traumatic event for women and their partners and is also stressful for clinical staff. Staff and resources may have to be obtained from other areas of clinical care. This should be undertaken as efficiently and effectively as possible, minimising anxiety and ensuring the safety of the mother and her baby.

For category 1 CS there is a recognised urgency to deliver as quickly as is reasonably possible. The majority of research in this area is quantitative and looks at the impact of the decision-to-delivery interval on various aspects of fetal and maternal outcomes rather than the interplay of factors that can affect this time period itself. Much of this evidence is retrospective. Although some work has been conducted in the UK to examine where the systematic delays lie and how to avoid them (Tuffnell et al., 2001), more work is needed to determine how to optimise the decision-to-delivery interval. This work should use qualitative as well as quantitative research methods to assess which factors influence the decision-to-delivery interval for a category 1 CS. Evaluation of these factors could be used to inform future NICE guidance, for example specific guidance for management of category 1 CS. Such information could also be used by hospitals for maternity services planning and at a team level would assist with audit and ongoing evaluation and training of the multidisciplinary team.

A large amount of NHS and other state funding is used to provide continuing care for

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\(^{10}\) Category 1 CS is when there is immediate threat to the life of the woman or fetus, and category 2 CS is when there is maternal or fetal compromise which is not immediately life-threatening.
infants who are disabled as a result of birth asphyxia and in providing lifelong support for the child and their family. In addition, large sums of public money are spent on litigation and compensation in some of these cases through the Clinical Negligence Scheme for Trusts (CNST). If research helped to minimise the impact of birth asphyxia this would reduce the costs of continuing care to the state and the burden to the child, their family and the wider community.

More realistic and more relevant expectations for the decision-to-delivery interval based on evidence would inform debate in the legal system and may help to reduce the cost to the state of related litigation.

A prospective study to determine whether the decision-to-delivery interval has an impact on maternal and neonatal outcomes when there is a category 2 level of urgency for CS.

Important primary outcomes would be
- fetal wellbeing (such as cord blood gases, Apgar score at 5 minutes, hypoxic encephalopathy, neonatal respiratory problems, unanticipated admission to neonatal intensive care unit (NICU), duration of stay in the NICU)
- maternal wellbeing (such as haemoglobin levels on day 2, need for blood transfusion, duration of hospital stay controlled for prolonged neonatal stay and general health/wellbeing).

Valuable secondary outcomes could include:
- fetal trauma at delivery
- iatrogenic maternal bladder or bowel injury
- postoperative maternal infectious morbidity
- establishment of breast-feeding
- psychological outcomes for women, such as the development of postnatal depression/post-traumatic stress disorder.

Why this is important
This research is important to inform the ongoing debate about the management of category 2 CS. The ‘continuum of risk’ in this setting has been recognised. However, the majority of work in this area, looking at maternal and fetal outcomes, generally considers unplanned caesarean sections as a whole group without making any distinction between degrees of urgency. Furthermore much of this work is retrospective. The majority of women who undergo intrapartum CS fall into the category 2 level of urgency (Thomas et al., 2001) and therefore specific information for this group could affect and benefit many women and contribute to the delivery of equity of care.

Delay in delivery with a compromised fetus may result in major and long-term harm including cerebral palsy and other major long-term disability. The immediate and long-term effect on a family of the birth of a baby requiring life-long specialised care and support is enormous. If such harm could be avoided by appropriate haste this would be an important improvement in outcome. However, if such haste is of no benefit then any related risk of adverse maternal outcome needs to be minimised.

A large amount of NHS and other state funding is used to provide continuing care for infants who are disabled as a result of delay in delivery and in providing lifelong support for the child and their family. In addition, large sums of public money are spent on litigation and compensation in some of these cases through the Clinical Negligence Scheme for Trusts (CNST). If research helped to minimise the impact of delay in delivery this would reduce the costs of continuing care to the state and the burden to the child, their family and the wider community.

More realistic and more relevant expectations for the decision-to-delivery interval based on evidence would inform debate within the legal system and may help to reduce the cost to the state of related litigation.
Repeat of the National Caesarean Section Sentinel Audit

The original CS guideline included a set of ‘auditable standards’. It would be a straightforward task to produce an updated set of auditable standards based on the important topics covered in the updated guideline. These could include:

- consent
- indications (including maternal request)
- procedural aspects
- maternal and fetal outcomes.

Many of the outcomes documented in a new CS audit would relate directly to recommendations in this CS guideline update. Researchers may also want to consider categorising different reasons underlying maternal request for CS such as previous poor childbirth experience, longstanding fear of childbirth, belief that CS is safer for the baby etc.

An additional useful feature of the audit would be to record key related data, such as the proportion of CS for a breech presentation that had an attempted external cephalic version.

Why this is important

During the 10 years since the National Caesarean Section Sentinel Audit was undertaken (2000–2001), many of the findings may have changed significantly. The audit examined who was having a CS and why, as well as the views of women having babies and the obstetricians looking after them. The audit found that a 20% CS rate was considered too high by 51% of obstetricians. UK CS rates now average about 25%.

A repeat of the CS Sentinel Audit would reveal any changes in indications and the views of women and obstetricians. The current literature does not adequately address the issue of maternal request for CS and this is one aspect the audit may address. Women’s views on maternal request for CS for when there are no obstetric indications are particularly relevant. Such requests may be on the rise and the reasons are not always clearly expressed or documented.

The methodology of the audit is established, making a repeat feasible. This should be given high priority because the benefit to the NHS would be significant.

7.4 Preoperative testing and preparation for CS

Full blood count and haemoglobin

Recommendations for antenatal screening include measuring haemoglobin (Hb) at booking and repeating this at 28 weeks of gestation to screen for anaemia. Pregnancy increases maternal iron requirements and antenatal screening enables women who have anaemia to receive appropriate treatment before birth. Women who are anaemic at the time of birth are likely to be less able to tolerate blood loss. Overall it is estimated that about 1.3% of all women giving birth have blood loss in excess of 1000 ml, while 0.7% have blood loss in excess of 1500 ml however measurements of blood loss at birth are reliant on visual estimations and are usually underestimations. In the NSCSA 32% of women who had CS had an estimated blood loss between 500–1000 ml, while for 4% it was in excess of 1000 ml. Haemorrhage remains an important cause of maternal mortality.

Two pragmatic RCTs comparing planned CS to planned vaginal birth report blood loss as an outcome measure. (n = 2281) No difference in blood loss greater than 1000 ml or 1500 ml between the two groups was detected (0.5% planned CS; 0.7% planned vaginal birth group, pooled RR 0.80, 95% CI 0.29 to 2.18. For blood loss greater than 1500 ml, pooled RR 1.32, 95% CI 0.39 to 4.42. [evidence level 1a] Non intention to treat analysis (by actual rather than intended mode of delivery) indicate that...
blood loss greater than 1000 ml occurred in 2.7% of women who had CS and 1.6% of women who had vaginal birth. Blood loss greater than 1500 ml occurred in 2% of women who had CS compared to none of the women who had vaginal birth.44 [evidence level 2]

A large UK cohort study291 reported that compared to women who had spontaneous vaginal deliveries, the risk of blood loss in excess of 1000ml was greater among women who had either planned CS (RR 3.94, 99% CI 2.52 to 6.17), CS in labour (RR 8.84, 99% CI 6.74 to 11.6) or assisted vaginal birth (RR 2.39, 95% CI 1.64 to 3.48). Compared with women who had planned CS, risk of blood loss in excess of 1000 ml was higher among women who had CS in labour (RR 2.24, 95% CI 1.43 to 3.53) [evidence level 2b]. However these relative risks do not take into account any other factors that may also affect blood loss, for example the reasons for performing CS in labour such as placental abruption or ante partum haemorrhage.

No studies have evaluated the effect of preoperative Hb or full blood count (FBC) on management or maternal health outcomes. Guidelines for preoperative testing in general surgery have been developed.293 [evidence level 3] The guideline divides surgical procedures into four grades; minor, intermediate, major, major+, neurosurgery and cardiovascular surgery. CS would be classed as major surgery. Patients are then classified according to American Society of Anaesthesia (ASA) grades. In most instances women having CS are ASA grade 1; that is a normal healthy patient without any co-morbidity. The recommendations in the guideline are based on case series, indirect evidence and consensus methodology. The guideline recommends full blood count before major surgery in healthy adults aged 16–40 years.

**Availability of blood and group and saving of serum**

Blood transfusion may be necessary in cases of severe obstetric haemorrhage and is a surrogate marker for heavy blood loss. Six RCTs report on the need for blood transfusion as an outcome measure42,44–48 (n = 2469). 1.4% of women in the planned CS group compared to 1.8% in the planned vaginal birth group required blood transfusion. No difference was detected in this outcome measure between the two groups (pooled RR 0.86, 95% CI 0.48 to 1.53). [evidence level 1a] Non intention to treat analysis (by actual rather than intended mode of delivery), indicate the rate of blood transfusion for women who had CS was 9–10% compared to 0–2% for women who had a vaginal birth.43,44 One cohort study reported on peripartum blood transfusion by mode of birth.294 The overall incidence of blood transfusion following birth was 0.99%. Compared to women who had spontaneous vaginal birth, the relative risk of blood transfusion for women who had CS was 5.6 (95% CI 2.9 to 10.8) and for women who had assisted vaginal birth it was increased (RR 15.5, 95% CI 8.3 to 29.0). [evidence level 2b]

National data on CS for the United Kingdom shows women who had CS for antepartum haemorrhage, placenta praevia or uterine rupture accounted for 21% of occurrences of blood loss greater than 1000 ml.4 [evidence level 3] Women with a prior diagnosis of placenta praevia, abruption, uterine rupture or APH are at increased risk of blood loss of more than 1000 ml (RR 5.31, 95% CI 4.67 to 6.04) compared with women without these conditions. Other predictive factors for haemorrhage during CS include pre-eclampsia, obesity, amnionitis and prolonged active phase of labour.295,296 [evidence level 3]

Haemorrhage is still an important cause of maternal mortality and it is recommended that all obstetric units should have a protocol for the management of obstetric haemorrhage and that women at high risk of haemorrhage should be delivered at a unit with a blood bank on site.95 [evidence level 3] The majority (95%) of maternity units in England and Wales report having on-site cross matching facilities at all times with 3% of maternity units cross matching facilities during the day only and the remainder keeping O-negative blood on labour ward at all times.5 [evidence level 3] There is also a wide range of blood ordering practices.297 [evidence level 3] Blood transfusion service guidelines do not address preoperative cross matching, rather provide recommendations for safer blood transfusion practices.298 [evidence level 4]

We did not identify any studies that looked at whether all women having CS should have group and save taken preoperatively. Women who are at high risk of having a blood loss of greater than 1000 ml at CS should be delivered at a site with blood transfusion services. Studies set in circumstances where there are no blood transfusion services suggest that availability of blood is of importance in reducing the morbidity associated with haemorrhage.299 [evidence level 3]
Other blood tests

We did not identify any evidence on the value of clotting screen or other blood tests prior to CS. Extrapolation from the preoperative testing guideline for major surgery mentioned previously would not recommend clotting screen or other tests such as urea and electrolytes prior to CS.\[293] [evidence level 3]

Routine ultrasound before CS

Preoperative ultrasound has been proposed for placental localisation, presentation and as a method of predicting the integrity of a previous CS scar. A cohort study looked at whether routine preoperative ultrasound at CS impacted on CS outcomes. The study performed preoperative ultrasound scans on 124 women and compared them with matched controls, retrospectively. The outcomes they considered were incidence of incision through the placenta, blood loss of more than 1000 ml, difficult birth; injury of the infant, injury to the cord or to other adjacent structures. No difference in these outcomes was detected between the two groups.\[300] [evidence level 2b]

It has been reported that about a quarter (28%) of transverse uterine scars can be seen on ultrasound, vertical uterine scars cannot be visualised on ultrasound.\[301] [evidence level 2b] The clinical usefulness of this is not clear.\[302] [evidence level 2a].\[303] [evidence level 1b]

Ultrasound has been used for the antenatal diagnosis of placenta accreta however the predictive value of this remains uncertain.\[304,305] [evidence level 3]

Urinary catheter use at CS

A UK survey of obstetricians reports that for CS with epidural anaesthesia the majority (82%) use an indwelling urethral catheter for both the procedure and postoperatively, a minority would use an indwelling catheter for either the duration of the procedure only (10.6%) or an in–out catheter (7.3%). This was similar for both unplanned or planned CS and for CS with general anaesthesia.\[306] [evidence level 3]

An RCT (n = 50) of women undergoing planned caesarean section under epidural analgesia who were randomised prospectively to be catheterised with an 'in–out' or an indwelling urethral catheter removed the after the CS. Of the women who had catheterisation for the time of surgery alone 44% subsequently required re-catheterisation, whereas all women with indwelling catheters voided spontaneously on their removal. The frequency of significant bacteriuria was the same in both groups.\[307] [evidence level 1b]

Another RCT from Iran (n = 270) included women having a CS with general or regional anaesthesia. Women were randomised into two groups: group I were not catheterised but were encouraged to void urine immediately prior to the CS; group II had indwelling catheters removed the day after the CS. Outcomes measured were discomfort at first voiding post-CS, time of ambulation, time of hospital stay and need for re-catheterisation. Of women who were not catheterised 4% required catheterisation postoperatively. There was no difference in ambulation time and women who did not have an indwelling catheter had a slightly shorter hospital stay (17 hours).\[308] [evidence level 1b]

Preoperative shaving

No RCTs have compared pre-CS shaving of the abdomen to no shaving. A systematic review included 2 RCTs (n = 539) to assess the effects of routine perineal shaving on admission in labour on maternal and neonatal outcomes. In the earlier trial, 389 women were alternately allocated to receive either skin preparation and perineal shaving (control) or clipping of vulval hair only (experimental). In the second trial, which included 150 participants, perineal shaving was compared with the cutting of long hairs for procedures only. The primary outcome for both trials was maternal febrile morbidity. No differences were found (combined OR 1.26, 95% CI 0.75 to 2.12). In the smaller trial, fewer women who had not been shaved had gram negative bacterial colonisation compared with women who had been shaved (OR 0.43, 95% CI 0.20 to 0.92).\[309] [evidence level 1a]
Pregnant women should be offered a haemoglobin assessment before CS to identify those who have anaemia. Although blood loss of more than 1000 ml is infrequent after CS (it occurs in 4-8% of CS) it is a potentially serious complication. [C] [2004]

Pregnant women having CS for antepartum haemorrhage, abruption, uterine rupture and placenta praevia are at increased risk of blood loss of more than 1000 ml and should have the CS carried out at a maternity unit with on-site blood transfusion services. [C] [2004]

Pregnant women who are healthy and who have otherwise uncomplicated pregnancies should not routinely be offered the following tests before CS:

- grouping and saving of serum
- cross-matching of blood
- a clotting screen
- preoperative ultrasound for localisation of the placenta, because this does not improve CS morbidity outcomes (such as blood loss of more than 1000 ml, injury of the infant, and injury to the cord or to other adjacent structures). [C] [2004]

Women having CS with regional anaesthesia require an indwelling urinary catheter to prevent over-distension of the bladder because the anaesthetic block interferes with normal bladder function. [GPP] [2004]

### 7.5 Anaesthesia for CS

#### Planning post-CS analgesia

The different options for post-CS analgesia should be discussed with the woman before her CS using available obstetric anaesthesia and analgesia patient information booklets so that the individual analgesic needs of each woman can be met. [evidence level 3] Post-CS pain relief should be prescribed prior to discharge from the anaesthetic recovery area to a general ward. [evidence level 3]

#### General versus regional anaesthesia for CS

The NSCSA reported that 77% of unplanned and 91% of planned CS are performed using regional anaesthesia. Of the CS that were reported to be grade 1 urgency (immediate threat to the life of the woman or fetus), 41% were performed using general anaesthesia, 54% had regional anaesthesia and 3% had general anaesthesia following regional anaesthesia. A UK survey of anaesthetic techniques for CS reported an overall failure rate of epidural anaesthesia of 7.1%, for combined spinal epidural’s it was 2% and for single shot spinal anaesthetic 1.9%. Failure of regional anaesthesia accounted for 10% of general anaesthetic cases for CS. [evidence level 3]

Three RCTs have compared the impact of general versus regional anaesthesia for CS on maternal and neonatal morbidity. One RCT (n = 341) randomised women into three groups: general anaesthesia, epidural anaesthesia or spinal anaesthetic. The maternal and neonatal outcomes were reported separately. [evidence level 1b] General anaesthesia resulted in increased blood loss, lower postoperative haematocrit and higher proportion of women with postoperative haematocrit of less than 30%. There was no difference in neonatal cord blood gas analysis, Apgar and a Neurologic Adaptive Capacity Score (4 hours after birth). [evidence level 1b] The second RCT (n = 47) randomised women to have either general or epidural anaesthesia, the trial measured neonatal outcomes only. No difference was detected in the incidence of low Apgar scores and umbilical artery gas analysis. [evidence level 1b] The third RCT (n = 104) randomised women having planned repeat CS to either general anaesthesia or spinal anaesthesia. The RCT measured short term neonatal outcomes only. This RCT is poorer quality because it has 20% loss to follow-up. Of the 84
infants followed up no difference was detected in neonatal outcomes between the two groups.\textsuperscript{316} All the RCTs are underpowered to look at neonatal outcomes.

A large observational study from the US (n = 3940) reported that infants born by CS with general anaesthesia are more likely to have an Apgar less than 7 and to need resuscitation compared to those born by CS with regional anaesthesia (1-minute Apgar less than 7: RR 3.13, 95% CI 2.5 to 3.88. 5-minute Apgar less than 7: RR 3.6, 95% CI 1.81 to 7.00. Need for resuscitation RR 2.02, 95% CI 1.39 to 2.9).\textsuperscript{317} [evidence level 3]

Two RCTs compared regional and general anaesthetic for specific clinical conditions; severe pre-eclampsia and placenta praevia. One RCT (n = 80) compared general, epidural or combined spinal epidural anaesthetic for CS in women with severe pre-eclampsia. They found no significant difference in maternal (BP or urine output) or fetal complications (umbilical artery pH, Apgar score) between the three groups.\textsuperscript{318} [evidence level 1b] The second RCT (n = 25) randomised women having CS for placenta praevia to receive either general or epidural anaesthesia. Women who received general anaesthesia had lower postoperative haematocrit (28.1% versus 32.5%) and were more likely to need blood transfusion (42% versus 15%; RR 2.71, 95% CI 0.64 to 11.4). There was no difference in neonatal outcomes.\textsuperscript{319} [evidence level 1b] Two large scale retrospective surveys comparing regional to general anaesthesia for CS for placenta praevia showed that general anaesthesia was an independent predictor for increased blood loss, decreased postoperative haemoglobin and increased need for blood transfusion. One of the surveys was conducted in the USA (514 women)\textsuperscript{320} [evidence level 3] and one in the UK (350 women).\textsuperscript{321} [evidence level 3]

A UK-based retrospective survey of 137 women reported that the mean time for surgical readiness for regional anaesthesia 27.6 minutes (range 13–55 minutes) compared with 15.4 minutes (range 2–44 minutes) for general anaesthesia, p < 0.01. Time for surgical readiness is defined as time between leaving the delivery room to skin incision.\textsuperscript{322} [evidence level 3]

**Monitoring during anaesthesia for CS**

For CS under regional block the following monitoring is recommended; continuous pulse oximetry, non-invasive blood pressure capable of one minute cycles (preferably with printout) and continuous ECG are required during induction, maintenance and recovery. The fetal heart rate should be recorded during the initiation of regional block and until the abdominal skin preparation is begun in unplanned CS.\textsuperscript{323} [evidence level 4]

During general anaesthesia, the woman should be monitored in accordance with the recommendations of the Association of Anaesthetists of Great Britain and Ireland (AAGBI) guidelines for obstetric anaesthesia services. The recommendations include continual assessment of the patient’s physiological state, depth of anaesthesia and function of equipment. Monitoring devices supplement clinical observations.\textsuperscript{324} [evidence level 4]

No economic studies comparing the cost effectiveness of general and regional anaesthesia for CS were identified. However we identified one economic study from America using effectiveness data from a case note review comparing spinal and epidural anaesthesia for planned CS. Spinal anaesthesia took up less operating time, required less intraoperative analgesia, and led to fewer complications than epidural. The only dimension which was not different between spinal anaesthesia and epidural was in the need for postoperative analgesia. Therefore spinal anaesthesia was associated with lower cost than epidural anaesthesia (postoperative analgesia was not included in the costs). A full cost-effectiveness analysis was not undertaken.\textsuperscript{325}

**Place of induction of anaesthesia**

There are no RCTs looking at the use of anaesthetic rooms in obstetric anaesthesia. One RCT (n = 100) patients having minor or intermediate operative procedures who were randomised to induction of anaesthesia in an anaesthetic room versus in theatre. The outcomes included patient anxiety assessed using physical parameters (such as heart rate) and questionnaire. There was no difference detected between the two groups.\textsuperscript{326} [evidence level 1b]

A survey of 115 women having a planned CS under regional anaesthesia in the UK reported that stress scores were higher in theatre. Women reported this to be due to anxiety about pain and the well being of themselves and their babies and not from the environment.\textsuperscript{327} [evidence level 3]
Anaesthetic rooms for induction of anaesthesia have been used in the United Kingdom for many years and are currently more commonly used than theatre for induction (4% of UK hospitals induce anaesthesia in theatre).\textsuperscript{328} [evidence level 3]

Converting epidural analgesia to anaesthesia for CS

There were no studies that addressed the issue of place of top-up. A survey of current UK practice is being conducted.\textsuperscript{329} Key issues in relation to the place of topping up of epidural or spinal are monitoring and safety. Two RCTs have compared different drugs to convert epidural analgesia for labour to epidural anaesthesia for CS. One RCT (n = 90) compared 3 groups. Group 1: bupivacaine 0.5% alone, group 2: bupivacaine 0.5% with lignocaine 2% and adrenalin and group 3: lignocaine 2% with adrenalin. The outcome was time to adequate block (loss of cold sensation to T4). No difference was detected between the groups but group 3 had 6 adverse events (3 high blocks and 3 patients requiring general anaesthesia).\textsuperscript{330} [evidence level 1b] Another RCT (n = 84) compared epidural conversion with or without alkalinising agents (bicarbonate v saline). Outcome assessed was time to adequate surgical block. Time to adequate block was less in the alkalinated group (mean difference 4.5 minutes).\textsuperscript{331} [evidence level 1b].

Procedures to avoid hypotension

Current practice in the UK includes the use of lateral tilt and intravenous ephedrine infusion to prevent and manage hypotension. Pre-loading and leg binders are not commonly used.\textsuperscript{312} [evidence level 3] Lateral tilt of the operating table at CS is used to decrease compression of the inferior vena cava by a gravid uterus and resultant hypotension.\textsuperscript{332} [evidence level 3] Lateral tilt is standard practice in UK units for CS.\textsuperscript{312} [evidence level 3] A systematic review that includes 3 RCTs (n = 293) has evaluated the effect of lateral tilt at CS on Apgar scores or umbilical artery pH measurements. All of the RCTs were methodologically poor with inadequate allocation concealment and poorly reported randomisation methods. All of the RCTs were conducted in the 1970s. Meta analysis was limited as different outcomes were measured. There were no differences in low Apgar scores (Peto OR 0.53, 95% CI 0.25 to 1.16) or umbilical artery pH measurements (weighted mean difference 0.03, 95% CI 0.01 to 0.04) when lateral tilt was used.\textsuperscript{333} [evidence level 1a]

We identified one RCT published after the most recent update of the review. In this RCT fetal heart rate patterns, uterine activity, umbilical artery, acid base status, newborn evaluation and maternal parameters were compared between left lateral tilt and no tilt at “emergency” CS. No difference was found when lateral tilt was used.\textsuperscript{334} [evidence level 1b]

A 15° wedge under the women’s right hip is sometimes used instead of lateral tilt at CS. Two RCTs (n = 100) considered the effect of lateral tilt versus a 15° wedge on aortocaval compression as measured by incidence of hypotension after spinal anaesthetic for CS. No difference was detected between the methods.\textsuperscript{335,336} [evidence level 1b]

A systematic review that included 20 RCTs evaluating techniques for preventing hypotension during spinal anaesthesia for CS reported that the following interventions reduce the incidence of hypotension under spinal anaesthesia for CS: pre load with crystalloid 20 ml/kg vs. control (1 RCT, n = 140; RR 0.78, 95% CI 0.6 to 1.0); pre emptive colloid vs. crystalloid (4 RCTs, n = 126; RR 0.54, 95% CI 0.37 to 0.78); ephedrine vs. control (3 RCTs, n = 146; RR 0.70, 95% CI 0.57 to 0.85); lower limb compression vs. control (5 RCTs, 181 women, RR 0.75, 95% CI 0.59 to 0.94). No difference in maternal or neonatal side effects were reported, however the RCTs were not large enough to evaluate these. A further 26 were excluded from this review for the main reason given for these exclusions was that the spinal technique was uncontrolled.\textsuperscript{337} [evidence level 1a]

Subsequent to the review two RCTs have been published that evaluate the use of elastic stockings for prevention of hypotension (n = 20).\textsuperscript{338} [evidence level 1b] and elastic stockings plus a sequential compression device (n = 100).\textsuperscript{339} [evidence level 1b] Neither RCT detected a difference in the incidence of hypotension with the use of elastic stockings alone or together with a compression device. The RCTs used different outcome measure to the RCTs included in the systematic review and therefore could not be added to the meta-analysis.

The use of bolus phenylephrine is a suggested alternative to ephedrine in maintaining maternal arterial blood pressure during regional anaesthesia. This was evaluated in an RCT (n = 38) which
reported maternal blood pressure was similar in both groups.\cite{340} [Evidence level 1b] A further RCT (n = 50) looked at the use of prophylactic epidural ephedrine to decrease the incidence of hypotension. They did not detect a difference in the incidence of hypotension between the groups.\cite{341} [Evidence level 1b]

**Procedures to manage hypotension**

Despite methods to prevent hypotension it does still occur. A systematic review of 7 RCTs (n = 292) compare the use of ephedrine to phenylephrine for the management of hypotension during spinal anaesthesia for CS. The review did not detect a difference between the two vasopressors for the management of hypotension (RR 1.00, 95% CI 0.96 to 1.06). Maternal bradycardia was more common with phenylephrine (RR 4.79, 95% CI 1.47 to 15.6) and neonates born to women given phenylephrine less likely to be acidic (RR 0.78, 95% CI 0.16 to 3.92).\cite{342} [Evidence level 1a].

A further RCT published since the review (n = 30) also compared intravenous ephedrine infusion with bolus ephedrine if hypotension developed. They reported a reduced incidence of hypotension when ephedrine infusion was used and less nausea and vomiting. There was no difference in neonatal heart rate or blood pressure.\cite{343} [Evidence level 1b] Current guidelines advise that maternity departments should have guidelines for management of hypotension.\cite{323} [Evidence level 4]

**Failed intubation**

Failed intubation remains a cause of maternal death.\cite{95} A survey of cases of failed tracheal intubation for the six year period 1993 to 1998 reports 36 cases of failed intubation in 8790 obstetric anaesthetics (incidence 1/249).\cite{344} This incidence was constant for the six year period. In the majority of cases there had been no preoperative assessment of the patient for intubation risk. There is no single test that on its own has a high predictive value for difficult intubation. Use of two or more abnormal airway findings are needed for prediction of difficult intubation and in this situation regional anaesthesia should be considered although that is no guarantee that intubation will not be needed.\cite{345} [Evidence level 4]

A number of opinion-based papers have proposed the use of laryngeal masks in cases of failed intubation with CS.\cite{346,347} [Evidence level 4] We identified a case series of 1067 women undergoing planned CS which used laryngeal masks instead of endotracheal intubation. They reported that an effective airway was obtained in 99% of women at the first attempt, 7% required intubation during the CS and there were no episodes of hypoxia, aspiration, regurgitation or laryngospasm.\cite{348} [Evidence level 3]

National anaesthetic obstetric guidelines recommend that each unit has their own drill for failed intubation such as described in recent literature.\cite{349-351} This together with predictive tools and innovative training tools such as anaesthetic emergency simulators\cite{352} should reduce mortality associated with failed intubation. [Evidence level 4]

**Use of antacids before CS**

Antacid prophylaxis forms part of routine practice at most units in the UK. NSCSA reports that 99% of UK units routinely use antacids and drugs to reduce the gastric volume and acidity for planned CS and 98% for unplanned CS. Ninety eight percent use histamine H2 receptor blockers (ranitidine or cimetidine), 2% proton pump inhibitors (omeprazole) and 99% a non-particulate antacid such as sodium citrate.\cite{353} [Evidence level 3] Ranitidine currently costs £0.64 and omeprazole £2.04 per dose to reduce acidity of gastric contents.\cite{353}

The risk of developing acid aspiration syndrome is increased when the volume aspirated into the lungs exceeds 25 ml and has an acidic pH (less than 2.5).\cite{354} [Evidence level 3] No studies have used maternal aspiration pneumonitis as an outcome measure as this is rare and would require large numbers of women to be included. Antacids are used to decrease the acidity of gastric contents. An RCT (n = 32) comparing sodium citrate with no antacid reported reduced acidity and no difference in gastric volume.\cite{355} [Evidence level 1a] A study of 20 women undergoing CS reported that women who received cimetidine preoperatively had an average pH of 5.05 compared to pH 2.97 in women who did not receive antacid. There was no difference in gastric volume measured by intraoperative aspiration of stomach contents.\cite{356} [Evidence level 2b]
An RCT (n = 595) compared ranitidine with sodium citrate to sodium citrate alone. Women who had acidic gastric contents (pH < 3.5) or a gastric volume > 25ml were defined as “at risk of aspiration”. The “risk of aspiration” was reduced in the group who had ranitidine and sodium citrate compared to sodium citrate alone (5.6% vs. 0.3%, p < 0.05) [evidence level 1b]. Another RCT (n = 541) compared omeprazole to placebo on the same “risk of aspiration” outcome. They reported a reduction in the women “at risk of aspiration” (4.3% vs. 1.4% OR 3.08 95% CI 1.02 to 9.29) [evidence level 1b]. A further 3 RCTs have compared ranitidine to omeprazole. Omeprazole results in a higher mean pH than ranitidine, however cost issues make ranitidine with sodium citrate a more cost effective option. [evidence level 1b]

Use of antiemetics

Nausea and vomiting commonly occur during CS due to aortocaval compression and resultant hypotension (see section on procedures to avoid hypotension during CS).

Routine practice in UK maternity units includes using an antacid and metoclopramide (a phenothiazine like antiemetic). [evidence level 3] An early RCT (n = 58) in women undergoing planned CS with general anaesthetic compared using metoclopramide to no treatment. The RCT did not detect a difference in gastric volume between the groups. [evidence level 1b] Later RCTs in women having CS with spinal anaesthesia show reduced incidence of nausea and vomiting in women who were given metoclopramide before induction of anaesthesia (14% vs. 81%). [evidence level 1b]

We identified five RCTs comparing different antiemetics to placebo: propofol, granisetron, droperidol and metoclopramide; ondansetron and droperidol; metoclopramide and ondansetron. Meta-analysis of these RCTs showed compared to placebo, any antiemetic reduced nausea and vomiting. [evidence level 1b] Ondansetron appears to be more effective than metoclopramide in reducing nausea (2 RCTs. RR 0.54, 95% CI 0.33 to 0.87). No difference was detected between ondansetron and droperidol in reducing nausea (2 RCTs. RR 1.0, 95% CI 0.44 to 2.27). However considering cost and safety in prescribing the cost of metoclopramide £0.28 per 10mg parenteral dose. Metoclopramide is not known to be harmful but its use should be limited to situation where there is known benefit. 5HT3 antagonists (ondansetron) is £12.89 per 8 mg parenteral dose, it is advised to avoid use during pregnancy and breastfeeding. Therefore metoclopramide should be offered if a pharmacological antiemetic is used during CS.

One RCT (n = 75) compared acupressure with placebo and metoclopramide for the prevention of nausea and vomiting during CS. Compared to placebo either acupressure or metoclopramide reduced nausea. No difference was detected between acupressure and metoclopramide (RR 1.5, 95% CI 0.5 to 4.7) [evidence level 1b]

Use of pre-oxygenation, rapid sequence induction and cricoid pressure

Standard UK practice for unplanned CS includes pre oxygenation, rapid-sequence induction and cricoid pressure for CS under general anaesthetic. We did not identify any RCT that compared use of these interventions to non use. A number of discussion papers were identified which included results of experimental work but no outcomes based studies. [evidence level 4]

<table>
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<tr>
<th>Number</th>
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<tbody>
<tr>
<td>62</td>
<td>Pregnant women having a CS should be given information on different types of post-CS analgesia so that analgesia best suited to their needs can be offered see recommendation 104). [GPP] [2004]</td>
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<tr>
<td>63</td>
<td>Women who are having a CS should be offered regional anaesthesia because it is safer and results in less maternal and neonatal morbidity than general anaesthesia. This includes women who have a diagnosis of placenta praevia. [A] [2004]</td>
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<tr>
<td>64</td>
<td>Women who are having induction of regional anaesthesia for CS should be cared for in theatre because this does not increase women’s anxiety. [B] [2004]</td>
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</table>
Women who are having a CS under regional anaesthesia should be offered intravenous ephedrine or phenylephrine, and volume pre-loading with crystalloid or colloid to reduce the risk of hypotension occurring during CS. [A] [2004]

Each maternity unit should have a drill for failed intubation during obstetric anaesthesia. [D] [2004]

To reduce the risk of aspiration pneumonitis women should be offered antacids and drugs (such as H₂ receptor antagonists or proton pump inhibitors) to reduce gastric volumes and acidity before CS. [B] [2004]

Women having a CS should be offered antiemetics (either pharmacological or acupressure) to reduce nausea and vomiting during CS. [A] [2004]

General anaesthesia for unplanned CS should include preoxygenation, cricoid pressure and rapid sequence induction to reduce the risk of aspiration. [GPP] [2004]

Intravenous ephedrine or phenylephrine should be used in the management of hypotension during CS. [A] [2004]

The operating table for CS should have a lateral tilt of 15°, because this reduces maternal hypotension. [A] [2004]

7.6 Surgical techniques for CS

A national survey of surgical techniques used during CS in the UK reports a wide range of surgical techniques being used in practice. [evidence level 3] This section presents the evidence on surgical techniques for lower segment CS in uncomplicated first procedures. Discussion of surgical techniques for specific clinical situations such as CS for preterm birth (classic uterine incision) or CS in women with previous CS (bladder adhesions) are outside the scope of this guideline.

Methods to prevent HIV transmission in theatre

Prevention of transmission of HIV from a woman undergoing CS who is known to be HIV-positive to staff carrying out the CS has been evaluated using a mathematical model and current UK HIV data, the estimated cumulative probability of occupationally acquired HIV infection is less than 1%. This is calculated at a skin puncture rate of 0.025 per procedure. However this estimate does not take into account the more common mode of contact with contaminated blood in obstetrics which is face contamination. One paper estimated the incidence of face shield contamination during CS as 50%. The incidence of cases of definite occupational acquisition of HIV in the United Kingdom has been small (1 in 319 percutaneous exposures and 1 in 3000 mucocutaneous exposures). [evidence level 3]

The use of surgical pass trays and double gloving have been tested in RCTs to determine whether their use decreases the risk of glove perforation and hence risk of infection. The use of surgical pass trays was considered in an RCT (n = 192 CS, 444 pairs of gloves) that did not detect any difference in the number of glove perforations (19% vs. 16.1% of gloves perforated, RR 1.2, 95% CI 0.8 to 1.8). [evidence level 1b]

A systematic review of wearing double gloves to reduce surgical cross infection included 18 RCTs that looked a glove perforation as an indirect measure of surgical infection. The results of the review showed that double latex gloving reduces the number of perforations to the innermost glove (OR 3.72, 95% CI 2.82, 4.91). [evidence level 1a]

In addition to the above evidence there are recommendations for safer surgical practices in general which include post exposure prophylaxis [evidence level 4]
Number | Recommendation
--- | ---
72 | Healthcare professionals should wear double gloves when performing or assisting at CS on women who have tested positive for HIV, to reduce the risk of HIV infection of healthcare professionals during surgery. [A] [2004]
73 | General recommendations for safe surgical practice should be followed at CS to reduce the risk of HIV infection of staff. [C] [2004]

### Use of adhesive drapes

We identified two RCTs on the use of adhesive drapes. Both studies addressed the impact of the use of adhesive drapes only on the incidence of postoperative wound infection. Other issues such as staff safety in the operating theatre related to spillage of blood were not addressed in these RCTs. One study described the use of adhesive drapes at CS as an isolated intervention and found the incidence of post-CS wound infection to be unchanged by their use. [378] [evidence level 1b] The other RCT described the use of adhesive drapes together with repeat disinfection of the skin before skin closure. This RCT did not find any decrease in the incidence of wound infection with the use of adhesive drapes. [380] [evidence level 1b] Neither RCT commented on the HIV status of the women that were included in the studies.

### Abdominal-wall incision

Vertical incisions for CS are uncommon in the UK (less than 1% of skin incisions are vertical) and have been replaced by transverse incisions. [306] [evidence level 3] No RCTs have compared midline to transverse incisions for CS. A meta-analysis of general surgical RCTs has compared midline, oblique and transverse incisions for their effect on postoperative pain, wound infection rates, incisional hernias and wound dehiscence. [381] Seven RCTs included postoperative pain as an outcome measure. Two RCTs (n = 209) compared midline and transverse incisions and found that the group with transverse incisions had lower pain scores and required less pethidine for analgesia (p < 0.001). Ten RCTs (n = 3586) reported on the incidence of wound infection and found no difference between the different types of incisions. Wound dehiscence and incisional hernias were reported in 9 RCTs (n = 2551) and there was no difference detected for these outcomes. [381] [evidence level 1a]

A case–control study of 48 cases of fascial dehiscence after CS described risk factors for dehiscence using stepwise logistic regression and did not find transverse incisions to have a lower risk of dehiscence than vertical incisions. [382] [evidence level 3]

An observational study (n = 89) reported on women’s perceptions of the cosmetic outcome of scar formation after either percutaneous or subcuticular sutures for CS. They found that the factor that impacted most on women’s perception of scar appearance was whether the scar was midline or transverse with transverse being more favoured. [383] [evidence level 2b]

Pfannenstiel, Maylard and Joel Cohen all described transverse abdominal wall incisions used for CS. The Pfannenstiel incision consists of a curved skin incision, two fingers breadths above the symphysis pubis, transverse incision of the sheath, rectus muscles are separated bluntly and the parietal peritoneum is incised is the midline. Maylard incision is similar but the rectus muscles are cut transversely with a knife. The Joel Cohen incision is a straight skin incision 3 cm above the pubic symphysis, then subsequent layers are opened bluntly and if necessary extended with scissors and not a knife. [384]
Four RCTs have compared different transverse incisions for CS. Two RCTs compared Pfannenstiel incision with the Joel Cohen incision. Both RCTs reported that the Joel Cohen incision is associated with shorter operating time (SMD –0.29 minutes, 95% CI –0.54 to –0.04385; SMD –0.87 minutes, 95% CI –1.28 to –0.46). Both RCTs also reported reduced postoperative febrile morbidity with the Joel Cohen incision (Pooled RR 0.35, 95% CI 0.19 to 0.64). Two RCTs compared Pfannenstiel with Maylard incisions and showed no difference in terms of operative and postoperative morbidity.

**Number** | **Recommendation**
--- | ---
74 | CS should be performed using a transverse abdominal incision because this is associated with less postoperative pain and an improved cosmetic effect compared with a midline incision. [B][2004]
75 | The transverse incision of choice should be the Joel Cohen incision (a straight skin incision, 3 cm above the symphysis pubis; subsequent tissue layers are opened bluntly and, if necessary, extended with scissors and not a knife), because it is associated with shorter operating times and reduced postoperative febrile morbidity. [A][2004]

**Instruments for skin incision**
No RCTs have addressed which instruments should be used for skin incision at CS. An RCT that included patients undergoing elective general surgical compared ‘one versus two scalpels’ technique (first scalpel for the skin and the second scalpel for deeper tissue) (n = 277). This RCT did not detect any difference in wound infection. [evidence level 1b] No other outcomes were reported. An experimental study showed that scalpels remained sterile after skin incision supporting the view that there was no need to discard the skin scalpel to prevent wound infection. [evidence level 3]

Two general surgical RCTs comparing abdominal entry using a scalpel with electrocautery did not detect any difference in any wound outcomes such as infection and strength. However the time required for the incision and incisional blood loss was less with electrocautery. [evidence level 1b]

Another RCT compared incision using a surgical knife with diathermy at cholecystectomy (n = 200). The results from this RCT showed that postoperative pain at 4, 8, 12, 16 and 24 hours and the need for morphine analgesia was less in the diathermy group. [evidence level 1b] This RCT did not assess the impact of diathermy on time to surgically open the abdomen.

**Number** | **Recommendation**
--- | ---
76 | The use of separate surgical knives to incise the skin and the deeper tissues at CS is not recommended because it does not decrease wound infection. [B][2004]

**Extension of the uterine incision**
In the UK 53% of clinicians use blunt dissection to extend the uterine incision and 47% use sharp dissection. [evidence level 3] Two RCTs have compared sharp versus blunt extension of the uterine incision at CS. [evidence level 1b]
One RCT (n = 945) reports that sharp extension is associated with greater estimated blood loss (886 ml versus 843 ml, p = 0.001); greater change in haematocrit (6.1% vs. 5.5%, p = 0.003); incidence in postpartum haemorrhage (13% vs. 9%, RR 1.23, 95% CI 1.03 to 1.46) and need for transfusion (2% vs. 0.4%, RR 1.65, 95% CI 1.25 to 2.21). The other RCT (n = 286) found no difference between sharp and blunt extension for the outcomes of unintended extension, postoperative endometritis, duration of surgery or estimated blood loss. This RCT was however underpowered to detect a difference in these outcomes. It was not possible to meta-analyse the data from these two RCTs because the outcomes are measured and reported differently in the trials.

Stapling devices can be used during incision of the uterus to decrease the blood loss from the cut edges of the uterine wall. They are not commonly used in the United Kingdom. A systematic review that included four RCTs (n = 526 women) reported no difference in the total operating time between the groups which used a stapling device and those that did not (weighted mean difference: 1.17 minutes, 95% CI –3.57 minutes to 1.22 minutes). However stapling devices increased the time to deliver the baby (weighted mean difference 0.85 minutes, 95% CI 0.48 minutes to 1.23 minutes). Blood loss was less with the use of staples (weighted mean difference 41.22 ml, 95% CI –50.63 ml to –31.8 ml). There was no difference for other perinatal outcomes. These RCTs were funded by the manufacturers of surgical staples.

### Number 77

**Recommendation**

When there is a well formed lower uterine segment, blunt rather than sharp extension of the uterine incision should be used because it reduces blood loss, incidence of postpartum haemorrhage and the need for transfusion at CS. ([A] 2004)

### Fetal laceration

The RCTs comparing sharp to blunt extension of the uterine incision do not report on incidence of trauma to the neonate, however three descriptive studies report on the incidence of fetal lacerations at CS. One study was from the UK [evidence level 3] and two of the studies were from the US (115). The UK study reports an incidence of fetal lacerations of 1.5% which is similar to the US studies (1.9% and 0.74% respectively). The UK study reported that the incidence of lacerations was independent of type of CS (unplanned or planned), fetal presentation cervical dilatation and operator grade. One US study reported that the incidence of lacerations increased to 6% with a non-cephalic presentation. ([evidence level 3])

### Number 78

**Recommendation**

Women who are having a CS should be informed that the risk of fetal lacerations is about 2%. ([C] 2004)

### Use of forceps

The use of forceps at CS has been suggested as a method of easing delivery of the fetal head, particularly for preterm infants or when the lower segment of the uterus is poorly formed. ([evidence level 3])

A small RCT (n = 44) of women undergoing planned repeat CS were randomised to vacuum, forceps or manual delivery of the fetal head. [evidence level 1b] There was no difference detected between the groups in the incidence of extension of the uterine scar, maternal blood loss or neonatal outcomes (including neonatal injuries). However women in the vacuum group reported less pain. The trial is however underpowered to evaluate these outcomes. ([evidence level 1b])
Number Recommendation

79 Forceps should only be used at CS if there is difficulty delivering the baby’s head. The effect on neonatal morbidity of the routine use of forceps at CS remains uncertain. [C] [2004]

Cord clamping

Suggested benefits of delayed cord clamping include decreased neonatal anaemia; better systemic and pulmonary perfusion; and better breastfeeding outcomes. Possible harms are polycythaemia, hyperviscosity, hyperbilirubinaemia, transient tachypnoea of the newborn and risk of maternal fetal transfusion in rhesus negative women.

One RCT based in the UK randomised women having a vaginal birth to either early or delayed cord clamping (n = 554). There was no difference detected in the duration of cord adherence, neonatal or maternal outcomes.

Two RCTs have compared the likelihood of infant anaemia between delayed and early cord clamping in preterm neonates delivered by CS. The trials use different outcome measures. One of the RCTs, from Germany (n = 40) reports that delayed cord clamping of 45 seconds results in a reduced need for packed cell transfusions during the first six weeks of life (RR 3.33, 95% CI 1.07 to 10.03). The second RCT from Australia (n = 46) found no difference in infant haematocrit between the two groups. Both RCTs found delayed cord clamping to be feasible at CS. Both RCTs were underpowered for the outcomes measured.

Number Research Recommendation

RR 33 RCTs are needed to determine the effect of delayed cord clamping on neonatal outcomes including transient tachypnoea of the newborn and risk of maternal fetal transfusion in rhesus negative women for term and preterm births.

Use of uterotonics

The licensed dose of oxytocin for CS is 5 iu by slow intravenous injection. Oxytocin is used to ensure uterine contraction, minimise delay in delivering the placenta, reduce intra operative blood loss and prevent postpartum haemorrhage. A survey of UK lead obstetric anaesthetists (n = 179) reports that 87% gave 10 units at CS, half of them administered this by rapid bolus injection. [evidence level 3] The risks of Syntocinon® (oxytocin), especially given by rapid injection, have been highlighted. Oxytocin has a direct relaxant effect on vascular smooth muscle. Under normal circumstances there is a reflex tachycardia and increased cardiac output that accompanies the transient decrease in blood pressure. The hypovolaemic woman may not respond in the normal way and in some circumstances profound hypotension may occur with resultant compromise of cardiac function.

Five RCTs have compared the use of different uterotonics at CS. Uterotonics used in these RCTs include oxytocin, oxytocin with ergometrine, misoprostol and prostaglandin F2a. No placebo controlled RCTs were identified. The use of ergometrine at uncomplicated CS is not common practice in the UK and therefore the RCT that included ergometrine is not discussed further. [evidence level 3]

One RCT (n = 40) compared oxytocin administered as an intravenous bolus of 5 iu compared with intramyometrial injection of 20 iu. This is not a licensed dose or route of administration. The intramyometrial injection was associated with more hypotension (mean decrease in systolic blood pressure one minute after oxytocin was 8.4mmHg in the intravenous group and 14.6mmHg in the intramyometrial group, p < 0.001). [evidence level 1b]
Another RCT (n = 321) compared different oxytocin infusion concentrations (20 iu/l versus 160 iu/l). The results showed that the lower concentration group had more need for additional uterotonics (39% vs. 19%, p < 0.001). There was no difference in the incidence of hypotension between the two groups.\textsuperscript{408} [evidence level 1b]

One small RCT (n = 40) compared oxytocin to misoprostol orally and found no difference between the two uterotonics.\textsuperscript{409} Misoprostol has not been found to be as effective as oxytocin for preventing postpartum haemorrhage after vaginal birth in large multicentred RCTs.\textsuperscript{410} [evidence level 1b]

Another RCT (n = 60) compared prophylactic administration of intravenous oxytocin and intramyometrial prostaglandin and detected no difference in mean estimated blood loss between the two uterotonics.\textsuperscript{411} [evidence level 1b]

Oxytocin (Syntocinon) has a short half life (4–10 minutes). Carbetocin is an oxytocin derivative which has a longer half life of 40 minutes.\textsuperscript{412} Two published RCTs (n = 694 + n = 40) have compared 100 microgrammes carbetocin with an 8-hour oxytocin infusion.\textsuperscript{413,414} The oxytocin regimen is not that recommended within this guideline. Only 1 RCT (n = 57) measured estimated blood loss and there was no difference detected between the groups.\textsuperscript{413} [evidence level 1b] The other RCT reported surrogate measures such as need for additional oxytocic.\textsuperscript{414} The RCTs were funded by the companies that produce carbetocin. Carbetocin is licensed in the UK but is yet to be launched. The basic NHS price is expected to be in the region of £12–15 per vial (information supplied by manufacturers) this compares to oxytocin which costs about £1.40 for a 5-iu or 10-iu vial.\textsuperscript{435}

Excessive haemorrhage or uterine atony can occur at CS despite the use of prophylactic uterotonics. Haemorrhage is an important cause of maternal mortality. However it is outside the scope of this guideline to address the management of obstetric haemorrhage.

### Number Recommendation

| 80 | Oxytocin 5 IU by slow intravenous injection should be used at CS to encourage contraction of the uterus and to decrease blood loss. [C][2004] |

### Method of placental removal

Nine RCTs have studied the effect of method of placental removal. Three of these are included in a systematic review.\textsuperscript{415} Eight of the RCTs considered blood loss and endometritis\textsuperscript{416,417} and one RCT only looked at fetomaternal haemorrhage\textsuperscript{418}. Feto-maternal transfusion does not appear increased by manual removal of the placenta (RR 0.37, 95% CI 0.13 to 1.07).\textsuperscript{418} [evidence level 1b]

The methods of placental removal described in each of the RCTs are manual removal of the placenta compared to controlled cord traction or spontaneous separation of the placenta. In current UK practice, the controlled cord traction is used more frequently (73%) compared to manual removal of the placenta (25%).\textsuperscript{306} [evidence level 3]

A meta-analysis of five of the RCT that reported data for endometritis was undertaken. The meta-analysis showed an increased incidence of endometritis with manual removal of the placenta compared to spontaneous separation (RR 1.54, 95% CI 1.23 to 1.92) [evidence level 1a]. The definition of endometritis was similar across the different RCTs (temperature of greater than 38°C, tender uterus, raised leucocyte count and offensive lochia). In four of the six RCTs all women received prophylactic antibiotics. In one RCT no antibiotics were given\textsuperscript{417} [evidence level 1b] and in the other RCT there was variable use of antibiotics.\textsuperscript{419} [evidence level 1b] All of these RCTs used routine administration of intra operative uterotonics.\textsuperscript{417,419–422} [evidence level 1b]

Three RCTs reported blood loss as an outcome measure.\textsuperscript{416,417,421} Meta-analysis of these RCTs showed no difference between manual removal and spontaneous separation of the placenta (SMD 0.62ml, 95% CI –1.17ml to 2.4 ml) [evidence level 1b].

Three RCTs reported on the effect of changing gloves after manual removal of the placenta and found no difference in the likelihood of post-CS endometritis (RR 1.1, 95% CI 0.75 to 1.47,\textsuperscript{423} RR of 1.0, 95% CI 0.79 to 1.3,\textsuperscript{422} and RR 1.2, 95% CI 0.5 to 2.8).\textsuperscript{419} [evidence level 1b].
Exteriorisation of the uterus

A survey of current surgical practice in the UK reports that 69% of surgeons rarely exteriorise the uterus for repair at CS, 18% ‘sometimes do so’ and 13% usually exteriorise the uterus. [evidence level 3] Four RCTs compare exteriorisation to intraperitoneal repair, two of the RCTs are included in a systematic review, the other two RCTs were published after the systematic review. All four RCTs report on blood loss and wound infection however this is measured differently across the trials (such as total units of blood transfused in each group, mean change in haematocrit per group, peri-operative change in haemoglobin and mean drop in haemoglobin between the two groups) Three RCTs detected no difference in blood loss between the groups. [evidence level 1b] The fourth RCT detected a reduction in haemoglobin drop if the uterus is exteriorised (SMD 0.2 g/dl 95% CI 0.03 g/dl to 0.51 g/dl) however there was no difference in blood transfusion rates or surgeon’s estimates of blood loss. [evidence level 1b]

Two RCTs reported on the proportion of women in each group that had blood transfusion. The meta-analysis of this outcome showed no difference in rate of blood transfusion between the two groups (RR 1.17, 95% CI 0.43 to 3.19). [evidence level 1b]

Three RCTs reported on wound infection. The meta-analysis showed no difference in wound infection between the two groups (RR 0.48 95% CI 0.18 to 1.29). [evidence level 1b]

One RCT assessed nausea, vomiting, sensation of tugging and pain scores at the end of the procedure and found no difference between the two groups. All of the women had CS under regional anaesthesia. However two women in the exteriorised group had their epidural converted to general anaesthetic due to pain. [evidence level 1b] The other RCT reported intra operative nausea, vomiting and intra operative pain and found no difference in these outcomes between the groups. Daily pain scores were measured from day 1 to day 5 postoperatively. Pain scores were higher in the exteriorisation group on day 3. A postal questionnaire was used to assess pain scores and satisfaction with the CS experience at six weeks. No difference was found in mean satisfaction scores or persistent pain. [evidence level 1b]

One- vs. two-layer closure of uterus

One-layer closure of the uterus at CS has been suggested as a means of decreasing operating time with no associated or subsequent increase in morbidity. Current practice in the UK reports that 96% of surgeons use a double layer closure and 3% a single layer. [evidence level 3]

A systematic review compares single versus two-layer suturing for closing the uterine incision at CS. [evidence level 1a] Two RCTs were included in the review (n = 1006). These RCTs measured different outcomes. One RCT (n = 906) analysed operating time and number of haemostatic sutures. [evidence level 1b] The results showed a shorter mean operating time of 5.6 minutes (43.8 versus 47.5 minutes, p = 0.0003) and fewer haemostatic sutures in the one layer closure group.

In the second RCT all the women had hysterography to determine integrity of the uterine scar 3 months after the CS in the first half of the menstrual cycle. [evidence level 1b] In the control group (two-layer closure) 82% of cases had either a major or minor scar deformity and in the intervention...
group (one layer closure) scar deformity was lower (26%). The method of randomisation in this RCT is unclear and the clinical significance of the hysterography findings as an outcome measure is uncertain.

The two RCTs have been published after the systematic review. Both assessed operating time as an outcome measure. One RCT (n = 188) found no difference in operating time\(^\text{[432]}\) [evidence level 1b] and the other RCT (n = 200) found a decrease in operating time with single layer closure of the uterus, the absolute difference was 12 minutes.\(^\text{[433]}\) [evidence level 1b]

These four RCTs used slightly different methods of single layer closure; two RCTs describing the use of continuous unlocked suture of the uterus, one RCT used continuous locked sutures while another RCT used interrupted sutures. The two later RCTs both used vicryl suture material, one of the earlier RCTs used chromic catgut and one RCT did not describe what suture material was used. None of the RCTs directly compared locked versus unlocked sutures.

Concern about the use of single layer closure of the uterus and scar rupture in future pregnancies have been raised by a cohort study (n = 2142) that reported an increase likelihood of uterine rupture in women who had had a single layer closure of the uterus (OR 3.95, 95% CI 1.35 to 11.49).\(^\text{[434]}\) [evidence level 2b] Follow up of the women recruited in one of these RCTs has also been reported.\(^\text{[435]}\) Of 164 subsequent births, 19 women had planned repeat CS and 145 experienced labour. Length of labour, mode of birth, incidence of uterine scar dehiscence and other labour outcomes were not significantly different between those women who had had previous one or two layer closure.\(^\text{[435]}\) [evidence level 2a] Closure of the uterus is currently being studied in a large UK RCT (CAESAR).\(^\text{[436]}\)

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<td>83</td>
<td>The effectiveness and safety of single layer closure of the uterine incision is uncertain. Except within a research context, the uterine incision should be sutured with two layers. [B] [2004]</td>
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**Closure of the peritoneum**

Closure of the peritoneum (visceral and parietal) has formed part of standard surgical practice and aimed to restore anatomy, reapproximate the tissues and reduce infection by forming an anatomical barrier. Current UK practice reports that 66% of surgeons do not close the parietal peritoneum while 34% do close the parietal peritoneum.\(^\text{[306]}\) [evidence level 3] A systematic review comparing non-closure with closure of the peritoneum at CS includes four RCTs (n = 1194).\(^\text{[437]}\) [evidence level 1a] Two RCTs compared closure to non-closure of both visceral and parietal peritoneum,\(^\text{[438,439]}\) one RCT compared closure to non-closure of the visceral peritoneum only\(^\text{[440]}\) and one RCT compared closure with non-closure of the parietal peritoneum only.\(^\text{[441]}\) Overall, non-closure of the peritoneum saved operating time (weighted mean difference of 6.12 minutes, 95% CI −8.00 to −4.27) with no significant differences detected in postoperative morbidity, analgesic requirements or length of hospital stay [evidence level 1a].

Since the review 7 RCTs comparing closure of both visceral and parietal peritoneum with non-closure of peritoneum have been published.\(^\text{[441-447]}\) [evidence level 1b] Four RCTs (n = 845 women) considered a wide range of morbidity measures as well as operating times.\(^\text{[442-445]}\) All consistently found operating times to be less with non-closure of the peritoneum. Three RCTs found no difference in morbidity measures between the closure and non-closure groups.\(^\text{[442,444,445]}\) One RCT suggested fewer postoperative complications.\(^\text{[443]}\) Three RCTs assess the effect on postoperative pain.\(^\text{[441,446,447]}\) [evidence level 1b] All three trials report no difference in postoperative pain (assessed using a visual analogue scoring (VAS),\(^\text{[441,446,447]}\) decreased use of analgesia after 24 hours with non-closure\(^\text{[441]}\) and increased maternal satisfaction.\(^\text{[447]}\) None of the RCTs reported long term outcomes related to healing and scarring or implications for future surgery.
Number  Recommendation

84 Neither the visceral nor the parietal peritoneum should be sutured at CS because this reduces operating time and the need for postoperative analgesia, and improves maternal satisfaction. [A] [2004]

Closure of the abdominal wall

We did not identify any RCTs that looked at closure of rectus sheath at CS. A meta-analysis (15 RCTs) has evaluated methods of abdominal-wall closure for midline incisions in general surgical patients (n = 6566). The main outcome measures were incidence of hernias, wound dehiscence, wound infection, wound pain and suture sinus formation. Incisional hernias were less common with continuous slowly absorbable sutures compared with continuous rapidly absorbable suture or non absorbable suture. Wound pain and sinus formation was more common with non absorbable sutures. [evidence level 1a]

A meta-analysis of RCTs comparing mass versus layered closure of midline incisions in general surgical patients found less incisional hernias and dehiscence to be less common with mass closures. [evidence level 1a] Midline incisions are not commonly used for CS, however there is no direct evidence on this issue so for midline incisions at CS we have extrapolated the research evidence from general surgical trials. Further research is needed on this topic for transverse abdominal incisions.

Number  Recommendation

85 In the rare circumstances that a midline abdominal incision is used at CS, mass closure with slowly absorbable continuous sutures should be used because this results in fewer incisional hernias and less dehiscence than layered closure. [B] [2004]

Number  Research Recommendation

RR 34 RCTs are required to determine the effectiveness of mass closure compared to layered closure of the abdominal wall incision at CS particularly for transverse abdominal incisions.

RR 35 Research is required to assess the effect of the various surgical techniques for CS on future surgery such as repeat CS and the incidence of complications during future surgery such as hysterectomy and urogynaecological procedures.

Closure of subcutaneous tissue

Current practice in the UK for closure of the subcutaneous layer varies between obstetricians: 42% never close it; 21% always close; 8% only close if the layer is thin; 28% close if the layer is thick. [evidence level 3]

Four RCTs have compared suturing of the subcutaneous tissue with no suturing at CS. Two RCTs randomised all women undergoing CS to suture or non-suture of the subcutaneous tissue space. One RCT found no difference in terms of wound infection or risk of wound separation. [evidence level 1b] The other RCT reported suturing to be protective against wound separation (0.36, 95% CI 0.14 to 0.91) however the method of randomisation and hence the quality of the RCT is not clear. [evidence level 1b]
Two further RCTs\(^5\) (n = 76, n = 91) randomised women with at least 2 cm subcutaneous fat.

Meta-analysis of these RCTs showed that closure of the subcutaneous space decreased the incidence of wound complications (RR 0.42, 95% CI 0.22 to 0.81). [evidence level 1a]

### Use of superficial wound drains

Five RCTs (n = 1211) have compared the routine use of superficial wound drains in CS to their selective use. [evidence level 1b] Each RCT measured slightly different parameters for the outcomes of infection and blood loss. There was no significant difference in wound infection, formation of haematoma, duration of hospital stay or need for analgesia between the groups.

One small RCT (n = 76) included women with more than 2 cm of subcutaneous fat randomised into three groups. Group 1 had suture closure of subcutaneous tissue, group 2 had a subcutaneous closed suction drain and group 3 the control group had neither. Use of a subcutaneous drain was associated with reduced incidence of wound complications compared with controls (RR 10.2, 95% CI 1.4 to 72.9) and reduced incidence of wound infection or separation (RR 7.4, 1.0 to 54.8). This is a small trial and these findings could be due to chance. [evidence level 1b]

We did not identify any evidence on the routine use of subrectus drains at CS.

### Closure of the skin

A systematic review that includes one RCT (n = 66) compares subcuticular polyglycolic suture with staples for closure of a Pfannenstiel skin incision. [evidence level 1b] They found that women with wounds closed using staples had more postoperative pain and the cosmetic effect was seen as less favourable by women. Staples took less time than subcuticular sutures (47 seconds versus 605 seconds, p < 0.001).

A nonrandomised controlled study compared percutaneous with intracutaneous (subcuticular) sutures and reported women's perceptions of the cosmetic appearance of scar formation after CS. They found that there was no difference between percutaneous and intracutaneous (subcuticular) sutures and that the factor that impacted most on women's perception of scar appearance was whether the scar was midline or transverse and the transverse scar was preferable. [evidence level 2a]

We did not identify any studies that looked at removal of staples or sutures or wound suture pain.
Obstetricians should be aware that the effects of different suture materials or methods of skin closure at CS are not certain. [C] [2004]

More RCTs are needed to determine the effect of staples compared to subcuticular sutures for skin closure at CS on postoperative pain, cosmetic appearance and removal of sutures and staples.

Umbilical artery pH measurement

Umbilical artery pH, neonatal Apgar and neonatal encephalopathy are the most reliable short term markers of poor longer term outcome such as neurodevelopment disability, cerebral palsy and perinatal death. Guidelines on electronic fetal monitoring recommend that umbilical artery pH is assessed following unplanned CS and paired umbilical artery and vein measurements are taken. [evidence level 4] This information can be used to review fetal wellbeing and to guide on-going care. It is can also be used for risk management and audit purposes.

Umbilical artery pH should be performed after all CS for suspected fetal compromise, to allow review of fetal wellbeing and guide ongoing care of the baby. [B] [2004]

Infectious complications after birth are an important cause of maternal morbidity and can prolong length of hospital stay. These include wound infection, postpartum endometritis and urinary tract infection.

Six RCTs (n = 2566) that compare planned CS to planned vaginal birth report on infection as a maternal morbidity outcome measure. The incidence of infection was 6.4% for women in the planned CS group compared with 4.9% in the planned vaginal birth group. In the largest RCT the protocol suggested prophylactic antibiotics should be used at CS. There was no information on the use of antibiotics in the other RCTs. No difference was detected in rate of infection between the two groups (pooled RR 1.29, 95% CI 0.97 to 1.72). [evidence level 1a]

Five RCTs comparing planned CS with planned vaginal birth reported on maternal puerperal pyrexia. This was defined in one of the RCTs as temperature above 38°C. Pyrexia can occur after any operative procedure and a low grade fever following a CS may not necessarily be a marker of infection. The pooled relative risk of puerperal pyrexia for women in the planned CS group was 1.96 (95% CI 1.36 to 2.84). [evidence level 1a]

Two cohort studies conducted in Israel (n = 75,947) and the USA (n = 33,251) examined the risk of infection according to mode of birth. In one study, the risk of infection was higher among women who had CS (7.9%) compared to those who had vaginal birth (1.8%) (RR 4.51 95% CI 4.00 to 5.09). [evidence level 2b] The majority of infections were endometritis; wound infection among women who had CS. In the other cohort, the incidence of postpartum endometritis among women who had CS was 2.6% compared to 0.2% among those who had vaginal births (RR 14.97, 95% CI 11.96 to 18.74). [evidence level 2b] The incidence of wound infection following CS in this study was 4.0%. The rates of wound infection were higher among women with gestational diabetes and those who had previous CS.
In the UK 85% of surgeons usually administer prophylactic antibiotics, 12% do so if other factors are present and 3% rarely use them.\textsuperscript{306} [evidence level 3]

One systematic review evaluates the use of antibiotic prophylaxis at CS on infectious complications.\textsuperscript{463} This review included 81 RCTs (n = 11,957) of which 12 RCTs included women having planned CS (n = 2037), 23 RCTs included women having unplanned CS (n = 2132), 48 RCTs included women having either planned or unplanned CS (n = 6788). In most trials antibiotic prophylaxis was administered intravenously after clamping of the umbilical cord. Overall the use of prophylactic antibiotics with CS results in a reduction in the incidence of episodes of fever (RR 0.45, 95% CI 0.39 to 0.52), endometritis (RR 0.39, 95% CI 0.34 to 0.43), wound infection (RR 0.41, 95% CI 0.35 to 0.48), urinary tract infection (RR 0.54, 95% CI 0.46 to 0.64) and serious infection (RR 0.42, CI 0.28 to 0.65). [evidence level 1a]

Maternal side effects were not consistently collected across the RCTs. There were 3 possible episodes in the placebo group and 16 in the antibiotic group, such as phlebitis or rash at the intravenous infusion site. No serious drug reactions were reported. The effect on breast feeding and thrush in newborns being breastfed was not reported in any of the RCTs included in the systematic review.

Another systematic review\textsuperscript{464} investigated the effectiveness of different antibiotic regimens. Fifty one RCTs were included. There is no advantage in using a multiple dose regimen compared with a single dose (OR 0.92, 95% CI 0.70 to 1.23). There was no difference in the efficacy of ampicillin compared with first generation cephalosporins (OR 1.27, 95% CI 0.84 to 1.93), nor was there any difference between first generation compared with second or third generation cephalosporins (OR 1.21, 95% CI 0.97 to 1.51). [evidence level 1a]

Other methods to reduce infectious morbidity at CS have been investigated including RCTs on the use of intra abdominal lavage with saline,\textsuperscript{465} intrauterine lavage with antibiotics,\textsuperscript{466} preoperative skin preparation\textsuperscript{467} and vaginal preparation with povidone iodine\textsuperscript{468} none of which showed a difference in infectious morbidity [evidence level 1b]. Pelvic irrigation with antibiotic solution\textsuperscript{467} and the use of intravaginal metronidazole\textsuperscript{469} did show some difference in infectious morbidity but the numbers were small. [evidence level 1b] We did not find any RCTs looking at the postoperative prophylactic use of antibiotics after CS.

**Economic considerations for the use prophylactic antibiotics at CS**

Where two antibiotics have the same efficacy, the less expensive antibiotic should be offered since there is no justification for the use of more expensive regimens. There is some economic evidence that a single dose of antibiotic is as effective as two- and three-dose regimens\textsuperscript{470} and since the efficacy is the same, the lower cost regimen should be offered.

An economic evaluation study undertaken in the United Kingdom in the late 1980s suggested that there might be significant savings from the use of prophylactic antibiotics.\textsuperscript{471} This evaluation was based on a model that used post-CS wound infection rates of 8.4% and 50–70% reduction in odds of wound infection with the use of prophylactic antibiotics. Using these assumptions in an economic model, the estimated additional average cost of hospital postnatal care for women with wound infection (compared with women who had had CS and no wound infection) was £716. Introducing routine prophylaxis with antibiotics would reduce average costs of postnatal care by between £1,300 and £3,900 per 100 CS, depending on the cost of the antibiotic used and its effectiveness. This analysis supports the use of prophylactic antibiotics after CS since this strategy dominates a no antibiotic strategy (due to lower cost, greater effectiveness).

A cost-effectiveness analysis of the cost per post-CS infection averted has not been undertaken in a United Kingdom setting.

**Timing of antibiotic administration**

**Introduction**

Antibiotic prophylaxis at the time of a CS is proven to reduce post-operative maternal postnatal infective morbidity rates. Traditionally, the antibiotics are not administered until after the umbilical cord is clamped so that the unborn baby is not unduly exposed to antibiotics administered to the mother,
with potential adverse effects. This contrasts with the general timing of antibiotic administration for surgical site infection prophylaxis (see below). This section examines whether administration of antibiotics before cord clamping is associated with lower maternal infective morbidity compared to administration post-clamping without imposing known additional risks to the neonate.

**Review question**
What is the effectiveness of antibiotics given prior to clamping of the cord compared to antibiotics given after clamping of the cord during a planned or unplanned caesarean section?

**Existing NICE guidance**
The Surgical site infection guideline (NICE, 2008) recommends that:

- antibiotic prophylaxis should be given in clean-contaminated surgery (such as caesarean section)
- giving a single dose of antibiotic prophylaxis intravenously on starting anaesthesia should be considered or earlier for operations in which a tourniquet is used.
- before giving antibiotic prophylaxis the timing, pharmacokinetics (for example, the serum half-life) and necessary infusion time of the antibiotic should be considered. A repeat dose of antibiotic prophylaxis should be given when the operation is longer than the half-life of the antibiotic.
- patients should be informed before the operation, whenever possible, if they will need antibiotic prophylaxis, and afterwards if they have been given antibiotics during their operation.

**Overview of evidence**
Six reports of RCTs were included in this review (Gordon et al., 1979; Sullivan, 2007; Thigpen et al., 2005; Wax et al., 1997; Nokiani, 2009; Yildirim, 2009).

Four studies were conducted in the USA (Gordon et al., 1979; Sullivan, 2007; Thigpen et al., 2005; Wax et al., 1997), one in Iran (Nokiani, 2009) and one in Turkey (Yildirim, 2009). Two of the American studies reported their participants to be high risk for subsequent infection (Sullivan, 2007; Thigpen et al., 2005). In total, 1503 women participated in these studies, with 805 receiving antibiotics prior to cord clamping and 698 receiving antibiotics after the cord was clamped. Two studies (Thigpen et al., 2005; Wax et al., 1997) intentionally included women who were in labour and had unplanned CS, three studies included planned caesarean cases (Sullivan, 2007; Yildirim, 2009; Gordon et al., 1979) and the remaining study (Nokiani, 2009) included mostly women undergoing planned CS, although significantly more women in the post-clamping group had an intrapartum CS after the onset of labour, despite the investigators’ efforts to recruit only participants undergoing planned CS.

One study examined the timing of a 1g dose of intravenous [IV] ampicillin (Gordon et al., 1979) whilst the remaining studies examined the timing of cefazolin administered in a 1g IV dose (Sullivan, 2007; Yildirim, 2009; Wax et al., 1997) or 2g IV dose (Thigpen et al., 2005; Nokiani, 2009).

Six maternal and three neonatal outcomes were chosen by the GDG as being of priority to inform recommendations and the results for these are presented in the table below.

**Table 7.3 GRADE findings comparing pre-clamp vs. post-clamp administration of antibiotics**

<table>
<thead>
<tr>
<th>Number of studies</th>
<th>Number of women/babies (%) or number of hours</th>
<th>Effect</th>
<th>Quality</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Pre cord-clamp antibiotics</td>
<td>Post cord-clamp antibiotics during CS</td>
<td>Relative (95% CI)</td>
</tr>
<tr>
<td>Overall/total maternal infectious morbidity</td>
<td>5 studies</td>
<td>55/609 (9%)</td>
<td>84/607 (13.8%)</td>
</tr>
</tbody>
</table>

Caesarean section: full guideline DRAFT (September 2011)  
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### Maternal wound infection

<table>
<thead>
<tr>
<th>Study Details</th>
<th>Cases (RR)</th>
<th>Controls (RR)</th>
<th>RR</th>
<th>Pooled RR</th>
<th>Summary</th>
</tr>
</thead>
<tbody>
<tr>
<td>5 studies (Gordon et al., 1979; Sullivan, 2007; Thigpen et al., 2005; Wax et al., 1997; Yildirim, 2009)</td>
<td>18/609 (3%)</td>
<td>29/607 (4.8%)</td>
<td>0.63 (0.35 to 1.11)</td>
<td>18 fewer per 1000 (from 31 fewer to 5 more)</td>
<td>Moderate</td>
</tr>
</tbody>
</table>

### Surgical site opening

<table>
<thead>
<tr>
<th>Study Details</th>
<th>Cases (RR)</th>
<th>Controls (RR)</th>
<th>RR</th>
<th>Pooled RR</th>
<th>Summary</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 study (Nokiani, 2009)</td>
<td>0/196 (0%)</td>
<td>1/91 (1.1%)</td>
<td>0.16 (0.01 to 3.78)</td>
<td>9 fewer per 1000 (from 11 fewer to 31 more)</td>
<td>Low</td>
</tr>
</tbody>
</table>

### Total maternal fever

<table>
<thead>
<tr>
<th>Study Details</th>
<th>Cases (RR)</th>
<th>Controls (RR)</th>
<th>RR</th>
<th>Pooled RR</th>
<th>Summary</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 study (Nokiani, 2009)</td>
<td>10/196 (5.1%)</td>
<td>3/91 (3.3%)</td>
<td>1.55 (0.44 to 5.49)</td>
<td>18 more per 1000 (from 18 fewer to 148 more)</td>
<td>Low</td>
</tr>
</tbody>
</table>

### Maternal urinary tract infection [UTI]

<table>
<thead>
<tr>
<th>Study Details</th>
<th>Cases (RR)</th>
<th>Controls (RR)</th>
<th>RR</th>
<th>Pooled RR</th>
<th>Summary</th>
</tr>
</thead>
<tbody>
<tr>
<td>3 studies (Gordon et al., 1979; Wax et al., 1997; Yildirim, 2009)</td>
<td>3/281 (1.1%)</td>
<td>6/276 (2.2%)</td>
<td>0.55 (0.15 to 1.98)</td>
<td>10 fewer per 1000 (from 18 fewer to 21 more)</td>
<td>Moderate</td>
</tr>
</tbody>
</table>

### Endometritis or endomyometritis

<table>
<thead>
<tr>
<th>Study Details</th>
<th>Cases (RR)</th>
<th>Controls (RR)</th>
<th>RR</th>
<th>Pooled RR</th>
<th>Summary</th>
</tr>
</thead>
<tbody>
<tr>
<td>5 studies (Gordon et al., 1979; Sullivan, 2007; Thigpen et al., 2005; Wax et al., 1997; Yildirim, 2009)</td>
<td>24/609 (3.9%)</td>
<td>42/607 (6.9%)</td>
<td>0.57 (0.35 to 0.92)</td>
<td>30 fewer per 1000 (from 6 fewer to 45 fewer)</td>
<td>High</td>
</tr>
</tbody>
</table>

### Endometritis

<table>
<thead>
<tr>
<th>Study Details</th>
<th>Cases (RR)</th>
<th>Controls (RR)</th>
<th>Summary</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 study (Nokiani, 2009)</td>
<td>0/196 (0%)</td>
<td>0/91 (0%)</td>
<td>not pooled</td>
</tr>
</tbody>
</table>

### Maternal pneumonia or respiratory tract infection [RTI]

<table>
<thead>
<tr>
<th>Study Details</th>
<th>Cases (RR)</th>
<th>Controls (RR)</th>
<th>Summary</th>
</tr>
</thead>
<tbody>
<tr>
<td>2 studies</td>
<td>0/243</td>
<td>0/236</td>
<td>not pooled</td>
</tr>
</tbody>
</table>
### Neonatal sepsis or infection

<table>
<thead>
<tr>
<th>Study Details</th>
<th>Neonatal sepsis or infection</th>
<th>Sepsis or infection</th>
<th>Mean</th>
<th>Sepsis or infection</th>
<th>Mean</th>
<th>Sepsis or infection</th>
<th>Mean</th>
<th>Sepsis or infection</th>
<th>Mean</th>
<th>Sepsis or infection</th>
<th>Mean</th>
</tr>
</thead>
<tbody>
<tr>
<td>4 studies (Sullivan, 2007, Thigpen et al., 2005, Wax et al., 1997; Yildirim, 2009)</td>
<td>37/588 (6.3%)</td>
<td>41/582 (7%)</td>
<td>RR 0.89 (0.58 to 1.35)</td>
<td>8 fewer per 1000 (from 30 fewer to 25 more)</td>
<td>Moderate</td>
<td></td>
<td></td>
<td></td>
<td></td>
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<td></td>
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</tbody>
</table>

### Neonatal sepsis

<table>
<thead>
<tr>
<th>Study Details</th>
<th>Sepsis or infection</th>
<th>Mean</th>
<th>Sepsis or infection</th>
<th>Mean</th>
<th>Sepsis or infection</th>
<th>Mean</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 study (Nokiani, 2009)</td>
<td>4/196 (2%)</td>
<td>1/91 (1.1%)</td>
<td>RR 1.86 (0.21 to 16.38)</td>
<td>9 more per 1000 (from 9 fewer to 169 more)</td>
<td>Low</td>
<td></td>
</tr>
</tbody>
</table>

### Mean neonatal length of stay

<table>
<thead>
<tr>
<th>Study Details</th>
<th>Mean</th>
<th>Sepsis or infection</th>
<th>Mean</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 study (Sullivan, 2007)</td>
<td>6.6 ± 9.9 (n=185)</td>
<td>8.5 ± 15.8 (n=194)</td>
<td>NC</td>
</tr>
<tr>
<td></td>
<td>MD 1.9 hours shorter (4.54 shorter to 0.74 longer)</td>
<td>Moderate</td>
<td></td>
</tr>
</tbody>
</table>

### Mean neonatal length of stay

<table>
<thead>
<tr>
<th>Study Details</th>
<th>Mean</th>
<th>Sepsis or infection</th>
<th>Mean</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 study (Nokiani, 2009)</td>
<td>2.99 ± 0.07 (n=196)</td>
<td>2.99 ± 0.11 (n=191)</td>
<td>NC</td>
</tr>
<tr>
<td></td>
<td>MD 0.0 hours shorter (0.02 shorter to 0.02 longer)</td>
<td>Low</td>
<td></td>
</tr>
</tbody>
</table>

### Mean neonatal intensive care unit [NICU] length of stay

<table>
<thead>
<tr>
<th>Study Details</th>
<th>Mean</th>
<th>Sepsis or infection</th>
<th>Mean</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 study (Sullivan, 2007)</td>
<td>14.2 ± 15.8 (n=185)</td>
<td>19.7 ± 24.9 (n=194)</td>
<td>NC</td>
</tr>
<tr>
<td></td>
<td>MD 5.50 shorter (9.68 shorter to -1.32 shorter)</td>
<td>Moderate</td>
<td></td>
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</table>

### Mean NICU length of stay

<table>
<thead>
<tr>
<th>Study Details</th>
<th>Mean</th>
<th>Sepsis or infection</th>
<th>Mean</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 study (Yildirim, 2009)</td>
<td>8.25 ± 2.62 (n=201)</td>
<td>5.66 ± 2.58 (n=198)</td>
<td>NC</td>
</tr>
<tr>
<td></td>
<td>MD 2.59 longer (2.08 longer to 3.10 longer)</td>
<td>Moderate</td>
<td></td>
</tr>
</tbody>
</table>

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### Evidence statements

#### Maternal outcomes

**Overall/total maternal infectious morbidity**

A meta-analysis of five RCTs found that antibiotics given before cord-clamping reduces the incidence of total maternal infectious morbidity compared to antibiotics given after cord-clamping. This finding was statistically significant. The evidence for this outcome was of high quality.

**Maternal wound infection**

A meta-analysis of five RCTs did not find a statistically significant difference in the rate of maternal wound infection when antibiotics were given before cord-clamping when compared with occasions when the antibiotics were given after cord-clamping. The evidence for this outcome was of moderate quality.
Surgical site opening

One study (with a serious design limitation) did not find a statistically significant difference in incidence rates of surgical site opening when antibiotics were given before cord-clamping when compared with occasions when the antibiotics were given after cord-clamping, although only one observation in 281 women undergoing CS was reported. The evidence for this outcome was of low quality.

Total maternal fever

One RCT did not find a statistically significant difference in the incidence of total maternal fever according to timing of antibiotic administration. The evidence for this outcome was of low quality.

Maternal urinary tract infection

A meta-analysis of three RCTs did not find a statistically significant difference in the incidence of maternal urinary tract infection according to the timing of antibiotic administration. The evidence for this outcome was of moderate quality.

Endometritis or endomyometritis

A meta-analysis of five RCTs found that administration of antibiotics before cord-clamping reduces the incidence of endometritis or endomyometritis compared to antibiotics given after cord-clamping. This finding was statistically significant. The evidence for this outcome was of high quality. One study (with a serious design limitation) found no incidences of endometritis in 281 women undergoing CS. This finding was not statistically significant. The evidence for this outcome was of low quality.

Maternal pneumonia or respiratory tract infection

Two RCTs examined the effects of timing of antibiotic prophylaxis on the incidence of maternal pneumonia or RTI. No events were seen in either study (n=479). The evidence for this outcome was of low quality.

Neonatal outcomes

Neonatal sepsis or infection

A meta-analysis of four RCTs did not find a statistically significant difference in the rate of neonatal infection or sepsis when antibiotics were given before cord-clamping compared with occasions when antibiotics were given after cord-clamping. The evidence for this outcome was of moderate quality. One study (with a serious design limitation) did not find a statistically significant difference in the rate of neonatal sepsis when antibiotics were given before cord-clamping compared with occasions when antibiotics were given after cord-clamping. The evidence for this outcome was of low quality.

Neonatal length of stay

Two RCTs did not find a statistically significant difference in the mean neonatal length of hospital stay when antibiotics were given before cord-clamping compared with occasions when antibiotics were given after cord-clamping. The evidence for this outcome was of moderate quality in the first study and low quality in the second study.

Neonatal intensive care unit (NICU) length of stay

One RCT found that administration of antibiotics after cord-clamping reduced NICU length of stay compared to antibiotics administered prior to cord-clamping. This finding was statistically significant. One RCT found that administration of antibiotics before cord-clamping reduced NICU length of stay compared to antibiotics administered after cord-clamping. This finding was statistically significant. The evidence for this outcome was of moderate quality.

Evidence to recommendations

Relative value placed on the outcomes considered

In developing the recommendations, it was necessary to consider the outcomes for both the woman and her baby. The group agreed that the most relevant outcomes to consider were measures such as sepsis which could be determined objectively. Antibiotics are given at CS to prevent maternal infection associated with surgery. Infection is reported in the literature in different ways e.g. different types of infection may be reported in combination as an overall infection rate or different infections...
may be reported individually. It is also the case that infection is reported using different definitions across studies e.g. wound infection vs. surgical site opening. The GDG chose overall infectious morbidity as the most useful outcome to inform recommendations. Results were compiled into an overall rate where this was possible. The GDG acknowledged that different infections would contribute to this overall score in each study. The GDG also thought it was important to consider the rates of particular types of maternal infection and identified 5 maternal outcomes relating to these (see evidence profile above).

The GDG considered confirmed estimates of neonatal infection to be the most informative neonatal outcome. The proxy estimate of neonatal length of stay in intensive care was also used although it was acknowledged that this outcome is less useful as it is also affected by other factors such as hospital policy.

**Trade-off between clinical benefit and harms**

The GDG agreed that prevention of maternal infection was the most important outcome when considering timing of antibiotic administration but that it was important to take into consideration potential harmful effects on the neonate. The group agreed that the strongest and clearest evidence showed that there was a reduction in maternal morbidity when antibiotics were administered prior to an incision being made. Furthermore, the group felt from their clinical experience that the benefits of administering antibiotics prior to incision would be even more pronounced in women with a longer incision-to-delivery time (such as obese women) as their chance of being exposed to infection is greater.

The evidence for neonates was more equivocal and the group did not feel able to discern an effect of the timing of antibiotics on benefits or harms for this group. The evidence showed no difference in rates of neonatal infection between the early administration and later administration groups. In addition, the GDG recognised that whilst administering antibiotics prior to incision would expose the baby to the antibiotic, in breast-fed babies antibiotics would be passed to the baby anyway, regardless of the timing of their administration. Whilst 2 RCTs showed a reduction in neonatal intensive care length of stay for the delayed administration group, this difference was not apparent in 2 other trials and the GDG felt the effects of confounding variables on this outcome meant it was less valid as an indicator of neonatal wellbeing compared with overall infection rates.

**Trade-off between net health-benefits and resource use**

Although no formal cost effectiveness modelling was carried out for this question, the group noted that with relatively strong evidence for a reduction in maternal infections with pre-incision administration of antibiotics, and no clear evidence either way for an effect on neonatal outcomes, it was likely that pre-incision administration of antibiotics would be cost effective compared with administration after cord clamping.

**Quality of evidence**

The evidence considered in this review was of mixed quality. The group noted a particular problem with one study which included twice as many women with an unplanned CS in one arm of the study than the other (Nokiani F.A. 2009). They felt that this was likely to impact on the results and so did not feel that it was appropriate to use the study in developing recommendations.

The group also noted that in one study which reported on mean length of NICU stay, the antibiotics were infused for 45 minutes prior to incision. As this practice would only be possible with planned caesareans, the group did not feel that these results were generalisable.

Overall, the group felt that the evidence which showed a clear difference in outcomes (overall maternal infectious morbidity and rates of endometritis or endomyometritis) was of a good quality.

**Other considerations**

All of the evidence reviewed for this section related to cefazolin (5 studies) and ampicillin (1 study). The group was aware that despite the recommendations from the original guideline, these antibiotics are not commonly used in the UK. The group was also aware of findings from the ORACLE studies which looked at the use of antibiotics both in women with pre-labour rupture of membranes and in women thought to be in preterm labour. The studies found an increased risk of necrotising enterocolitis in babies if they were exposed in utero to co-amoxiclav. This risk did not vary between babies born to the groups of women and nor did it vary whether birth was whilst still
being exposed to co-amoxiclav or whether the exposure had ceased. Given this increased risk, and given the neonate’s chance of exposure to the antibiotic when giving prophylaxis before skin incision or cord clamping, the group agreed that the use co-amoxiclav should be discouraged, particularly as there are a number of acceptable alternative antibiotics available.

<table>
<thead>
<tr>
<th>Number</th>
<th>Recommendation</th>
</tr>
</thead>
<tbody>
<tr>
<td>90</td>
<td>Offer women prophylactic antibiotics at CS before skin incision. Inform them that this reduces the risk of maternal infection more than prophylactic antibiotics given after skin incision, and that no effect on the baby has been demonstrated. [new 2011]</td>
</tr>
<tr>
<td>91</td>
<td>Women having a CS should be offered prophylactic antibiotics, such as a single dose of first-generation cephalosporin or ampicillin, to reduce the risk of postoperative infections (such as endometritis, urinary tract and wound infection), which occur in about 8% of women who have had a CS. [A] [2004]</td>
</tr>
<tr>
<td>92</td>
<td>Do not use co-amoxiclav when giving antibiotics before skin incision. [new 2011]</td>
</tr>
</tbody>
</table>

### Thromboprophylaxis for CS

Pregnancy is a risk factor for thromboembolic disease. The reported incidence of pulmonary thromboembolism is 6 per 10,000 maternities, this varies according to risk factors such as maternal age, obesity, smoking. Pulmonary embolism is the leading direct cause of maternal death in the UK (estimate mortality rate of 1.45 per 100,000 maternities). Thromboembolic disease is rare and is reported as an outcome measure in only one RCT of planned CS compared with planned vaginal birth, however within this trial there were no events in either group.

A population-based cohort study evaluated the risk of thromboembolism by mode of birth (n = 1,003,489) (1987–1995). The risk of pulmonary embolism was increased for women who had CS compared with those who had vaginal birth (unadjusted RR 3.8 95% CI 2.0 to 4.9). Within this cohort it is not known how many women in this study would have received thromboprophylaxis.

A systematic review of thromboprophylaxis during pregnancy and the early postnatal period was identified. The review included eight RCTs (n = 649) of which only four studies address the issue of thromboprophylaxis for CS (n = 350). The interventions evaluated in these trials include hydroxyethyl starch, heparin and placebo. Thromboembolic events are relatively rare so that although no differences were detected between the intervention and control groups this is probably because the trials are too small to evaluate these outcomes. There is a large RCT of thromboprophylaxis after CS in progress.

Currently available publications to guide practice on this issue recommend thromboprophylaxis for CS based on assessment of risk (such as unplanned versus planned CS, maternal age over 35 years, weight greater than 80 kg, medical complication). Recommended thromboprophylaxis includes hydration, early mobilisation, graduated elastic compression stockings and low-molecular-weight heparin. [evidence level 4] Data from the NSCSA shows that in current practice, thromboprophylaxis is used in 89% of unplanned CS and 87% of planned CS.

<table>
<thead>
<tr>
<th>Number</th>
<th>Recommendation</th>
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<tbody>
<tr>
<td>93</td>
<td>Women having a CS should be offered thromboprophylaxis because they are at increased risk of venous thromboembolism. The choice of method of prophylaxis (for example, graduated stockings, hydration, early mobilisation, low molecular weight</td>
</tr>
</tbody>
</table>
Caesarean section: full guideline DRAFT (September 2011)

<table>
<thead>
<tr>
<th>Number</th>
<th>Research Recommendation</th>
</tr>
</thead>
<tbody>
<tr>
<td>RR 38</td>
<td>What is the most effective antibiotic to prevent maternal infectious morbidity post-CS when given prior to incision</td>
</tr>
<tr>
<td>RR 39</td>
<td>What is the physical, psychological and social impact of maternal infectious morbidity post-CS?</td>
</tr>
</tbody>
</table>

### Need for further surgery (including hysterectomy)

Surgery immediately following birth can include manual removal of placenta, uterine curettage, and laparotomy (with or without hysterectomy). In the UK, the reported rate of peripartum hysterectomy is 6–7 per 10,000 deliveries. In other well resourced countries the incidence (excluding elective hysterectomy) range from 4–15 per 10,000. These rates vary according to parity, number of previous CS and other conditions e.g. placenta praevia. In one UK survey about 2% of women required further surgery. The need for dilatation and curettage was reported in one RCT (n = 2082) that compared planned CS with planned vaginal birth. Dilatation and curettage was reduced in the planned CS group (0.3%) compared to the planned vaginal birth group (0.4% RR 0.75, 95% CI 0.17 to 3.34). [evidence level 1b]. Hysterectomy was reported in two RCTs. In one RCT there were no events in either group. In the other RCT (n = 208), 1.1% of women in the planned CS group and none of the women in the planned vaginal birth group were reported to have this outcome.

One Australian cohort study (n = 29,488) evaluated need for further surgery following childbirth. The return to theatre rate for women who had a CS was 0.5% compared to 0.03% of women who had vaginal birth (unadjusted RR 17.53, 95% CI 9.37 to 32.1). [evidence level 2a] The main reason for further surgery in both groups was severe obstetric haemorrhage. 80% of women who had further surgery for haemorrhage following CS required a laparotomy compared to 27% of women who required surgery after vaginal birth for severe haemorrhage. The majority (73%) of women who had a vaginal birth with severe haemorrhage requiring surgery had uterine curettage.

Two cohort studies conducted in the USA have compared rates of hysterectomy for women according to mode of birth. The rate of peripartum hysterectomy was higher among women who had CS (0.7 to 0.8%) compared with 0.01 to 0.02% among women who had vaginal birth (unadjusted RR 95.5, 95% CI 67.7 to 136.9, unadjusted RR 43.97 95% CI 22.52 to 85.85). The RR adjusted for placenta praevia was reported to be 10.8 (95% CI 7.8 to 15.4). [evidence level 2a] In one of these studies, 19% cases of peripartum hysterectomy were in women who were in their first pregnancy. Data on rates of peripartum hysterectomy following primary CS were not reported in either of these studies.

### Maternal satisfaction during CS

A number of practices have been suggested to improve women’s satisfaction with CS birth. These include seeing the baby born via a lowered screen; music playing in theatre; silence at moment of birth in theatre so the mother’s voice is the first the baby hears and lowering the lights at the moment of birth. We did not identify any RCT that evaluated the effectiveness of these changes in practice. Although no papers discuss the use of music during CS one experimental study (n = 65) describes the use of medical resonant music therapy as preoperative preparation for CS compared with women who received sedatives. The experimental group receiving music therapy had lower cortisol levels and noted better sleep and less need for analgesics postoperatively. [evidence level 2b]

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11 For more information see ‘Venous thromboembolism: reducing the risk’ (NICE clinical guideline 92).
Case reports\textsuperscript{488} [evidence level 3] and case series\textsuperscript{489} [evidence level 3] report positive maternal attitudes towards music during labour in terms of pain relief and satisfaction. A non-systematic review of literature on the efficacy of music therapy for premature infants suggest that music is associated with reduced length of hospital stay, improved weight gain and oxygen saturation level.\textsuperscript{490} [evidence level 3]

A number of studies relate to hearing ‘mother’s voice’ were identified. One (n = 10 babies) experiment showed that neonates were ‘more likely to work’ to produce their mother’s voice than other female voices\textsuperscript{491} [evidence level 3] and another experimental study (40 neonates) found that neonates responded more to their mother’s voice than other female voices even when there was no postnatal experience of the mother’s voice.\textsuperscript{492} [evidence level 3]

No other published evidence was found on other changes in practice to improve woman’s satisfaction of CS birth. Personal communication from consumer groups suggest that this is an area that warrants further research due to woman’s perceptions of the benefit of these practices.\textsuperscript{493}

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<th>Number</th>
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<tbody>
<tr>
<td>94</td>
<td>Women’s preferences for the birth, such as music playing in theatre, lowering the screen to see the baby born, or silence so that the mother’s voice is the first the baby hears, should be accommodated where possible. [GPP] [2004]</td>
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<th>Number</th>
<th>Research Recommendation</th>
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<tbody>
<tr>
<td>RR 40</td>
<td>More evaluation of interventions such as seeing baby born via a lowered screen; music playing in theatre; silence in theatre so mother’s voice is the first baby hears and lowering the lights in theatre during CS are needed.</td>
</tr>
</tbody>
</table>

Caesarean section: full guideline DRAFT (September 2011)
8 Care of the baby born by CS

The perinatal mortality rate in England and Wales is 7.9 per 1000 total births.\(^7\) The effect of CS on baby outcome is not a simple reciprocal relationship.\(^{494,495}\) Perinatal mortality rate can decline in the presence of a low and stable CS rate or remain stable while the CS rate increases.\(^{494,496}\) [evidence level 4] A cohort study (n = 11,702) reported neonatal mortality. No difference was detected in neonatal mortality between vaginal birth and CS however the study is underpowered to evaluate this outcome (RR 1.09, 95% CI 0.14 to 8.38).\(^{497}\) [evidence level 2b]

8.1 Presence of paediatrician at CS

One cohort study reported of infants delivered by CS (using regional anaesthesia) were more likely to have a 1 minute Apgar of less than 4, (6.3%) compared with infants delivered vaginally (1.3%. RR 3.04 95% CI 1.80 to 5.13).\(^{497}\) [evidence level 2b] Two descriptive studies list CS as one of the situations that require a paediatrician to be present at birth.\(^{498,499}\) [evidence level 3] A series of 460 deliveries showed that there was higher incidence of neonatal resuscitation with planned CS deliveries compared to vaginal births. Similar results were found in two other studies as well.\(^{500,501}\) Of the 59 “emergency” CS, 24 were for fetal distress of which 12 needed resuscitation. There is no difference in the need for resuscitation between babies with cephalic presentation born by CS (1.8%) and vaginal birth (2.7%) with no evidence of fetal distress.\(^{502}\) [evidence level 3]

A large observational study from the USA (n = 3940) reported that infants born by CS with general anaesthesia are at an increased risk of having 1- and 5-minute Apgar scores of less than 7 when compared with those born by CS with regional anaesthesia (1-minute Apgar less than 7 RR 3.13, 95% CI 2.5 to 3.88, 5-minute Apgar RR 3.6, 95% CI 1.81 to 7.00) and the need for resuscitation (RR 2.02, 95% CI 1.39 to 2.9)\(^{517}\) [evidence level 3]. These findings are consistent with those in the NSCSA.\(^{290}\)

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<th>Number</th>
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<tr>
<td>95</td>
<td>An appropriately trained practitioner skilled in the resuscitation of the newborn should be present at CS performed under general anaesthesia or where there is evidence of fetal compromise. [C] [2004]</td>
</tr>
</tbody>
</table>

8.2 Neonatal encephalopathy and cerebral palsy

There are a number of causes of cerebral palsy and probably only about 10% are related to intrapartum events.\(^{503}\) The majority of neurological pathologies causing cerebral palsy occur as a result of multi factorial and mostly unpreventable reasons during either fetal development or the neonatal period.\(^{504,503}\) It is therefore not surprising that ecological studies do not show an association between high CS rates and low cerebral palsy rates.\(^{505}\) [evidence level 3] The impact of CS on cerebral palsy was assessed in a systematic review. The review identified 10 studies none of which found a difference in the rates of cerebral palsy, abnormal neurological development between children born by CS or vaginal birth. The studies were in groups at “high risk” of these outcomes (such as preterm birth, breech).\(^{505}\) [evidence level 3]
Another cohort study considers the effect of CS on severe neurological morbidity including cerebral palsy.\textsuperscript{100} There was an increased risk of severe neurological morbidity in those delivered by CS (unadjusted RR 1.81, 95% CI 1.56 to 2.11). \textsuperscript{[evidence level 2b]} A case–control study compared 164 babies with neonatal encephalopathy compared with 400 babies that did not have neonatal encephalopathy (controls). Babies that had neonatal encephalopathy were more likely to have had instrumental vaginal delivery (OR 2.34, 95% CI 1.16 to 4.70), “emergency” CS (OR 2.17, 95% CI 1.01 to 4.64) and less likely to have had “elective” CS (OR 0.17, 95% CI 0.05 to 0.56).\textsuperscript{506} \textsuperscript{[evidence level 3]}

### Number Research Recommendation

| RR 41 | Further evaluation of the long and short term risks and benefits of CS compared with vaginal birth for babies is required. |

#### 8.3 Birth injuries

The benefits of CS for specific groups such as term breech, or preterm birth are discussed in Chapter 5. The evidence on the comparative risk of birth injuries in term singleton cephalic infants is limited to one large audit of birth records looking at mode of birth and intracranial injury\textsuperscript{507} and one case–control study looking at brachial plexus injuries.\textsuperscript{508} In the audit,\textsuperscript{583,340} live born singleton infants born to nulliparous women, weighing between 2500 g and 4000 g over a two year period were studied. Breech presentations were excluded. Neonates were grouped according to mode of birth. The incidence of intracranial haemorrhages was 0.01% in the ‘CS during labour’ group compared to 0.05% in the ‘spontaneous’ vaginal birth group (OR 2.1, 95% CI 1.6 to 2.7). It was 0.04% in the ‘CS before labour’ group (OR 0.7, 95% CI 0.4 to 1.3).\textsuperscript{507} \textsuperscript{[evidence level 3]}

The case–control study compared all modes of birth including assisted vaginal deliveries\textsuperscript{508} for risk of brachial plexus injury in 106 cases of Erb’s palsy and 382 controls. No difference between CS and vaginal birth could be found for brachial plexus injuries once controlled for birth weight and presentation (OR 0.5, 95% CI 0.1 to 1.9). \textsuperscript{[evidence level 3]}

#### 8.4 Thermal care for babies born by CS

Descriptive studies report that babies born by CS have lower body temperatures\textsuperscript{509,510} \textsuperscript{[evidence level 3]}. Standard care includes a warm environment for the newborn. We did not identify any studies that address the specific requirements for thermal care for babies born by CS. One RCT showed that fathers can effectively achieve heat conservation in healthy newborn infants.\textsuperscript{511} \textsuperscript{[evidence level 1b]}

Skin-to-skin contact for women and their newborn babies is addressed in Section 8.5.

### Number Recommendation

| 96 | Babies born by CS are more likely to have a lower temperature, and thermal care should be in accordance with good practice for thermal care of the newborn baby. \[GPP] [2004] |

### Number Research Recommendation

| RR 42 | Research is required to establish the thermal care requirements for babies born by CS. |
8.5 Maternal contact (skin-to-skin)

A systematic review was identified that looked at early skin-to-skin contact for women and their healthy babies. Sixteen RCTs and one quasi-randomised trial were included (n = 806). Two of these RCTs included women having CS. The methodological quality of 12 of the included RCTs was poor. Overall, early skin-to-skin contact was associated with higher rates and longer duration of breastfeeding (OR 2.15, 95% CI 1.10 to 4.22. WMD 41.99, 95% CI 13.97 to 70.0) reduced infant crying (OR 21.89, 95% CI 5.2 to 92.3) and higher score summary score of maternal affection. There were no apparent negative effects. One RCT included only women having CS and used three different instruments to evaluate the impact of early contact (within 12 hours of birth) on maternal perceptions of their infant, mothering skills and maternal behaviour. They found significant differences between the groups that had early versus late or limited (after 12 hours) contact and found early skin-to-skin contact to be of benefit. However, these differences were less marked one month after birth.513

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<th>Number</th>
<th>Recommendation</th>
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<tbody>
<tr>
<td>97</td>
<td>Early skin-to-skin contact between the woman and her baby should be encouraged and facilitated because it improves maternal perceptions of the infant, mothering skills, maternal behaviour, and breastfeeding outcomes, and reduces infant crying. [A] [2004]</td>
</tr>
</tbody>
</table>

8.6 Breastfeeding

At least 70% of women express a preference for a birth that would give them the best start to breastfeeding.4 The RCTs that compare planned vaginal birth with planned CS include only women with small, preterm or term breech babies. Three RCTs40,42,514 measure uptake of breastfeeding either as rates of breastfeeding at discharge from hospital or as “any attempt at breastfeeding”.40,42,514 Overall, no difference was detected between the two groups (Pooled RR 0.94, 95% CI 0.89 to 1.00). [evidence level 1a].

One RCT514 also surveyed women at three months to ask if breastfeeding had been initiated at any time and if they were currently breastfeeding. At three months no difference in breastfeeding rates was detected between the groups. (Planned CS group 68%, planned vaginal birth group 70% RR 0.98, 95% CI 0.92 to 1.05). [evidence level 1b]

In the non intention to treat analysis, 73–77% of women who had a vaginal birth and 65–67% of those who had CS, had breastfed at three months after birth.514

Six relevant population studies were identified.515–520 These included diverse populations from several countries including one from the UK.515 In this latter study (n = 202), breastfeeding rates were 76% among those who delivered vaginally and 39% among those who had a CS. [evidence level 2a] Rates of breastfeeding vary markedly between countries from around 30% in Hong Kong518 to more than 90% in Scandinavia519,520 [evidence level 2a] In all studies rates of initiation of breastfeeding were higher in women who had had a vaginal birth compared to those having a CS. Two of the studies517,519 followed women up for 3 months, and one519 followed women up for 6 months. There was no difference in breastfeeding rates according to mode of birth at either 3 or 6 months. [evidence level 2a]

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<tbody>
<tr>
<td>98</td>
<td>Women who have had a CS should be offered additional support to help them to start breastfeeding as soon as possible after the birth of their baby. This is because women who have had a CS are less likely to start breastfeeding in the first few hours after the birth, but, when breastfeeding is established, they are as likely to continue as women who have a vaginal birth. [A] [2004]</td>
</tr>
</tbody>
</table>
9 Care of the woman after CS

Common complications and the estimated frequency with which they occur are shown in Table 4.5.

High dependency unit/intensive therapy unit admission

Maternal mortality is rare. In the UK it is 11.4/100,000 maternities,\textsuperscript{95} [evidence level 3] the direct maternal mortality rate from all causes is 1/20,000 maternities. The mortality rate for women who have vaginal deliveries is 16.9/million compared to 82.3 per million for women who have CS (RR 4.9, 95% CI 2.96 to 7.97).\textsuperscript{95} However it was not possible to determine the proportion of the increased risk that is attributable to antecedent conditions or the procedure itself. The incidence of severe morbidity for women giving birth has been reported to be 12 per 1000 deliveries.\textsuperscript{292} A small proportion of women (0.1–0.9%) develop complications of pregnancy that require admission to an Intensive Therapy Unit (ITU).\textsuperscript{521} HDU/ITU admission was not reported as an outcome in any of the RCTs.

In the NSCSA, 10% of women who had CS required special care postoperatively within a high dependency unit, 3.5% of these women were transferred to an intensive care unit.\textsuperscript{4} [evidence level 3]

Table 9.1 shows the proportion of women who had CS and required admission to an intensive care unit according to the reason for the CS.

We identified one case control study that examined risk factors associated with intensive care unit admission during hospital stay for childbirth among women in USA between 1984 and 1997 (n = 2046).\textsuperscript{522} The overall rate of admission to ICU was 0.13%. The odds of admission to ICU was significantly higher for women who had CS compared with those who had vaginal birth, after adjustment for socio demographic factors (age and ethnicity) and type of hospital (OR 9.0, 95% CI 7.24 to 11.16).\textsuperscript{[evidence level 3]} However it is not possible to disentangle the effect of CS from the reasons for CS when interpreting these results. A UK study that evaluated the risk of severe obstetric morbidity has not been included here because the comparison groups are between women who had unplanned CS to women who had either planned CS or vaginal births.\textsuperscript{292} [evidence level 3]

<table>
<thead>
<tr>
<th>Reason for CS</th>
<th>Admission to ICU (%)</th>
<th>OR (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Breech</td>
<td>0.2</td>
<td>1.00</td>
</tr>
<tr>
<td>Placenta praevia, actively bleeding</td>
<td>2.5</td>
<td>16.6 (5.3 to 52.2)</td>
</tr>
<tr>
<td>Placenta praevia, not actively bleeding</td>
<td>1.1</td>
<td>7.0 (2.2 to 22.1)</td>
</tr>
<tr>
<td>Placental abruption</td>
<td>1.1</td>
<td>7.2 (1.7 to 30.4)</td>
</tr>
<tr>
<td>Pre-eclampsia/eclampsia/HELLP</td>
<td>1.9</td>
<td>12.4 (4.3 to 35.5)</td>
</tr>
<tr>
<td>Maternal medical disease</td>
<td>2.7</td>
<td>17.8 (6.4 to 49.2)</td>
</tr>
<tr>
<td>Uterine rupture</td>
<td>6.4</td>
<td>43.3 (9.9 to 189.5)</td>
</tr>
</tbody>
</table>
99  Healthcare professionals caring for women after CS should be aware that, although it is rare for women to need intensive care following childbirth, this occurs more frequently after CS (about 9 per 1000). [B] [2004]

9.1 Routine monitoring after CS

There were 3 deaths in the last CEMD triennium report in which poor postoperative care was a contributing factor. The importance of monitoring the patient adequately postoperatively was emphasised. [evidence level 3] Earlier triennial reports recommended electronic monitoring of oxygen saturation levels. [evidence level 3] UK obstetric anaesthesia guidelines suggest that the postoperative care of a CS patient should be in accordance with the care of any postoperative patient as laid out in guidelines for postanaesthetic recovery. [evidence level 4]

After CS, women should be observed on a one-to-one basis by an anaesthetist, recovery nurse, midwife or other properly trained member of staff until they have regained airway control and cardiorespiratory stability and are able to communicate. All recovery rooms must be staffed to a level which allows this to be routine practice. Women must be kept under clinical observation at all times and all measurements must be recorded. The introduction of automatic recording systems is encouraged. The frequency of recordings will depend on the stage of recovery and clinical condition of the patient. As a minimum non-invasive blood pressure, heart rate and rhythm, respiratory rate and continuous pulse oximetry every 5 minutes for the first 30 minutes in recovery (‘recovery’ refers to any area where the patient is cared for immediately postoperatively and is not limited to a specific recovery room. The following information should be recorded:

- level of consciousness
- haemoglobin saturation and oxygen administration
- blood pressure
- respiratory frequency
- heart rate and rhythm
- pain intensity e.g. verbal rating scale
- intravenous infusions
- drugs administered.

Other parameters depending on circumstances e.g. temperature, urinary output, central venous pressure, end tidal CO₂, surgical drainage.

For all women, the name, hospital number, time of admission, time of discharge and destination should be recorded in a central register.

Women with epidural or intrathecal analgesia will need additional observations including pain and sedation scores, respiratory rate and mobility which should be laid out in individual hospital protocol. This recording will normally be continued after discharge from the recovery area. It is generally accepted that after discharge from the recovery area to the ward, observations (respiratory rate, heart rate, blood pressure, pain and sedation) should be continued every half hour for two hours and hourly thereafter provided that the observations are stable or satisfactory. If these observations are not stable, more frequent observations and medical review are recommended.

For women who have had intrathecal opioids, there should be a minimum hourly observation of respiratory rate, sedation and pain scores for at least 12 hours for diamorphine and 24 hours for morphine. For epidural opioids and opioid PCA, there should be routine hourly monitoring of the latter throughout the duration of the treatment plus a further period of at least 2 hours after discontinuation.
An ECG, nerve stimulator, thermometer and capnograph should be readily available as well as facilities for resuscitation and emergencies. Women should only be discharged from the recovery area once they have been assessed by a trained recovery staff member and should be taken to the postoperative ward with all of their case notes. In addition no patient should be returned to a general ward unless control of emesis and postoperative pain is satisfactory. After the first 30 minutes postoperatively if the patient is stable then observations are carried out and documented half hourly, 2 hourly and then 4 hourly.\textsuperscript{524,524} [evidence level 4]

<table>
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<tr>
<th>Number</th>
<th>Recommendation</th>
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<tbody>
<tr>
<td>100</td>
<td>After CS, women should be observed on a one-to-one basis by a properly trained member of staff until they have regained airway control and cardiorespiratory stability and are able to communicate. [D] \textsuperscript{[2004]}</td>
</tr>
<tr>
<td>101</td>
<td>After recovery from anaesthesia, observations (respiratory rate, heart rate, blood pressure, pain and sedation) should be continued every half hour for 2 hours, and hourly thereafter provided that the observations are stable or satisfactory. If these observations are not stable, more frequent observations and medical review are recommended. [GPP] \textsuperscript{[2004]}</td>
</tr>
<tr>
<td>102</td>
<td>For women who have had intrathecal opioids, there should be a minimum hourly observation of respiratory rate, sedation and pain scores for at least 12 hours for diamorphine and 24 hours for morphine. [GPP] \textsuperscript{[2004]}</td>
</tr>
<tr>
<td>103</td>
<td>For women who have had epidural opioids or patient-controlled analgesia with opioids, there should be routine hourly monitoring of respiratory rate, sedation and pain scores throughout treatment and for at least 2 hours after discontinuation of treatment. [GPP] \textsuperscript{[2004]}</td>
</tr>
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</table>

9.2 Pain management after CS

In the UK, intrathecal analgesia, patient controlled analgesia, local anaesthetic wound infiltration and nonsteroidal anti-inflammatory agents are commonly used for analgesia post-CS.

Intrathecal analgesia

Key issues related to intrathecal analgesia post-CS are which drug and dose to use as most side effects (particularly with morphine) are dose related.\textsuperscript{525} [evidence level 3] Morphine was commonly used until diamorphine was shown to be a useful alternative.\textsuperscript{526} [evidence level 1b] One RCT comparing intrathecal morphine with normal saline (n = 60) reported that the group given intrathecal morphine had less pain as measured by visual analogue scale (VAS) at 4 and 24 hours postoperative (p < 0.05) and morphine consumption was lower (p < 0.01).\textsuperscript{527} [evidence level 1b] The documented side effects of intrathecal morphine include itching, nausea and vomiting. Alternative intrathecal opioids have been used more frequently because they have fewer reported side effects. One RCT (n = 40) comparing intrathecal diamorphine with intrathecal morphine reported no difference in VAS for pain or overall PCA morphine use. However VAS for itching and drowsiness were higher in the morphine group.\textsuperscript{528} [evidence level 1b] The second RCT (n = 40) used 0.3 mg of intrathecal diamorphine post spinal anaesthesia for CS. The women then used morphine PCA. The median amount of PCA morphine used over 24 hours was less in the group that received the intrathecal...
diamorphine (5 mg versus 45 mg, p < 0.05) and the time to request for first morphine dose was less
(340 minutes versus 80 minutes, p = 0.0006).\textsuperscript{530} [evidence level 1b]

One RCT (n = 80) of women undergoing planned CS with spinal anaesthesia randomised to receive
one of four doses of intrathecal diamorphine for post-CS analgesia (0.125 mg, 0.25 mg, 0.375 mg, or
saline). The optimal intrathecal dose of diamorphine for intrathecal post-CS analgesia is reported to
be between 0.25 mg and 0.375 mg. Nausea and pruritus increased as dose increased.\textsuperscript{531} [evidence
level 1b] Higher doses than this have also been suggested because the minimum dose of intrathecal
diamorphine required to prevent intraoperative supplementation of spinal anaesthesia for CS is 0.4
mg.\textsuperscript{532}

Epidural diamorphine 2.5 mg to 5 mg is an alternative to intrathecal diamorphine as a significant
proportion of unplanned CS (34\%) are carried out using epidural anaesthesia.\textsuperscript{312} [evidence level 3]
This has been evaluated in 2 RCTs. One RCT (n = 50) showed no difference in the duration and
quality of analgesia between intrathecal and epidural diamorphine. There was no difference in the
incidence of pruritus between the two groups but there was a higher incidence of nausea and
vomiting in the epidural group (24\% vs. 4\%, p < 0.05).\textsuperscript{533} [evidence level 1b] The other RCT (n = 53)
comparing epidural with intrathecal diamorphine reported that time to first request for morphine and
side effects were similar between the two groups but VAS pain scores and additional morphine
consumption was higher in the intrathecal group (p = 0.03 and p = 0.03 respectively).\textsuperscript{534} [evidence
level 1b]

Another RCT compared intramuscular administration of diamorphine and four epidural regimens for
the administration of diamorphine. Time to next analgesia was shorter in the intramuscular group
when compared to any of the epidural groups (3.53 hours vs. 5.7, p = 0.007).\textsuperscript{535} [evidence level 1b]

A small American cost-effectiveness study was identified that evaluated the addition of intrathecal
morphine to a regimen of oral analgesia. The effectiveness data was gathered retrospectively for 55
patients. The comparator was patient controlled analgesia. There was no synthesis of costs and
benefits. The authors reported that the mean intrathecal morphine cost US$15 (1997 prices)
compared with US$35 for patient controlled analgesia. Nursing time was also significantly reduced for
intrathecal analgesia patients. Since there were no reported differences in pain control or side-effects,
the study concluded that the addition of intrathecal morphine was a less expensive and less time
consuming and therefore the more cost-effective option.\textsuperscript{536}

**Patient-controlled analgesia**

Patient-controlled analgesia (PCA) is either epidural patient-controlled analgesia (EPCA) or via an
infusion pump device. In the UK EPCA is not common practice post-CS and hence is not considered
here

We did not identify any RCTs that evaluate the effectiveness of PCA compared to other forms of
analgesia after CS. We identified two RCTs that compared different drugs for PCA after CS using a
pump device. One RCT (n = 77) compared morphine alone to morphine combined with alfentanil for
PCA. The group with alfentanil and morphine scored higher on a written questionnaire in terms of
speed of onset of effectiveness of analgesia but there were no differences in terms of grading for
duration of analgesia or overall satisfaction.\textsuperscript{537} [evidence level 1b] The other RCT compared morphine
to fentanyl and found no difference in patient satisfaction or provision of effective analgesia over 37
hours.\textsuperscript{538} [evidence level 1b]

**Wound infiltration with local anaesthetic**

Three RCTs evaluated the use of wound infiltration and nerve blocks for post-CS analgesia
specifically. One RCT (n = 45) used 20 ml of 0.1% bupivacaine infiltrated into the CS wound. They
randomised the women into three groups: one group had general anaesthetic and wound infiltration;
one group regional anaesthetic and wound infiltration and one group general anaesthetic only. They
reported that the two groups that had wound infiltration did not use any pethidine in the first 6 hours
postoperatively compared to the group with no infiltration in which all the women needed at least one
dose of pethidine within the first 6 hours.\textsuperscript{539} [evidence level 1b]

Another RCT (n = 62) compared the effectiveness of bilateral ilioinguinal nerve block and wound
infiltration with 0.5% bupivacaine for postoperative analgesia after CS. Mean VAS scores and mean
papaveretum (morphine derivative) requirements were compared at 4, 8, 12, 16, 20 and 24 hours post-CS. Mean VAS scores for the ilioinguinal block group were reduced compared to control at 4, 8, 12, 20 and 24 hours and papaveretum requirements were less at 4, 8, 12 and 20 hours. Mean VAS scores for the wound infiltration group were reduced compared with the control group at 4 and 12 hours and papaveretum requirements less at 4, 8 and 12 hours (p < 0.05). A review of 26 RCTs, (n = 1211) evaluating the effectiveness of wound infiltration with local anaesthetic in a range of general surgical abdominal operations. Outcome measures were pain scores, supplementary analgesics and time to first analgesic requirements. Overall the study did not find any effect of local anaesthetic wound infiltration for postoperative pain.

**Non-steroidal anti-inflammatory analgesia**

Non-steroidal anti-inflammatory drugs (NSAIDs) are used together with other modalities of pain relief after CS mainly to reduce the need for morphine based analgesics. We considered evidence on NSAID preparations available in the UK.

Two RCTs looked at the analgesic sparing effect of rectal NSAIDs suppository (diclofenac) administered immediately post-CS. In one RCT (n = 50) there was no difference in the VAS scores but the time to request for first analgesia was prolonged with rectal NSAID from 13 hours 45 minutes in the placebo group to 18 hours 58 minutes in the study group (p < 0.03). The other RCT (n = 45) used the amount of PCEA as an outcome measure as well as VAS scores of pain. Women who received the rectal NSAID used less PCEA local anaesthetic solution (52.8 ml) compared to the control group (74 ml). There was no difference in VAS pain scores.

Another RCT (n = 50) administered the NSAID (75 mg diclofenac) intramuscularly to women who were using morphine based PCA post-CS. The women who had the NSAID consumed less morphine via the PCA than the control group (mean at 18 hours post-CS was 61.4 mg compared with 91.4 mg).

An audit of epidural related complications from Australia reports rates of complications for regional anaesthesia as follows: need for re insertion of epidural catheter 4.7%; hypotension after epidural for CS 28%; inadequate block 1.7%; conversion to general anaesthetic 0.5%. Serious complications are relatively rare: unexpected high block 0.07%; high block requiring intubation 0.02% respiratory depression 0.06% and local anaesthetic toxicity 0.04%.

<table>
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<tr>
<th>Number</th>
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<tbody>
<tr>
<td>104</td>
<td>Women should be offered diamorphine (0.3–0.4 mg intrathecally) for intra- and postoperative analgesia because it reduces the need for supplemental analgesia after a CS. Epidural diamorphine (2.5–5 mg) is a suitable alternative. [A] [2004]</td>
</tr>
<tr>
<td>105</td>
<td>Patient-controlled analgesia using opioid analgesics should be offered after CS because it improves pain relief. [GPP] [2004]</td>
</tr>
</tbody>
</table>
Providing there is no contraindication, non-steroidal anti-inflammatory drugs should be offered post-CS as an adjunct to other analgesics, because they reduce the need for opioids.  

[A] [2004]

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**Number**  
**Research Recommendation**

RR 43  
Further research is needed to determine the effect of wound infiltration with local anaesthetic at CS on the need for post-CS analgesia.

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### 9.3 Early eating and drinking after CS

A systematic review compared early with delayed oral fluids and food after CS and included 6 RCTs. Three RCTs were limited to CS with regional anaesthesia; the other 3 RCTs included both regional and general anaesthesia. The intervention groups varied (either allowing immediate access to fluids and food within 6–8 hours if the woman was hungry or thirsty). The comparison groups delayed oral intake for a minimum of 12 hours to 24 hours, or to the presence of bowel sounds and graduated intake. Early eating and drinking was associated with reduced time to return of bowel sounds (1 RCT, n = 118; weighted mean difference of –4.3 hours, 95% CI –6.78 hours to –1.82 hours) and reduced postoperative hospital stay (2 RCTs, n = 220). There was no difference between the intervention and control groups with respect to nausea, vomiting, abdominal distension, time to bowel action, paralytic ileus and number of analgesic doses.  

[evidence level 1a]

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**Number**  
**Recommendation**

107  
Women who are recovering well after CS and who do not have complications can eat and drink when they feel hungry or thirsty.  

[A] [2004]

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### 9.4 Urinary catheter removal after CS

Urinary bladder catheters are commonly used during CS to prevent damage to the bladder during surgery. The effect of urinary bladder catheterisation at CS has been described in a prospective survey (n = 8402) as a risk factor for postpartum urinary retention. Evidence to determine timing of removal of the urinary catheter and the value of routine indwelling catheterisation is currently under review. We identified two RCTs on this topic. One RCT compared immediate catheter removal to removal of an indwelling catheter the next day in women who had a CS under general anaesthetic (n = 107). They report no difference in incidence of urinary tract infection (RR 1.64, 95% CI 0.80 to 3.34) but more instances of urinary retention with intermittent catheterisation (39% vs. 0%). A small RCT compared urinary retention after CS with a general anaesthetic to urinary retention after CS with an epidural anaesthetic and found no difference.  

[evidence level 1b]

Another RCT (n = 78) compared removal of the urinary bladder catheter immediately post-operatively with removal the next day in women undergoing gynaecological (pelvic) surgery, 29 who had CS. They found no difference in the incidence of urinary tract infection, urinary retention or fever but the method of randomisation is unclear and data given in the paper is incomplete.  

[evidence level]

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**Number**  
**Recommendation**

108  
Removal of the urinary bladder catheter should be carried out once a woman is mobile after a regional anaesthetic and not sooner than 12 hours after the last epidural ‘top up’ dose.  

[D] [2004]
9.5 Respiratory physiotherapy after CS

One RCT (n = 120) has evaluated the effect of respiratory physiotherapy after CS under general anaesthesia. The RCT did not detect a difference between the intervention group who had post-CS respiratory physiotherapy and the control group for coughing, phlegm, body temperature, chest palpation and auscultation.\[555 [evidence level 1b]

<table>
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<tr>
<td>109</td>
<td>Routine respiratory physiotherapy does not need to be offered to women after a CS under general anaesthesia, because it does not improve respiratory outcomes such as coughing, phlegm, body temperature, chest palpation and auscultatory changes. [A] [2004]</td>
</tr>
</tbody>
</table>

9.6 Debriefing for women after CS

A longitudinal study, based in Australia suggests that a high level of obstetric intervention during childbirth, such as unplanned CS is associated with the development of acute traumatic symptoms in women postnatally.\[556 [evidence level 3] Midwife led debriefing has been proposed to be of value in reducing the incidence of depression and anxiety after birth. A systematic review (11 RCTs) evaluating the effect of psychological debriefing on the prevention of post traumatic stress disorder (PTSD) in the general population reported that single session individual debriefing did not affect the incidence of PTSD at 3 to 5 months (6 RCT, n = 387, OR 1.22, 95% CI 0.60 to 2.46) and increased the likelihood of long term PTSD (after one year 2 RCTs, n = 238, OR 2.04, 95% CI 0.92 to 4.53).\[557 [evidence level 1a] Only two of the included studies were in an obstetric setting.\[558,559 Of these two trials, one was UK based (n = 129) and included primigravid women who had a normal vaginal birth. Women who received midwife debriefing were less likely to have high anxiety and depression scores after birth than women who did not (anxiety score OR 13.5, 95% CI 0.41 to 56.9; depression OR 8.5, 95% CI 2.8 to 30.9).\[559 [evidence level 1b] The second RCT was from Australia (n = 1041) looked at the effect of midwife-led debriefing on maternal depression after operative childbirth. No difference was detected in depression scores (OR 1.24, 95% CI 0.87 to 1.77) or in the proportion of women who reported that depression had been a problem at six months after the birth (OR1.37, 95% CI 1.00, 1.86)\[558 [evidence level 1b]

Subsequently a further two RCTs have been published. One RCT (n = 103) tested opportunity to debrief at an initial postnatal interview (less than 72 hours postpartum) and 4–6 weeks postpartum to usual care. The RCT reported a high baseline prevalence of post-traumatic stress disorder (9.6% of women at 4 to 6 weeks postpartum). No difference was detected in the prevalence of symptom profile for PTSD immediately following debriefing or at 3 months. (RR 1.06, 95% CI 0.61 to 1.84). This is an RCT is underpowered to detect a 2% difference in prevalence of symptoms of post-traumatic stress disorder.\[560 [evidence level 1b] A recently published RCT (n = 1745) compared a midwife debriefing session within 72 hours of birth to usual care. No differences were detected between the groups for either stress disorders or depression (assessed EPDS and report of depressive illness).\[561 [evidence level 1b]
Number | Recommendation
--- | ---
1 | The recommendation for this review has been amended and updated following a new (2011) review. The new recommendation can be found in section 11.2

Number | Research Recommendation
--- | ---
RR 45 | More RCT evidence is required to determine the effect of midwifery-led debriefing following CS.

### 9.7 Length of hospital stay and readmission to hospital

Length of hospital stay after childbirth is declining; recent routine national statistics for England\(^{562}\) suggest that women who have a spontaneous vaginal delivery spend on average 1 day in hospital, women who have an instrumental delivery spend 1 or 2 days in hospital and women who have a CS spend 3 or 4 days in hospital.

In one RCT\(^{48}\) that compared planned CS with planned vaginal birth, the median length of hospital stay for women in the planned CS group was 4 days (5th centile 1.7 days, 95th centile 7.4 days). For women in the planned vaginal birth group it was 2.8 days (0.8, 6.9 days). The median length of stay reported in this RCT\(^{48}\) is compatible with routine maternity statistics for the U.K. In 3 RCTS\(^{38,39,42}\) the length of hospital stay was reported as either greater or less than 10 days. On pooling these results, the relative risk of length of hospital stay greater than 10 days for women in the planned CS group was 1.27 (95% CI 0.35 to 4.65). [evidence level 1a]

### Readmission to hospital

Infection and bleeding constitute the main reasons for readmission to hospital following birth.\(^{563}\) Two surveys of women in the postpartum period have estimated about 3% are readmitted to hospital for reasons related to their own health.\(^{563,564}\)

Readmission to hospital was not included as an outcome measure in the RCTs of planned CS versus planned vaginal birth.

One prospective cohort study in Australia\(^{564}\) examined rates of readmission to hospital within 8 weeks of birth. A higher proportion of women who had CS (5.3%) reported readmission to hospital compared to women who had vaginal birth (2.2%) (OR 2.46, 95% CI 1.11 to 5.43). [evidence level 2b] Similar findings were reported in a retrospective cohort study conducted in Washington USA\(^{565}\) (n = 256,795). The age adjusted relative risk for rehospitalisation among women who had CS compared to those who had vaginal birth was increased (RR 1.8, 95% CI 1.6 to 1.9). [evidence level 2b]

Discharge from hospital after CS usually occurs on day 3.\(^{562}\) [evidence level 3] A systematic review of early post natal discharge from hospital included eight RCTs but only two RCTs included women who had caesarean births, one of which is ongoing.\(^{566}\) [evidence level 1a] The RCT (n = 61) randomised women having CS to either early hospital discharge and home follow up or usual hospital discharge (requires the woman to be ambulatory, voiding, tolerating a normal diet, passing flatus, normal uterine involution, afebrile for 24 hours, uncomplicated wound healing, removal of skin sutures or staples and an adequate blood count). Women in the intervention group were discharged when they met the same criteria other than afebrile for 24 hours and staple or suture removal. They report no difference in maternal or infant rehospitalisations, maternal affect or overall maternal functional status. Women in the early discharge group were more satisfied with care and had a 29% reduction in health care requests.\(^{567}\) [evidence level 1b]
<table>
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<th>Recommendation</th>
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<tbody>
<tr>
<td>110</td>
<td>Length of hospital stay is likely to be longer after a CS (an average of 3–4 days) than after a vaginal birth (average 1–2 days). However, women who are recovering well, are apyrexial and do not have complications following CS should be offered early discharge (after 24 hours) from hospital and follow-up at home, because this is not associated with more infant or maternal readmissions.</td>
</tr>
</tbody>
</table>
Recovery following CS

Postnatal advice for women who have had a CS includes general and specific advice. Specific advice includes advice on CS wound care, analgesia at home, when to resume normal activities such as driving, exercise and sexual intercourse and the provision of detailed information on possible risks associated with CS birth and possible complications. Information on the risk and benefits of CS should have been discussed prior to CS however they should be reiterated again. It is outside the scope of this guideline to consider general post natal advice. General advice has been developed and published as part of RCT (IMPaCT study). [evidence level 3]

Pain

Antenatally about 60% women express a preference for a birth that is as pain free as possible and for a quick recovery. Assessment of pain during the immediate postoperative period is not reported in any of the RCTs. One RCT (n = 1596) report on abdominal, perineal and back pain at three months after birth. [evidence level 1b] Four cohort studies involving a total of 4749 women in Australia, USA and Scotland reported on pain between 2 weeks to 18 months after birth.

Three months after delivery women who had planned CS were more likely to report pain in the abdomen (RR 1.76, 95% CI 1.24 to 2.50), and pain deep inside the abdomen (RR 1.89, 95% CI 1.29 to 2.79) than women who had planned vaginal birth at three months after birth. Not surprisingly perineal pain is reduced in women who have planned CS (RR 0.32, 95% CI 0.18 to 0.58). [evidence level 1b] At three months after birth there is also no difference in reports of back pain (RR 0.93, 95% CI 0.71 to 1.22). [evidence level 1b] Back pain is common, 22% to 50% of women surveyed report having back pain at either 8, 16 or 24 weeks after birth. Mode of birth has not been found to affect rates of back pain. [evidence level 1b]

In cohort studies 60% of women who had a CS (either planned CS or CS in labour), reported having wound pain at 24 weeks after birth. [evidence level 2b]

There is little direct evidence to guide prescribing practice of analgesia after discharge from hospital for women who have had a CS with no complications. Current guidelines on post-CS wound care suggest that for mild post-CS pain paracetamol (1000 mg four times daily) should be prescribed, for moderate pain co-codamol (1 to 2 tablets four times daily) and for severe pain co-codamol with added ibuprofen (500 mg twice daily). [evidence level 3]

Wound care

General CS wound care advice for women includes encouraging women to take prescribed analgesia, to complete antibiotics if prescribed, to wear loose comfortable clothes and cotton underwear, to bathe or shower daily, to gently clean and dry the wound well (flannels or washcloths should be freshly laundered) and only apply dressings if advised by the doctor or midwife. [evidence level 3]

Infection

Evidence from cohort studies report an increased risk of postpartum endometritis among women who had CS compared to those who had spontaneous vaginal birth (RR 4.51, 95% CI 4.00 to 5.09). [evidence level 2b] For this reason prophylactic antibiotics are prescribed during CS. [evidence level 1a] Overall the impact of CS on risk of infection when antibiotics are used is less clear. No difference was detected in rates of infection between women randomised to have planned CS (6.4%) and planned vaginal birth (4.9%) (RR 1.29, 95% CI 0.97 to 1.72). [evidence level 1a]
Midwives and doctors involved in post natal care of women who have had a CS should retain a high index of suspicion for wound infection, urinary tract infection and endometritis; they should ask the woman about wellbeing and in particular any signs of fever; assess the wound for signs of infection, separation or dehiscence; discuss pain relief requirements and plan to remove sutures or clips when appropriate. [568] [evidence level 3]

**Urinary symptoms**

Urinary symptoms in women who have had a CS are commonly due to urinary tract infection, but can be due to stress incontinence or rarely due to urinary tract injury.

Pregnancy and childbirth are established risk factors for urinary incontinence. Urinary incontinence is the involuntary loss of urine that becomes a social or hygienic problem. [571] [evidence level 4] Women who have had a CS may have urinary incontinence but the risk of incontinence following CS is reduced compared to women who have had a vaginal birth. (3 months following birth planned CS 4.5%, planned vaginal birth 7.3% (RR, 95% CI 0.62 to 0.91). [514] [evidence level 1b] Five cohorts also report an increased risk of urinary incontinence among women who have vaginal deliveries compared to those who have CS. [572-576] [evidence level 2b]. One cohort (n = 149) did not detect any difference in urinary incontinence at 9 weeks by mode of birth. [577] [evidence level 2b] Risk of incontinence increases following pregnancy (10% in the nulliparous women, 16% after CS and 21% after vaginal birth) [574] [evidence level 2b].

The estimated incidence of bladder injury in women delivered by CS is 0.1% and 0.003% in women delivered vaginally (RR 36.59, 95% CI 10.43 to 128.38). Ureteric injury occurred in 0.03% of women who had CS and in 0.001% women who had vaginal birth (RR 25.22, 95% CI 2.63 to 243.50). [578] [evidence level 3] In other studies the frequency is reported to range between 16 per million to 1% [579,578,580,581] Risk factors include repeat CS and peripartum hysterectomy. [580,582,583] [evidence level 4] Two RCTs include bladder/bowel/ureteric injury as an outcome measure. [44,48] There were no events in either group in one RCT, [48] while in the other 1 of the 93 women in the planned CS group, and none of the 115 women in the planned vaginal birth group suffered this morbidity measure. [44] [evidence level 1b].

**Faecal incontinence**

Faecal or anal incontinence has been defined as the involuntary leakage of solid or liquid faeces or gas. [584] One RCT (n = 1596) asked women about symptoms of incontinence of faeces and flatus three months following birth. No difference was detected between the groups. (Incontinence of faeces 0.8% planned CS 1.5%, planned vaginal birth group RR 0.54, 95% CI 0.18 to 1.62. Incontinence of flatus 10.7% planned CS, 9.7% planned vaginal birth RR 1.10, 95% CI 0.79 to 1.54). [514] [evidence level 1b] Non-intention-to-treat analysis was also not different.

Four cohort studies evaluated faecal or anal incontinence according to mode of birth. In two of these studies [584,585] no difference was detected in the prevalence of faecal incontinence among women who had CS and those who had vaginal birth. In the other two studies [586,587] none of the women who had CS were reported to have faecal incontinence. The prevalence of faecal incontinence among women who had vaginal deliveries in these studies ranged from 1% to 23%.

**Resuming activities**

In one cohort study (n = 971) the extent to which bodily pain interfered with usual activities was measured 8 weeks after birth. Women who had CS were more likely to have bodily pain which interfered with usual activity. [570] [evidence level 2b] At six months pain limited physical activity among women who had either CS or assisted vaginal birth when compared with women who had spontaneous vaginal birth after birth. [evidence level 2b]

The Association of Chartered Physiotherapists in Women’s Health (ACPWH) suggests that women who have had a CS should wait 8 to 10 weeks before commencing vigorous exercise. We did not identify any other guidance on exercise after a CS. [568] [evidence level 4].

The Driver and Vehicle Licensing Agency (DVLA) in their guide for medical practitioners as to current medical standards of fitness to drive do not specifically provide guidance on driving after a CS. They provide a general statement on driving after any surgery that suggests that drivers wishing to drive...
after surgery ‘should establish with their own doctors when it is safe to do so’. They add that decisions regarding return to driving should consider recovery from the anaesthesia, distracting effect from the pain of the surgery and any resultant physical restrictions. [evidence level 4]

**Sexual intercourse**

A study of women in their first pregnancy reported the pre-pregnancy prevalence of sexual problems to be 38%. Sexual morbidity increased in the first three months after birth to 83%, declining to 64% at 6 months after birth. [evidence level 2b]

Sexual function after birth has been assessed in one RCT and 4 cohort studies. The measures used to assess this included resumption of sexual activity after birth, and dyspareunia following birth. One RCT evaluated sexual function at 3 months after birth and did not detect any difference between the two groups in the proportion of women who reported (i) not having sex since the birth (RR 1.12, 95% CI 0.89 to 1.42) or (ii) having pain during sex on the most recent occasion (RR 1.03, 95% CI 0.91 to 1.16). [evidence level 1b]

One cohort study (n = 971) included women in their first pregnancy. No difference was detected between women who had CS and those who had vaginal birth (assisted or unassisted). [evidence level 2b] A smaller study from the USA (n = 66) did not detect any difference in dyspareunia at 2–8 weeks postpartum between women who had CS and those who had vaginal birth. [evidence level 2b] The third study reported that one month after birth women who had CS were more likely to have resumed intercourse than women who delivered vaginally. [evidence level 2b] The fourth study reported that dyspareunia was associated with vaginal deliveries and previous experience of dyspareunia in the first 3 months after birth. At six months there was no difference detected in rates of dyspareunia according to mode of birth [evidence level 2b]

**Breastfeeding**

Rates of initiation of breastfeeding are higher among women who had vaginal birth compared with those who had CS. However, by three to six months after birth there is no difference in breast feeding rates between the two groups. [evidence level 1b]

**Postnatal depression**

The incidence of postnatal depression is estimated to be 13%. Self report measures tend to yield higher estimates of postpartum depression than interview-based methods. Depression following childbirth has been assessed by various scales including the Edinburgh Postnatal Depression Scale (EPDS), the Profile of Mood States (POMS), the Beck Depression Inventory, the Zung Depression Scale and the Center for Epidemiological studies Depression scale.

One RCT measured postnatal depression, at 6 weeks (n = 2086) and 3 months (n = 1596). Early postpartum depression occurred in 0.3% of women in the CS group and none in the planned vaginal birth group. It is therefore not possible to estimate a relative risk measure for this outcome. At 3 months no difference was detected in postnatal depression as defined by the Edinburgh Postnatal Depression scale (EPDS) between the groups (RR 0.93, 95% CI 0.70 to 1.24). [evidence level 1b]

Six observational studies have evaluated post natal depression and mode of birth. These studies were conducted in Scotland, Australia, USA and Finland. A variety of methods have been used to assess postnatal depression and the length of follow up varies between 2 weeks to 18 months. Two studies report a higher prevalence of postnatal depression among women who had a CS in the first two weeks after birth compared to those who had a vaginal birth. However, after 8 weeks postpartum, no difference was detected in the prevalence of postnatal depression between the two groups. [evidence level 2b]

**Post-traumatic stress disorder**

None of the RCTs on planned mode of birth have evaluated the impact of this on post-traumatic stress disorder. Two cohort studies from Sweden examined the prevalence of post-traumatic stress disorder between 1 month and 2 years postpartum. No difference was detected in the prevalence of post-traumatic stress disorder between women who had CS and vaginal birth. Compared with women...
who had vaginal birth, a higher proportion of women who had “emergency” CS (OR 6.3, 95% CI 2.0 to 20.2) and those who had assisted vaginal birth (OR 4.8, 95% CI 1.5 to 15.2) had post-traumatic stress disorder at 1–2 years after birth. \textsuperscript{598,599} [evidence level 2b]

**Maternal satisfaction**

One RCT asked women at three months after birth about their likes and dislikes regarding the childbirth experience.\textsuperscript{514} More women in the planned CS group indicated that they liked being able to schedule their birth (RR 1.99, 95% CI 1.66 to 2.40), liked that the childbirth experience was not very painful (RR 1.18, 95% CI 1.05 to 1.31) and felt reassured about their infant’s health (RR 1.13, 95% CI 1.06 to 1.20). However, fewer women in the planned CS group indicated that they ‘liked that birth was natural’ (RR 0.17, 95% CI 0.14 to 0.22), ‘liked actively participating in the birth’ (RR 0.37, 95% CI 0.31 to 0.44) and ‘liked that recovering from the childbirth experience was not difficult’ (RR 0.84, 95% CI 0.77 to 0.92). A similar proportion of women in both groups indicated that they ‘liked the method of birth that they had had’ or ‘felt reassured about their own health’. The proportion of women that reported that ‘there was nothing they liked about their childbirth experience’ was also similar in both groups. No difference was detected between the two groups with regards to either ‘ease in caring for their new infant’ or ‘adjusting to being a new mother’. Similar trends were seen for these outcomes in the non intention to treat analysis. [evidence level 1b]

One cross sectional study\textsuperscript{600} surveyed women within a week of birth in Dublin, Ireland. The CS rate in this study was 10%. 91% of women who had vaginal birth compared with 33% of those who had CS reported that they would like a similar mode of birth for future pregnancies. [evidence level 3]

**Prolapse**

The prevalence of genital prolapse around the menopause has been estimated at 5%. In a case control study (n = 21,449) women attending menopause clinics were examined for uterine prolapse. Previous CS was associated with a 40% reduction in the risk of developing uterine prolapse (OR 0.6, 95% CI 0.5 to 0.8).\textsuperscript{601} [evidence level 3] Another case control in the USA found that women who underwent surgery for uterovaginal prolapse were less likely to have had a CS.\textsuperscript{602} [evidence level 3]

### Number Recommendation

111 In addition to general postnatal care, women who have had a CS should be provided with:

- specific care related to recovery after CS
- care related to management of other complications during pregnancy or childbirth. \[GPP\] \[2004]\n
112 Women who have a CS should be prescribed and encouraged to take regular analgesia for postoperative pain, using:

- for severe pain, co-codamol with added ibuprofen
- for moderate pain, co-codamol
- for mild pain, paracetamol. \[D\] \[2004]\n
113 CS wound care should include:

- removing the dressing 24 hours after the CS
- specific monitoring for fever
- assessing the wound for signs of infection (such as increasing pain, redness or discharge), separation or dehiscence
- encouraging the woman to wear loose, comfortable clothes and cotton underwear
- gently cleaning and drying the wound daily
- if needed, planning the removal of sutures or clips. \[D\] \[2004]\n
114 Healthcare professionals caring for women who have had a CS and who have urinary symptoms should consider the possible diagnosis of:
- urinary tract infection
- stress incontinence (occurs in about 4% of women after CS)
- urinary tract injury (occurs in about 1 per 1000 CS). [D] [2004]

Healthcare professionals caring for women who have had a CS and who have heavy and/or irregular vaginal bleeding should consider that this is more likely to be due to endometritis than retained products of conception. [D] [2004]

Women who have had a CS are at increased risk of thromboembolic disease (both deep vein thrombosis and pulmonary embolism), so healthcare professionals need to pay particular attention to women who have chest symptoms (such as cough or shortness of breath) or leg symptoms (such as painful swollen calf). [D] [2004]

Women who have had a CS should resume activities such as driving a vehicle, carrying heavy items, formal exercise and sexual intercourse once they have fully recovered from the CS (including any physical restrictions or distracting effect due to pain). [GPP] [2004]

Healthcare professionals caring for women who have had a CS should inform women that after a CS they are not at increased risk of difficulties with breastfeeding, depression, post-traumatic stress symptoms, dyspareunia and faecal incontinence. [D] [2004]
Pregnancy and childbirth after CS

11.1 Implications of CS for future pregnancies

Infertility

Infertility is defined as failure to conceive within 1–2 years of unprotected sexual intercourse. Most studies however have measured birth interval, reflecting future live birth rates and not rates of conception. These studies may not have been able to adjust for confounding factors such as use of contraception.

We found one systematic review which included 8 cohort studies in Northern Europe and USA and one further cohort study conducted in England which had addressed this question. Follow-up period in most studies ranged between 3.5 to 6 years, however one study had a follow up period between 1–19 years. Register information or interviews examined outcomes of at least one pregnancy, at least one live birth, all pregnancies, all live births, and fecundity. Almost all studies report that fewer women having a CS will subsequently have children/or will have less children, due to a combination of a lessened desire for, or an incapability of having children. There is a 46% increase in the risk of having no more children five years after primary CS (RR 1.46, 95% CI 1.07 to 1.99). [evidence level 2b]

Sterilisation rates were higher after a CS in 3 studies. The increased risk ranged between 6% and 23%. [evidence level 2b]

Placenta praevia

We identified three recent cohort studies and an earlier systematic review. The incidence of placenta praevia in these studies ranges from 0.2% to 0.5% for women with a previous vaginal birth and 0.4% to 0.8% for women with a previous CS. These studies report a 30% to 60% increase in risk of placenta praevia in subsequent pregnancies for women who had had a previous CS compared to those who had had vaginal deliveries. Three case series have reported on the incidence of placenta praevia and placenta accreta in women who have had previous CS. Overall the incidence of placenta accreta is estimated to be 1 in 2500 pregnancies, however, there is no comparative data for the incidence in women who have not had previous CS.

The incidence of placenta praevia ranges from 0.2% to 0.5% for women with a previous vaginal birth and 0.4% to 0.8% for women with a previous CS. [evidence level 2b].

Stillbirth

A large retrospective cohort study in Scotland (n = 120,633) investigated the association between previous CS and risk of stillbirth in subsequent pregnancies. The risk of antepartum stillbirth among women who had no previous CS was 2 per 1000 compared to 4 per 1000 among women who had a previous CS (hazard ratio 1.64, 95% CI 1.17 to 2.30). The risk of unexplained stillbirth associated with previous CS differed with gestational age, the excess risk was apparent from 34 weeks (hazard ratio 2.23, 95% CI 1.48 to 3.36). [evidence level 2b]
11.2 Pregnancy and childbirth after CS

Introduction
A recent study of 146 NHS Trusts in England (Bragg et al., 2010) found that one in four women had a CS. The rise in primary CS rates has led to an increased proportion of women of reproductive age with a scarred uterus. Thus, the issue of the most appropriate mode of delivery following a CS continues to be the subject of continued research and debate.

This chapter presents the best available evidence to facilitate antenatal counselling and decision making when planning the mode of birth following one or more previous CS.

Review question
What are the risks and benefits of planned caesarean section compared with planned vaginal birth for both women and babies in women who have had a previous caesarean section?

Overview of evidence
Four studies were included in this review (Guise et al., 2010; Cahill et al., 2010; Tahseen & Griffiths, 2010; Law et al., 2010).

Of the four included studies in this review, one is a systematic review (Guise et al., 2010), one was conducted in the USA (Cahill et al., 2010), one in the UK (Tahseen & Griffiths, 2010) and one in Hong Kong (Law et al., 2010). One study is a large, rigorous systematic review of observational studies (Guise et al., 2010). One study (Tahseen & Griffiths, 2010) performed a systematic review of observational studies of success rate and adverse maternal and neonatal outcomes of vaginal birth after one or two CS and repeat CS.

One study (Cahill et al., 2010) reported maternal morbidity in women with three or more prior caesarean births who attempt a vaginal birth (VBAC).

One study (Law et al., 2010) reported maternal psychological status among women with one previous caesarean birth who were randomised to planned vaginal birth or planned CS.

Evidence profile
One evidence profile summarises maternal outcomes from one systematic review plus one randomised trial of the risks and benefits of "elective" repeat CS [ERCS] compared with trial of labour [TOL] (Guise et al., 2010, Law et al., 2010). One evidence profile summarises neonatal outcomes from one systematic review of the risks and benefits of ERCS compared with TOL (Guise et al., 2010). Three evidence profiles report maternal complications associated with repeat CS as reported by the same systematic review (Guise et al., 2010). One evidence profile reports maternal outcomes of vaginal birth or planned CS after two previous CS compared with vaginal birth or planned CS after one previous CS (Tahseen & Griffiths, 2010). Maternal morbidity in women who plan vaginal birth after three or more prior CS is detailed in one evidence profile reporting findings from one observational study (Cahill et al., 2010). All included studies were observational studies. Therefore, using the GRADE system, the quality of the evidence was moderate, low or very low for all studies.

Maternal outcomes
All of the data included in this table have been taken from one systematic review and one randomised trial and details outcomes for women who have had one previous CS. The number in the first column indicates the number of studies within the review that contribute data to that outcome.

Table 11.1 GRADE summary of findings comparing planned CS with planned vaginal birth in women with a previous CS (maternal outcomes)

<table>
<thead>
<tr>
<th>Number of studies</th>
<th>Number of women</th>
<th>Effect</th>
<th>Quality</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Planned CS</td>
<td>Planned vaginal birth</td>
<td>Relative (95% CI)</td>
</tr>
</tbody>
</table>

Caesarean section: full guideline DRAFT (September 2011)
### Maternal mortality (term)

| 4 studies | 17/225,239 (7.5 per 100,000) | 3/156,690 (1.9 per 100,000) | RR 3.94 (1.20 to 12.5)* | Absolute risk difference: 5.6 more deaths per 100,000 (from 1.2 more to 10.4 more) | Moderate |

### Maternal mortality (any gestational age)

| 12 studies | 19/229,635 (8.2 per 100,000) | 5/167,220 (3.0 per 100,000) | RR 2.76 (1.07 to 7.14)* | Absolute risk difference: 5.3 more deaths per 100,000 (from 0.4 more to 10.3 more) | Moderate |

### Uterine rupture (term)

| 2 studies | 4/18195 (0.22 per 1000) | 118/16250 (7.26 per 1000) | RR 0.03 (0.011 to 0.082)* | Absolute risk difference: 7.04 fewer per 1000 (from 8.5 fewer to 5.8 fewer)* | Very low |

### Uterine rupture (any gestational age)

| 4 studies | 6/26,535 (0.22 per 1000) | 148/20,717 (7.14 per 1000) | RR 0.031 (0.014 to 0.070)* | Absolute risk difference: 7 fewer per 1000* (Adjusted risk difference: 5.1 fewer per 1000* from 2.3 fewer to 11.2 fewer) | Very low |

### Blood transfusion (term)

| 4 studies | 607/227,960 (2.6 per 1000) | 547/156,690 (3.5 per 1000) | RR 0.76 (0.67 to 0.85)* | Absolute risk difference: 0.9 fewer per 1000* (Adjusted risk difference: 1.4 fewer per 1000 from 0.7 fewer to 2.2 fewer) | Very low |

### Blood transfusion (any gestational age)

<p>| 9 studies | 712/233,884 | 641/167,423 | RR 0.795 | Absolute risk | Very low |</p>
<table>
<thead>
<tr>
<th>(Guise et al., 2010)</th>
<th>(3 per 1000)</th>
<th>(3.8 per 1000)</th>
<th>(0.714 to 0.884)* difference: 0.8 fewer per 1000 (from 1.16 fewer to 0.41 fewer)*</th>
</tr>
</thead>
</table>

**Caesarean section**

**Hysterectomy (term)**

3 studies

<table>
<thead>
<tr>
<th>(Guise et al., 2010)</th>
<th>248/227,479 (1.09 per 1000)</th>
<th>174/155,763 (1.11 per 1000)</th>
<th>RR 0.97 (0.80 to 1.18)* Absolute risk difference: 0.02 fewer per 1000 (from 0.24 fewer to 0.18 more)*</th>
</tr>
</thead>
</table>

**Hysterectomy (any gestational age)**

8 studies

<table>
<thead>
<tr>
<th>(Guise et al., 2010)</th>
<th>280/234,349 (1.19 per 1000)</th>
<th>197/167,710 (1.17 per 1000)</th>
<th>RR 1.01 (0.84 to 1.22)* Absolute risk difference: 0.02 more per 1000 (from 0.19 fewer to 0.23 more)*</th>
</tr>
</thead>
</table>

**Infection: endometritis, chorioamnionitis, wound and other postpartum infections (any gestational age)**

10 studies

<table>
<thead>
<tr>
<th>(Guise et al., 2010)</th>
<th>32 per 1000</th>
<th>46 per 1000</th>
<th>Not calculable (NC) Absolute risk difference: 14 fewer per 1000*</th>
</tr>
</thead>
</table>

**Length of hospital stay (any gestational age)**

8 studies

<table>
<thead>
<tr>
<th>(Guise et al., 2010)</th>
<th>Mean 3.92 days</th>
<th>Mean 2.55 days</th>
<th>NC</th>
<th>MD 1.37 higher</th>
</tr>
</thead>
</table>

**Edinburgh Postnatal Depression Scale (6 months postpartum)**

1 study

<table>
<thead>
<tr>
<th>(Law et al., 2010)</th>
<th>Median 0.0 (inter-quartile range 0.0 – 4.0)</th>
<th>Median 0.5 (inter-quartile range 0.0 – 4.0)</th>
<th>NC</th>
<th>p = 0.766</th>
</tr>
</thead>
</table>

**Beck Depression Inventory (6 months postpartum)**

1 study

<table>
<thead>
<tr>
<th>(Law et al., 2010)</th>
<th>Median 1.5 (inter-quartile range 0.0 – 4.8)</th>
<th>Median 1.0 (inter-quartile range 0.0 – 4.3)</th>
<th>NC</th>
<th>p = 0.929</th>
</tr>
</thead>
</table>

**Client Satisfaction Questionnaire (6 months postpartum)**

1 study

<table>
<thead>
<tr>
<th>(Law et al., 2010)</th>
<th>Median 24.0 (inter-quartile range 22.0 – 25.0)</th>
<th>Median 23.0 (inter-quartile range 22.0 – 25.0)</th>
<th>NC</th>
<th>p = 0.433</th>
</tr>
</thead>
</table>

*Calculated by NCC-WCH technical team*
Repeat CS

Narrative discussions of maternal complications associated with multiple CS were reported in one systematic review (Guise et al., 2010). All participants gave birth by CS. The number of studies contributing to the outcome in question is reported in the first column of the table. For tables 11.2 to 11.4 just one study from within the systematic review (not the same study) contributed data to each outcome. All included studies involved women giving birth at any gestation.

Table 11.2 GRADE summary of findings for repeat CS (1 prior CS vs. 2 prior CS)

<table>
<thead>
<tr>
<th>Number of studies</th>
<th>Number of women</th>
<th>Effect</th>
<th>Quality</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1 prior CS</td>
<td>2 prior CS</td>
<td>Relative (95% CI)</td>
</tr>
<tr>
<td>Blood transfusion rates</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 study (Guise et al., 2010)</td>
<td>427/23,579 (1.8%)</td>
<td>202/7,902 (2.6%)</td>
<td>0.70 (0.60 to 0.83)*</td>
</tr>
<tr>
<td>Infection rates (endometritis)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 study (Guise et al., 2010)</td>
<td>404/14,808 (2.7%)</td>
<td>178/6,324 (2.8%)</td>
<td>0.96 (0.81 to 1.16)*</td>
</tr>
<tr>
<td>Wound complication (infection and wound dehiscence)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 study (Guise et al., 2010)</td>
<td>165/15,808 (1.0%)</td>
<td>107/5,324 (2.0%)</td>
<td>0.55 (0.43 to 0.70)*</td>
</tr>
<tr>
<td>Surgical (bladder) injuries rates</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 study (Guise et al., 2010)</td>
<td>15/15,808 (0.1%)</td>
<td>18/6,324 (0.3%)</td>
<td>0.33 (0.17 to 0.65)</td>
</tr>
</tbody>
</table>

*Calculated by NCC-WCH technical team

Table 11.3 GRADE summary of findings for repeat CS (1 prior CS vs. ≥ 2 prior CS)

<table>
<thead>
<tr>
<th>Number of studies</th>
<th>Number of women</th>
<th>Effect</th>
<th>Quality</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1 prior CS</td>
<td>≥ 2 prior CS</td>
<td>Relative</td>
</tr>
</tbody>
</table>

Caesarean section: full guideline DRAFT (September 2011)
### Blood transfusion rates

<table>
<thead>
<tr>
<th>Study</th>
<th>Number of women</th>
<th>Effect</th>
<th>Quality</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 study (Guise et al., 2010)</td>
<td>16/491 (3.3%)</td>
<td>22/277 (7.9%)</td>
<td>Absolute risk difference: 46 fewer per 1000 (from 56 fewer to 14 fewer)</td>
</tr>
</tbody>
</table>

### Hysterectomy rates

<table>
<thead>
<tr>
<th>Study</th>
<th>Number of women</th>
<th>Effect</th>
<th>Quality</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 study (Guise et al., 2010)</td>
<td>1/491 (0.20%)</td>
<td>3/277 (1.08%)</td>
<td>Absolute risk difference: 9 fewer per 1000 (from 29 fewer to 2 more)</td>
</tr>
</tbody>
</table>

*Calculated by NCC-WCH technical team

### Table 11.4 GRADE summary of findings for repeat CS (1 prior CS vs. 3 prior CS)

<table>
<thead>
<tr>
<th>Number of studies</th>
<th>Number of women</th>
<th>Effect</th>
<th>Quality</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1 prior CS</td>
<td>3 prior CS</td>
<td>Relative (95% CI)</td>
</tr>
<tr>
<td>Surgical (bladder) injuries rates (any gestational age)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 study (Guise et al., 2010)</td>
<td>15/15,808 (0.99%)</td>
<td>17/1452 (1.2%)</td>
<td>0.08 (0.04 to 0.15)</td>
</tr>
<tr>
<td>Infection (endometritis): (any gestational age)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 study (Guise et al., 2010)</td>
<td>404/15,808 (2.5%)</td>
<td>43/1452 (3.0%)</td>
<td>0.86 (0.63 to 1.17)</td>
</tr>
<tr>
<td>Wound complication (infection and wound dehiscence)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 study (Guise et al., 2010)</td>
<td>165/15,808 (1.0%)</td>
<td>22/1452 (1.5%)</td>
<td>0.68 (0.44 to 1.06)*</td>
</tr>
</tbody>
</table>

*Calculated by NCC-WCH technical team
Vaginal birth attempt following two or more CS

All of the data included in this section have been taken from two studies (Cahill et al., 2010; Tahseen & Griffiths, 2010) that reported maternal morbidity in women who attempted VBAC. The range of successful VBACs was 74% to 80% in one observational study (Cahill et al., 2010) and 72% to 76% in the other study (Tahseen & Griffiths, 2010). For the systematic review (Tahseen & Griffiths, 2010) the number of studies reported in the first column of the evidence profile corresponds to the number of included studies contributing findings to each reported outcome.

Table 11.5 GRADE summary of findings for planned VBAC after 2 prior CS versus planned repeat CS after 2 prior CS

<table>
<thead>
<tr>
<th>Number of studies</th>
<th>Number of women</th>
<th>Effect</th>
<th>Quality</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Planned vaginal birth 2 prior CS</td>
<td>Planned repeat CS 2 prior CS</td>
<td>Relative (95% CI)</td>
</tr>
<tr>
<td>Blood transfusion</td>
<td>6 studies (Tahseen &amp; Griffiths, 2010)</td>
<td>47/2,292 (2.1%)</td>
<td>172/10,277 (1.7%)</td>
</tr>
<tr>
<td>Febrile morbidity</td>
<td>6 studies (Tahseen &amp; Griffiths, 2010)</td>
<td>192/2,678 (7.2%)</td>
<td>630/9,858 (6.4%)</td>
</tr>
<tr>
<td>Hysterectomy</td>
<td>7 studies (Tahseen &amp; Griffiths, 2010)</td>
<td>9/1,747 (0.5%)</td>
<td>51/8,009 (0.6%)</td>
</tr>
</tbody>
</table>

*Calculated by NCC-WCH technical team

Table 11.6 GRADE summary of findings for planned VBAC after ≥ 3 prior CS versus planned repeat CS after ≥ 3 prior CS

<table>
<thead>
<tr>
<th>Number of studies</th>
<th>Number of women</th>
<th>Effect</th>
<th>Quality</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Planned vaginal birth ≥ 3 prior CS</td>
<td>Planned repeat CS ≥ 3 prior CS</td>
<td>Relative (95% CI)</td>
</tr>
<tr>
<td>Blood transfusion</td>
<td>1 study (Cahill et al., 2010)</td>
<td>2/89 (2.2%)</td>
<td>17/771 (2.2%)</td>
</tr>
</tbody>
</table>
1000 (from 21 fewer to 56 more)*

Fever

<table>
<thead>
<tr>
<th>Study</th>
<th>Number of neonates</th>
<th>Relative Risk (95% CI)</th>
<th>Absolute Risk Difference (95% CI)</th>
<th>Quality</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 study (Cahill et al., 2010)</td>
<td>14/89 (15.7%)</td>
<td>121/771 (15.7%)</td>
<td>RR 1.00 (0.60 to 1.67)</td>
<td>Absolute risk difference: 0.3 more per 1000 (from 67 fewer to 93 more)*</td>
</tr>
</tbody>
</table>

Bladder injury rates

<table>
<thead>
<tr>
<th>Study</th>
<th>Number of neonates</th>
<th>Relative Risk (95% CI)</th>
<th>Absolute Risk Difference (95% CI)</th>
<th>Quality</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 study (Cahill et al., 2010)</td>
<td>0/89 (1.6%)</td>
<td>12/771 (15.7%)</td>
<td>Not calculable (NC)</td>
<td>Absolute risk difference: 15 fewer per 1000 (from 27 fewer to 25 more)*</td>
</tr>
</tbody>
</table>

Surgical injury rates

<table>
<thead>
<tr>
<th>Study</th>
<th>Number of neonates</th>
<th>Relative Risk (95% CI)</th>
<th>Absolute Risk Difference (95% CI)</th>
<th>Quality</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 study (Cahill et al., 2010)</td>
<td>0/89 (0.9%)</td>
<td>7/771 (1.6%)</td>
<td>NC</td>
<td>Absolute risk difference: 9 fewer per 1000 (from 18 fewer to 32 more)*</td>
</tr>
</tbody>
</table>

Uterine rupture

<table>
<thead>
<tr>
<th>Study</th>
<th>Number of neonates</th>
<th>Relative Risk (95% CI)</th>
<th>Absolute Risk Difference (95% CI)</th>
<th>Quality</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 study (Cahill et al., 2010)</td>
<td>0/89</td>
<td>0/771</td>
<td>NC</td>
<td>NC</td>
</tr>
</tbody>
</table>

*Calculated by NCC-WCH technical team

Neonatal outcomes

All of the data included in this table have been taken from one systematic review (Guise et al., 2010). The number in the first column indicates the number of studies from that review which report on those outcomes.

Table 11.7 GRADE summary of findings comparing planned CS with planned vaginal birth in women with a previous CS (neonatal outcomes)

<table>
<thead>
<tr>
<th>Number of studies</th>
<th>Number of neonates</th>
<th>Effect</th>
<th>Quality</th>
</tr>
</thead>
<tbody>
<tr>
<td>Perinatal mortality (term)</td>
<td>Planned CS Planned vaginal birth</td>
<td>Relative (95% CI) Absolute (95% CI)</td>
<td></td>
</tr>
<tr>
<td>5 studies (Guise et al., 2010)</td>
<td>46/35,686 (0.12%) 72/41,213 (0.17%)</td>
<td>RR 0.73 (0.51 to 1.06)*</td>
<td>Absolute risk difference: 0.46 less deaths per 1000</td>
</tr>
</tbody>
</table>
### Neonatal mortality (term)

<table>
<thead>
<tr>
<th>Study</th>
<th>Unadjusted Rate</th>
<th>RR (95% CI)</th>
<th>Adjusted Rate</th>
<th>Risk Difference (95% CI)</th>
<th>Quality</th>
</tr>
</thead>
<tbody>
<tr>
<td>6 studies (Guise et al., 2010)</td>
<td>40/63,843 (0.06%)</td>
<td>RR 0.546 (0.36 to 0.82)*</td>
<td>51/44,485 (0.11%)</td>
<td>Absolute risk difference: 0.52 fewer deaths per 1000 (from 0.92 fewer to 0.17 fewer)*</td>
<td>Very low</td>
</tr>
</tbody>
</table>

### Bag and mask ventilation (term)

<table>
<thead>
<tr>
<th>Study</th>
<th>Unadjusted Rate</th>
<th>RR (95% CI)</th>
<th>Adjusted Rate</th>
<th>Risk Difference (95% CI)</th>
<th>Quality</th>
</tr>
</thead>
<tbody>
<tr>
<td>3 studies (Guise et al., 2010)</td>
<td>62/976 (6.3%)</td>
<td>RR 0.39 (0.30 to 0.52)*</td>
<td>183/1134 (16.1%)</td>
<td>Absolute risk difference: 98 fewer per 1000 (Calculated risk difference: 25 fewer per 1000 [from 7.7 fewer to 50 fewer])*</td>
<td>Very low</td>
</tr>
</tbody>
</table>

### Transient Tachypnea (term)

<table>
<thead>
<tr>
<th>Study</th>
<th>Unadjusted Rate</th>
<th>RR (95% CI)</th>
<th>Adjusted Rate</th>
<th>Risk Difference (95% CI)</th>
<th>Quality</th>
</tr>
</thead>
<tbody>
<tr>
<td>3 studies (Guise et al., 2010)</td>
<td>190/1476 (12.9%)</td>
<td>RR 1.04 (0.88 to 1.21)*</td>
<td>427/3451 (12.4%)</td>
<td>Absolute risk difference: 5 more per 1000 (Calculated risk difference: 8.3 more per 1000 [from 33 fewer to 17 more])*</td>
<td>Very low</td>
</tr>
</tbody>
</table>

1. * Calculated by NCC-WCH technical team
2. No pooled data was reported in the systematic review (Guise J.M. et al., 2010) for neonatal intensive care unit [NICU] admission, hypoxic-ischemic encephalopathy (HIE) and neonates’ Apgar score.
3. There were narrative discussions for these outcomes which are summarised here.
4. There was evidence that the rate of NICU admission was higher in neonates born by planned repeat CS compared with neonates born following planned vaginal birth (eight studies, low quality, pooled data not reported).
5. There was very low quality evidence from three studies that reported lower rates of HIE among neonates born by planned repeat CS compared with neonates born following planned vaginal birth (pooled data not reported).
6. There was low quality evidence from four studies that found no difference in the proportion of babies with an Apgar score of 7 or below at 5 minutes in neonates born by planned repeat CS compared with neonates born following a planned vaginal birth (pooled data not reported).
Evidence statements

Maternal outcomes following one CS

Maternal mortality

One systematic review found that the maternal mortality rate was higher in women who had undergone planned repeat CS at term compared with women who had undergone a planned vaginal birth at term. This finding was statistically significant. The evidence for this outcome was of moderate quality.

One systematic review found that the mortality rate was significantly higher in women who had undergone planned repeat CS at any gestational age compared with women who had undergone a planned vaginal birth at any gestational age. This finding was statistically significant. The evidence for this outcome was of moderate quality.

Uterine rupture

One systematic review found that the rate of uterine rupture was lower among women with planned repeat CS at term compared with women who had undergone a planned vaginal birth at term. This finding was statistically significant. The evidence for this outcome was of very low quality.

One systematic review found that the rate of uterine rupture was lower among women with planned repeat CS at any gestational age compared with women who had undergone a planned vaginal birth at any gestational age. This finding was statistically significant. The evidence for this outcome was of very low quality.

Blood transfusion

One systematic review found that the rate of blood transfusion was lower among women who had a planned repeat CS at term when compared with women who had undergone a planned vaginal birth at term. This finding was statistically significant. The evidence for this outcome was of very low quality.

One systematic review found that the rate of blood transfusion was lower among women who had a planned repeat CS at any gestational age when compared with women who had undergone a planned vaginal birth at any gestational age. This finding was statistically significant. The evidence for this outcome was of very low quality.

Hysterectomy

One systematic review did not find a statistically significant difference in the rates of hysterectomy among women who had a planned CS at term compared with women who had undergone a planned vaginal birth at term. The evidence for this outcome was of very low quality.

One systematic review did not find a statistically significant difference in the rates of hysterectomy among women who had a planned CS at any gestational age compared with women who had undergone a planned vaginal birth at any gestational age. The evidence for this outcome was of very low quality.

Infection (endometritis)

One systematic review found that the rate of infection was lower among women with planned repeat CS when compared with women who had a planned vaginal birth. However, the paper did not provide enough data to determine if this difference was statistically significant. The evidence for this outcome was of very low quality.

Length of hospital stay

One systematic review found that the mean length of hospital stay was longer among women with planned repeat CS when compared with women who had a planned vaginal birth. However, the paper did not provide enough data to determine if this difference was statistically significant. The evidence for this outcome was of very low quality.

Postnatal depression

One randomised trial did not find a statistically significant difference in the rates of postnatal depression 6 months postpartum among women who had a planned CS compared with women who had a planned vaginal birth using the Edinburgh Postnatal Depression Scale and Beck Depression Inventory scales. The evidence for this outcome was of very low quality.
Client satisfaction

One randomised trial did not find a statistically significant difference in the rates of client satisfaction 6 months postpartum among women who had a planned CS compared with women who had a planned vaginal birth using Client Satisfaction Questionnaires scores. The evidence for this outcome was of very low quality.

Repeat caesarean sections

No pooled data were reported in the systematic review (Guise et al 2010) for maternal complications associated with repeat CS. There were narrative discussions of each included study for the following outcomes:

1 prior CS versus 2 prior CS

Blood transfusion

One study found that the rate of blood transfusion was lower among women giving birth by CS following one prior CS compared with women who had undergone two prior CS at any gestational age. This finding was statistically significant. The evidence for this outcome was of low quality.

Infection (endometritis)

One study did not find a statistically significant difference in infection rates in women giving birth by CS following one prior CS compared with women who had undergone two prior CS at any gestational age. The evidence for this outcome was of low quality.

Wound complication

One study which found that the wound complication rate was lower among women giving birth by CS following one prior CS compared with women who had undergone two prior CS at any gestational age. This finding was statistically significant. The evidence for this outcome was of low quality.

Surgical injuries

One study found that the rate of surgical injuries was lower among women giving birth by CS following one prior CS at any gestational age when compared with women who had two or more prior CS. This finding was statistically significant. The evidence for this finding was of low quality.

1 prior CS versus ≥ 2 prior CS

Blood transfusion

One study found that fewer women giving birth by CS following one prior CS at any gestational age required a blood transfusion when compared with women who had two or more prior CS. This finding was statistically significant. The evidence for this finding was of low quality.

Hysterectomy

One study did not find a statistically significant difference in the rate of hysterectomy among women giving birth by CS following one prior CS at any gestational age when compared with women who had two or more prior CS. The evidence for this outcome was of low quality.

1 prior CS versus 3 prior CS

Surgical injuries

One study found that the rate of surgical injuries was lower among women giving birth by CS following one prior CS at any gestational age when compared with women who had three prior CS. This finding was statistically significant. The evidence for this outcome was of low quality.

Infection (endometritis)

One study did not find a statistically significant difference in the infection rates among women giving birth by CS following one prior CS compared with women who had undergone three prior CS at any gestational age. The evidence for this outcome was of low quality.

Wound complication

One study did not find a statistically significant difference in the wound complication rates among women giving birth by CS following one prior CS compared with women who had undergone three prior CS at any gestational age. The evidence for this outcome was of low quality.
Planned VBAC after 2 prior CS versus planned repeat CS after 2 prior CS

Blood transfusion
One systematic review did not find a statistically significant difference in the rate of blood transfusion among women who planned a vaginal birth following two prior CS compared with women who had planned CS following two prior CS. The evidence for this outcome was of very low quality.

Febrile morbidity
One systematic review did not find a statistically significant difference in the rate of febrile morbidity among women who planned a vaginal birth following two prior CS compared with women who had planned CS following two prior CS. The evidence for this outcome was of very low quality.

Hysterectomy
One systematic review did not find a statistically significant difference in the rate of hysterectomy in women who planned a vaginal birth following two prior CS compared with women who had planned CS following two prior CS. The evidence for this outcome was of very low quality.

Planned VBAC after ≥ 3 prior CS versus planned repeat CS after ≥ 3 prior CS

Blood transfusion
One study did not find a statistically significant difference in the blood transfusion rates among women who planned a vaginal birth following three or more prior CS compared with women who had planned CS following three or more prior CS. The evidence for this outcome was of very low quality.

Fever
One study did not find a statistically significant difference in fever rates in women who planned a vaginal birth following three or more prior CS compared with women who had planned CS following three or more prior CS. The evidence for this outcome was of very low quality.

Bladder injury
One study did not find a statistically significant difference in the rate of bladder injuries among women who planned a vaginal birth following three or more prior CS compared with women who had an planned CS following three or more prior CS. The evidence for this outcome was of very low quality.

Surgical injury
One study did not find a statistically significant difference in the rate of surgical injuries among women who planned a vaginal birth following three or more prior CS compared with women who had planned CS following three or more prior CS. The evidence for this outcome was of very low quality.

Uterine rupture
One study did not find a statistically significant difference in the rate of uterine rupture in women who planned a vaginal birth following three or more prior CS compared with women who had an planned CS following three or more prior CS. The evidence for this outcome was of very low quality.

Neonatal outcomes

Perinatal mortality
One systematic review did not find a statistically significant difference in the perinatal mortality rate among infants born to women who planned a repeat CS at term compared with infants born at term to women who planned a vaginal birth. The evidence for this outcome was of very low quality.

Neonatal mortality
One systematic review found that the neonatal mortality rate was lower for infants born at term to women who planned a repeat CS compared with infants born at term to women who planned a vaginal birth. This finding was statistically significant. The evidence for this outcome was of very low quality.

Bag and mask ventilation
One systematic review found that the use of bag and mask ventilation was lower among neonates born at term to women who planned a repeat CS compared with neonates born at term to women who planned a vaginal birth. This finding was statistically significant. The evidence for this outcome was of very low quality.
Transient tachypnea
One systematic review did not find a statistically significant difference in the incidence of neonatal transient tachypnoea between neonates born to women who planned a repeat CS and those born at term to women who planned a vaginal birth. The evidence for this outcome was of very low quality.

Health economics
A model was developed to compare the cost-effectiveness of VBAC (a trial of labour) versus a planned caesarean section in women with one previous caesarean section and with no plans for further children. A summary of this analysis is provided below (see Chapter 13. for further details).
In addition to the costs of birth the model also estimated “downstream” costs and QALYs based on the risk of adverse events for each planned mode of birth. The base case analyses considered only the outcomes that were reported in the review undertaken for this guideline to determine the risks and benefits of planned caesarean section compared with planned vaginal birth for both women and babies in women who have had a previous caesarean section. Secondary analyses, in addition, used outcomes that were only reported in the review which compared the risk and benefits of planned caesarean section compared with planned vaginal birth. However, it should be recognised that these risks are likely to be underestimated for this population and that the relative risk for these adverse outcomes may also be different in this population.

The results tended to show that VBAC was more likely to be cost-effective although this was a borderline finding and considerable uncertainty remains especially with respect to all the outcomes that may differ between the different modes of planned birth in this population.

Evidence to recommendations
Relative value placed on outcomes considered
It was noted that findings from studies of babies born at term were very similar to those that included babies born at any gestational age. This was thought to reflect the relatively low numbers of preterm babies included in the studies involving all gestational ages. The extra statistical power afforded by the larger numbers where studies of all gestational ages have been included meant the GDG were happy to consider this evidence when making recommendations.

Whilst maternal mortality is clearly a vitally important outcome at an individual level, in terms of informing decision-making for a whole population, the very low numbers of deaths reported in the studies (absolute numbers range from 1.9 to 7.5 per 100,000) mean this outcome does not necessarily drive the recommendations made. Other important outcomes including uterine rupture and neonatal mortality were more common, although still rare, meaning that although differences between study groups were statistically significant, the low incidence meant that they were also considered less clinically significant in terms of driving clinical practice and advice to women. In the context of low actual risk, then absolute risk will be a more important consideration than relative risk.
The reported neonatal outcomes of bag and mask ventilation and transient tachypnoea were felt to be of limited value as it was not possible to determine how these outcomes related to ongoing health problems or disability.

Trade-off between clinical benefits and harms
Whilst mortality-related outcomes are very rare it was noted that, for women planning birth following one previous CS, maternal mortality is statistically significantly higher for women planning a repeat CS whilst neonatal mortality is statistically significantly higher for women planning a vaginal birth. Perinatal mortality is not statistically different between the two groups.
The GDG agreed that whilst it is right to give women all available information when planning mode of birth, for women planning birth after one previous CS, the important thing to underline is that serious adverse outcomes, including maternal and neonatal mortality, uterine rupture, need for blood transfusion and hysterectomy are very rare, no matter whether planned repeat CS or VBAC is chosen.
It was noted that the relative risk of adverse outcomes may vary from woman to woman depending upon her obstetric history, including reasons for previous CS and whether or not a woman has
previously given birth vaginally. These individual considerations need to be taken into account when discussing mode of birth following previous CS.

When considering increasing numbers of previous CS the evidence showed no difference between planned vaginal birth and planned CS in rates of blood transfusion, fever and hysterectomy after two prior CS.

With an increasing number of CS, there is an increasing risk of need for blood transfusion, wound complications and injuries to the bladder, regardless of the mode of birth.

Trade-off between net health benefits and resource use

An economic model developed for this guideline to compare the cost effectiveness of planned CS versus planned vaginal birth in women who have had a previous CS did not strongly suggest a preferred mode of birth. As a result this model would, given the current state of evidence, support a recommendation allowing women to choose their preferred method of birth in consultation with the health care professionals responsible for her care. Considerations about any future pregnancies may be an important factor in the decisions made given the increased risks e.g. in incidence of placenta praevia and morbidly adherent placenta, associated with repeat CS.

Quality of evidence

The evidence for outcomes following one previous CS was drawn from one large meta-analysis of observational studies and one RCT and ranged from moderate to very low. The large sample sizes reported for the meta-analysis meant the GDG felt more confident in the validity of the findings regarding rare outcomes.

The evidence examining outcomes following more than one previous CS is of low and very low quality. While evidence comparing outcomes for women having repeat CS following 1, 2 or more previous CS is interesting and can be used to provide general information about increasing risks following 2 or more CS it does not help a woman decide the level of risk associated with her choice of mode of birth in the current pregnancy. However, this information is helpful for decision making about future births. The evidence comparing outcomes for women choosing a planned vaginal birth vs. those choosing a planned CS after 2 or more previous CS is useful in this respect as this reflects the choice women have. Unfortunately this evidence is of very low quality, thus lowering the validity of the reported findings.

No good quality evidence was available for women planning birth following 5 or more previous CS.

Other considerations

The GDG noted that many women leave hospital following a caesarean birth without understanding the implications for planning future pregnancies and births. It was felt that it is important to provide this information to women and their partners so that they can have an accurate picture of what this means for them planning their family, including options for future modes of birth. The GDG agreed that there is a benefit to providing this information to women and their partners prior to leaving the hospital because the medical records are easily available to refer to. As a result, the group recommended that this discussion take place with women after the CS. However, they also recognised that some women may prefer to have this discussion at a later date so highlighted that this discussion can be deferred. Due to the large amount of information women and their partners receive during the immediate postnatal period this information should be provided both verbally and in written formats. It is important to emphasise that, regardless of future choice of mode of birth, poor outcomes are very rare.

<table>
<thead>
<tr>
<th>Number</th>
<th>Recommendation</th>
</tr>
</thead>
<tbody>
<tr>
<td>119</td>
<td>When advising about the mode of birth after a previous CS consider:</td>
</tr>
<tr>
<td></td>
<td>• maternal preferences and priorities</td>
</tr>
<tr>
<td></td>
<td>• the risks and benefits of repeat CS</td>
</tr>
<tr>
<td></td>
<td>• the risks and benefits of planned vaginal birth after CS, including the risk of unplanned CS. [new 2011]</td>
</tr>
</tbody>
</table>
Inform women who have had up to and including four CS that the risk of fever, bladder injuries and surgical injuries does not vary with planned mode of birth and that uterine rupture is very rare. [new 2011]

Offer women planning a vaginal birth who have had a previous CS:

- electronic fetal monitoring during labour
- care during labour in a unit where there is immediate access to CS and on-site blood transfusion services. [GPP] [2011]

During induction of labour, women who have had a previous CS should be monitored closely, with access to electronic fetal monitoring and with immediate access to CS, because they are at increased risk of uterine rupture. [GPP [2004]

Pregnant women with both previous CS and a previous vaginal birth should be informed that they have an increased likelihood of achieving a vaginal birth than women who have had a previous CS but no previous vaginal birth. [B] [2004]

While women are in hospital after having a CS, give them the opportunity to discuss with healthcare professionals the reasons for the CS and provide both verbal and printed information about birth options for any future pregnancies. If the woman prefers, provide this at a later date. [new 2011]

For recommendations on methods of induction for women who have had a previous CS, see the ‘Induction of labour’ guideline (NICE, 2008)

<table>
<thead>
<tr>
<th>Number</th>
<th>Research Recommendation</th>
</tr>
</thead>
<tbody>
<tr>
<td>RR 46</td>
<td>A comparison of the long term psychological and physical outcomes between women who have chosen and/or been advised towards a VBAC or a planned repeat CS.</td>
</tr>
<tr>
<td>RR 47</td>
<td>An evaluation of the effectiveness of continuity of carer on the proportion of women planning and achieving a VBAC, and the short and long term psychological and physical outcomes of women following a planned VBAC.</td>
</tr>
</tbody>
</table>
12 Auditable standards

Table 12.1 Suggested audit criteria

<table>
<thead>
<tr>
<th>Criterion</th>
<th>Exception</th>
<th>Definition of terms</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Making the decision</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Percentage of women having CS that have a documented discussion on benefits and risks of CS compared with vaginal birth specific to the woman and her pregnancy</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Percentage of women requesting a CS that have a documented discussion on the reasons for the request</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Carrying out the procedure</strong></td>
<td>Percentage of CS carried out using a regional block</td>
<td>Regional block – spinal or epidural anaesthesia</td>
</tr>
<tr>
<td>Percentage of CS where the woman receives prophylactic antibiotics</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Percentage of CS where an appropriate method of thromboprophylaxis is used</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Percentage of CS where antacids are given prior to regional or general anaesthesia</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Percentage of CS where antiemetics are given prior to regional or general anaesthesia</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Percentage of planned CS carried out after 39 weeks</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Reducing the likelihood of CS</strong></td>
<td>Percentage of women who have an uncomplicated singleton breech pregnancy at 36 weeks’ gestation that have a documented offer of external cephalic version</td>
<td>Specific clinical indications</td>
</tr>
<tr>
<td>Hospitals should measure the overall CS rate as well as the percentage of CS performed for the four major determinants (presumed fetal compromise, failure to progress in labour, breech presentation, multiple pregnancy) and ‘maternal request’</td>
<td>Women in labour, women with a uterine scar or abnormality, fetal compromise, ruptured membranes, vaginal bleeding and medical conditions</td>
<td></td>
</tr>
<tr>
<td>Percentage of women in labour that have continuous support during labour, provided by women with or without prior training, for example, doulas, childbirth educators or a female relative.</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Percentage of women with uncomplicated pregnancies beyond 41 weeks with documented offer of induction of labour

Percentage of women in spontaneous labour with an uncomplicated singleton pregnancy at term monitored using a partogram with a 4-hour action line

Percentage of documented involvement of consultant obstetricians in the decision making for CS

Percentage of CS for abnormal fetal heart rate pattern, suspected fetal acidosis, in which fetal blood sampling is undertaken

Partogram – graphic representation of labour progress

Women not having a CS

Severely abnormal fetal heart rate pattern

Contraindications to fetal blood sampling
13 Health Economics

13.1 Introduction

The aims of the health economic input to the guideline were to inform the GDG of potential economic issues relating to caesarean section and to ensure that recommendations represented a cost-effective use of healthcare resources. Health economic evaluations aim to integrate data on benefits or harms (ideally in terms of quality adjusted life years [QALYs]) and costs of different care options.

The GDG prioritised the clinical questions where it was thought that economic considerations would be particularly important in formulating recommendations. For this guideline the areas prioritised for economic analysis were:

- Diagnosis of morbidly adherent placenta (see section 5.6 for summary and 13.2 for full details)
- Maternal request for caesarean section (see section 5.9 for summary and 13.3 for full details)
- Vaginal birth after caesarean section (see section 11.2 for summary and 13.4 for full details)

13.2 Cost effectiveness of diagnosis of morbidly adherent placenta

Introduction

In women with a previous caesarean section there is an increased risk of placenta praevia and this risk increases with the number of previous caesarean sections (Clark et al., 1985). In turn, women with placenta praevia from a previous caesarean section are at risk of a morbidly adherent placenta. Although this risk is small it increases with the number of previous caesarean sections (Silver et al. 2006). In addition to an increased maternal mortality risk, a morbidly adherent placenta also can lead to excessive blood loss, the need for a hysterectomy and surgical complications.

Practice for the diagnosis of a morbidly adherent placenta is not consistent across England and Wales. Furthermore, there is uncertainty about the accuracy of imaging techniques used to diagnose a morbidly adherent placenta and further uncertainty as to whether a diagnosis using these imaging techniques leads to improved outcomes. As a result this was considered an important issue for the update of the guideline.

A systematic review of the literature didn’t identify any papers addressing the cost-effectiveness of diagnostic imaging for morbidly adherent placenta. Therefore, a new model was developed for the purposes of this guideline.

Method

A decision analytic model was developed in Microsoft Excel® to compare the cost effectiveness of the following diagnostic strategies for morbidly adherent placenta.

i) None

ii) Ultrasound

iii) MRI

iv) Ultrasound followed by MRI in ultrasound test positives

The basic structure of the model is illustrated in Figure 13.1. In assessing the cost-effectiveness of these diagnostic strategies it is important not to overlook treatment as the two are highly interdependent. For example, a very effective and inexpensive treatment may not ultimately be cost-
effective if the costs of identifying patients who could benefit are prohibitively high. Similarly, a very accurate and cheap diagnostic test may not be worth doing if it has no bearing on patient outcomes. However, the clinical review did not identify any evidence of the relationship between a diagnosis of morbidly adherent placenta and patient outcomes. The GDG, however, thought there were likely to be advantages in terms of “being prepared”. These advantages could be in terms of “downstream” cost savings and possible improved outcomes for mother and baby. In addition the GDG considered there was a benefit to the mother in being prepared for some of the likely outcomes of her pregnancy and birth. In the absence of evidence to quantify these benefits this model adopts a “what-if” approach to determine the thresholds for cost-effectiveness.

Figure 13.1: The model structure

This analysis was undertaken from the perspective of the NHS and personal social services which is in accordance with NICE guidelines methodology (NICE, 2009). Costs and benefits are compared using standard methods of incremental analysis of costs and benefits. Costs were based on 2009/10 prices. A number of sensitivity analyses were undertaken to assess the importance of parameter uncertainty within the model.

Model inputs
The default model input values are shown in Table 13.1 to Table 13.3. Based on GDG opinion the costs of “being prepared”, in the event of a diagnosis of morbidly adherent placenta, would typically be two hours of a consultant anaesthetist’s time and having an additional four units of cross matched blood available. Diagnostic accuracy data was taken from the literature that was retrieved as part of the systematic review that was undertaken for the guideline.
The use of “adverse outcome” in this context of the model is purposefully vague given that it is not known to what extent having a diagnosis and “being prepared” leads to better outcomes. However, it is intended to capture the idea that treatment, which in this case is “being prepared”, could mitigate any adverse outcomes relative to a state of not being prepared.

The probabilities of an adverse outcome, with and without “being prepared”, are illustrative values. These values are used to provide a “what-if” with respect to treatment effectiveness. The QALY loss associated with an adverse outcome and the costs associated with an “adverse outcome” are set to zero in the model’s default setting but the impact of relaxing this assumption on model outcomes can readily be observed.

**Table 13.1 Model costs**

<table>
<thead>
<tr>
<th>Item</th>
<th>Value</th>
<th>Source</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ultrasound</td>
<td>£55</td>
<td>NHS Reference Costs 2009-10 Outpatient</td>
<td></td>
</tr>
<tr>
<td>MRI</td>
<td>£175</td>
<td>NHS Reference Costs 2009-10 Outpatient</td>
<td></td>
</tr>
<tr>
<td>Being prepared</td>
<td>£800</td>
<td><a href="http://www.hta.ac.uk/fullmono/mon1044.pdf">http://www.hta.ac.uk/fullmono/mon1044.pdf</a></td>
<td>Unit of blood £111.16 x 4 Matching £23.24</td>
</tr>
<tr>
<td>“Adverse outcome”</td>
<td>£0</td>
<td>n/a</td>
<td>Can be varied as part of a “what-if” analysis</td>
</tr>
</tbody>
</table>

**Table 13.2 Probabilities**

<table>
<thead>
<tr>
<th>Item</th>
<th>Value</th>
<th>Source</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Prevalence</td>
<td>3%</td>
<td>Silver et al., (2006)</td>
<td>In women with placenta praevia, the risk for placenta accreta was 3% for first repeat caesarean deliveries</td>
</tr>
<tr>
<td>Adverse outcome/prepared</td>
<td>10%</td>
<td>n/a</td>
<td>illustrative value</td>
</tr>
<tr>
<td>Adverse outcome/not prepared</td>
<td>40%</td>
<td>n/a</td>
<td>illustrative value</td>
</tr>
<tr>
<td>Ultrasound sensitivity</td>
<td>92%</td>
<td>Shih et al.(2009)</td>
<td>US Colour Doppler</td>
</tr>
<tr>
<td>Ultrasound specificity</td>
<td>68%</td>
<td>Shih et al.(2009)</td>
<td>US Colour Doppler</td>
</tr>
<tr>
<td>MRI sensitivity</td>
<td>100%</td>
<td>Masselli et al. (2008)</td>
<td></td>
</tr>
<tr>
<td>MRI specificity</td>
<td>100%</td>
<td>Masselli et al. (2008)</td>
<td></td>
</tr>
</tbody>
</table>

**Table 13.3 Cost-effectiveness**

<table>
<thead>
<tr>
<th>Item</th>
<th>Value</th>
<th>Source</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Willingness to pay for a</td>
<td>£20,000</td>
<td>NICE</td>
<td>manual</td>
</tr>
</tbody>
</table>
Results

The results using the base case inputs are shown in Table 13.4 and Figure 13.2. It must be remembered that there is an element of “what-if” in these results with some of the inputs being hypothetical illustrative values. These results essentially show the diagnostic costs of the respective strategies and do not consider any “downstream” savings that might arise as a result of being prepared or any health gains to the mother and baby.

Table 13.4 Results

<table>
<thead>
<tr>
<th>Strategy</th>
<th>Cost</th>
<th>Incremental cost</th>
</tr>
</thead>
<tbody>
<tr>
<td>Do Nothing</td>
<td>£0</td>
<td>-</td>
</tr>
<tr>
<td>Ultrasound+MRI</td>
<td>£136</td>
<td>£136</td>
</tr>
<tr>
<td>MRI</td>
<td>£199</td>
<td>£63</td>
</tr>
<tr>
<td>Ultrasound</td>
<td>£325</td>
<td>£126</td>
</tr>
</tbody>
</table>

Figure 13.2 Results with base case values

Sensitivity Analysis

i. Changing prevalence

There is less uncertainty surrounding the prevalence than there is for other parameter values. However, showing the impact of a different prevalence gives insights into the drivers of the model’s results. Figure 13.3 shows the results for a prevalence of 60% holding other model values constant at their base case values.
Here two “what-ifs” are explored, first if the “downstream” costs of an adverse outcome are assumed to be £5,000 and second if those costs were assumed to be £30,000. The results of this are shown in Figures 13.4 and 13.5 respectively.

**Figure 13.3** Results with prevalence set to a hypothetical 60%

**Figure 13.4** Results with “Adverse outcomes” set to £5,000
iii. Assuming an “adverse outcome” has QALY implications

In this analysis we assess the cost-effectiveness assuming that averting “adverse outcomes” has a QALY gain for the mother and/or baby. All other inputs in the model are held constant at their base case level, including the assumption that there are no “downstream” costs associated with an adverse outcome. Table 13.5 shows the results for a QALY loss per adverse outcome of 0.02 and Table 13.6 shows the results when a much greater QALY loss of 5.00 is assumed.

<table>
<thead>
<tr>
<th>Strategy</th>
<th>Cost</th>
<th>Incremental</th>
<th>QALY</th>
<th>Incremental</th>
<th>ICER</th>
</tr>
</thead>
<tbody>
<tr>
<td>Do nothing</td>
<td>£0</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Ultrasound + MRI</td>
<td>£136</td>
<td>£136</td>
<td>0.00011</td>
<td>0.00011</td>
<td>£822,645</td>
</tr>
<tr>
<td>MRI</td>
<td>£199</td>
<td>£63</td>
<td>0.00012</td>
<td>0.00001</td>
<td>£4,359,028</td>
</tr>
<tr>
<td>Ultrasound</td>
<td>£325</td>
<td>£126</td>
<td>0.00011</td>
<td>-0.00001</td>
<td>Dominated</td>
</tr>
</tbody>
</table>

Given the reported incremental cost then a strategy of ultrasound + MRI would have to generate 0.0068 QALYs per pregnancy for this to be cost-effective relative to ‘do nothing’. Similarly, MRI alone would have to produce an additional 0.0031 QALYs per woman compared do ultrasound + MRI in order to be considered cost-effective relative to that strategy.

<table>
<thead>
<tr>
<th>Strategy</th>
<th>Cost</th>
<th>Incremental</th>
<th>QALY</th>
<th>Incremental</th>
<th>ICER</th>
</tr>
</thead>
<tbody>
<tr>
<td>Do nothing</td>
<td>£0</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Ultrasound + MRI</td>
<td>£136</td>
<td>£136</td>
<td>0.0276</td>
<td>0.0276</td>
<td>£3,291</td>
</tr>
</tbody>
</table>
A threshold analysis was undertaken which showed that the QALY gain would have to be 0.8 or greater for ultrasound + MRI to be cost-effective relative to ‘do nothing’. A further threshold analysis found that MRI alone would be the most cost-effective strategy, at a willingness to pay threshold of £20,000 per QALY, if the QALY loss associated with an “adverse outcome” was 4.4. It should be remembered though that these threshold results are for when other model inputs are held constant at their base case value.

iv. Two-way sensitivity analysis varying the QALY loss and cost of an “adverse outcome”

Two very important unknowns in the model are the QALY loss from an “adverse outcome” and the “downstream” costs arising from them. In this analysis the QALY loss from an “adverse outcome” is varied between 0.0 and 10.0 and the “downstream” cost of an “adverse outcome” is varied between £0 and £10,000. Figure 13.6 shows the thresholds at which a strategy of Ultrasound+MRI is cost-effective relative to ‘do nothing’ for different QALY and “downstream” cost combinations. Figure 13.7 shows the thresholds for determining the cost-effectiveness of MRI alone relative to Ultrasound+MRI.

Figure 13.6 Cost-effectiveness thresholds for Ultrasound+MRI relative to ‘do nothing’

Figure 13.7 Cost-effectiveness thresholds for MRI relative to Ultrasound+MRI
v. Two-way sensitivity analysis varying the QALY loss and cost of an “adverse outcome” assuming a lower effectiveness from “being prepared”

In the previous analysis, the effectiveness of “being prepared” was assumed to be that used in the base-case analysis. However, there is a lack of evidence as to whether “being prepared” does lead to such a reduction in “adverse outcomes”. In this analysis we assume that the risk of an “adverse outcome” in a case of morbidly adherent placenta is 20% if not prepared as against a risk of 10% if prepared. In other words we assume here that correctly identifying cases has a smaller benefit in terms of averting “adverse outcomes”. The “downstream” cost of an “adverse outcome” is varied between £0 and £100,000 and the QALY loss from an “adverse outcome” is varied between 0.0 and 25.0. This analysis for Ultrasound+MRI relative to ‘do nothing’ is displayed in Figure 13.8 and for MRI alone relative to Ultrasound+MRI in Figure 13.9.

**Figure 13.8** Cost-effectiveness thresholds for Ultrasound+MRI relative to “do nothing” assuming that the risk of an “adverse outcome” when not prepared is 20%

**Figure 13.9** Cost-effectiveness thresholds for MRI relative to Ultrasound+MRI assuming that the risk of an “adverse outcome” when not prepared is 20%

**Discussion**

Caution needs to be exercised in interpreting the results of this analysis given the “what-if” approach. Nevertheless, it does give some insights into the cost-effectiveness of different diagnostic strategies.
for morbidly adherent placenta. First, there is good evidence on the prevalence of morbidly adherent placenta in women with placenta praevia. The prevalence is quite low and therefore the false positive rate is likely to be an important determinant of cost-effectiveness. As a result the strategy of ultrasound was dominated in most of the above analyses. This was because it had a relatively high false positive rate, which meant lower diagnostic costs were more than offset by higher “preparedness” costs, and its lower sensitivity meant that fewer “adverse outcomes” were prevented than in other diagnostic strategies. Only when the prevalence was set to an unrealistically high level, did ultrasound alone become the cheapest diagnostic strategy.

The results also suggest a potential advantage of the sequential Ultrasound+MRI strategy compared to MRI alone. By testing ultrasound positives with MRI to confirm the diagnosis, the false positives are removed. The additional costs of the MRI test are more than offset by the reduction in unnecessary “preparedness”. On the other hand the actual number of MRI tests undertaken is much less than occurs with a strategy based on MRI alone and because of the substantial difference in costs between an Ultrasound and MRI this means that the sequential strategy has markedly lower diagnostic costs even if the absolute number of tests undertaken is higher. Furthermore, although true positives are missed because of the lower sensitivity of ultrasound compared to MRI, the low prevalence and the fact that the ultrasound detects 92% of cases means that the absolute difference in missed cases is very small. Therefore, for MRI alone to be cost-effective “being prepared” would have to substantially reduce the risk of “adverse outcomes” and/or there would have to be large QALY losses and “downstream” costs associated with “adverse comes” in order to make the higher costs worthwhile. This is well illustrated in Figures 13.7 and 13.9. Figures 13.8 and 13.10 show that a much lower QALY gain and “downstream” saving is necessary for Ultrasound+MRI to be considered cost-effective relative to “do nothing”. Of course, this is predicated on the point estimates of test accuracy used in the model being a good approximation of their ‘true’ value. Sensitivity analysis was not undertaken on diagnostic test parameters as these inputs had at least some evidential basis and this was considered very much a second order level of uncertainty compared to the effectiveness of “being prepared”, the QALY loss and “downstream” cost of “adverse outcomes. Were the diagnostic accuracy values found to be substantially different then the “what-if” results would all differ. Nevertheless, it should be remembered that the “what-if” analyses do have some uncertainty associated with them for this reason.

Figure 13.5 and Figure 13.6 suggest that the model output is not that sensitive to the costs of an “adverse outcome”. This is principally due to the low prevalence of morbidly adherent placenta. Nevertheless, even at low prevalence, Tables 13.5 and 13.6 suggest that the QALY gained from an averted “adverse outcome” is likely to be an important determinant of cost-effectiveness. Clearly, any QALY gain also depends on the effectiveness of “being prepared” in averting “adverse outcomes”.

**Conclusion**

An absence of evidence on a number of key parameters means that firm conclusions about the cost-effectiveness of different diagnostic strategies cannot be reached from this model. However, the model does use a “what-if” approach to show the scenarios in which different strategies would be cost-effective. Clearly, if “being prepared” confers no benefit to mother and child, then no diagnostic strategy is likely to be cost-effective unless society places a very high value on providing information to the woman. At this stage there is no evidence to quantify the extent to which “being prepared” does improve outcomes but the expert and consensus opinion of the GDG is that some improvement in outcomes is likely to result from “being prepared”. However, even if “being prepared” does confer a benefit, that is a necessary but not sufficient condition for diagnosis to be cost-effective. In addition, the benefit must be worth the additional costs involved in which case advertised “adverse outcomes” must also yield a sufficiently large QALY gain and/or “downstream” saving. Again, this is an unknown but although current UK practice varies, it does involve diagnostic strategies to diagnose morbidly adherent placenta. Furthermore, it is plausible to anticipate that there would be some QALY gain and “downstream” saving from averting “adverse outcomes”. Therefore, the GDG recommendation of the Ultrasound+MRI strategy seems reasonable based on this analysis. Ultrasound alone is likely to be dominated because of the high false positive rate associated with a test specificity of 68%. On the other hand MRI alone is more expensive than a sequential test strategy and sensitivity analysis suggested large treatment effect sizes and QALY losses and “downstream” saving from “adverse outcomes” would be necessary to justify the additional.
diagnostic and “preparedness” cost. Furthermore, there could be capacity constraints in recommending this strategy and therefore such a recommendation would be difficult to justify in the absence of good cost-effectiveness evidence. On the other hand, much smaller benefit from diagnosis is required for Ultrasound+MRI to be cost-effective relative to no diagnosis. Therefore, whilst further evidence is required, a recommendation of Ultrasound+MRI seems to make pragmatic sense given current practice, GDG opinion and the insights available from this “what-if” modelling approach.

13.3 Cost effectiveness of planned vaginal birth versus maternal request caesarean section

Introduction

Many developed countries have experienced rising rates of caesarean section. In England caesarean section rates have increased from 9.0% in 1980 to 24.6% in 2008-09 (Bragg et al., 2010). Whilst some of the change is likely to be explained by changes in the case-mix of women giving birth – increasing maternal age at first pregnancy for example – and improvements in technology making the operation safer, there is also evidence of a dramatic increase in caesarean section rates among women with no indicated medical risk (Zupancic, 2008). A number of commentators have expressed concern with the economic implications of these trends and the issue is potentially an important one given scarce resources (Zupancic, 2008; Druzin and El-Sayed, 2006). Data from NHS Reference Costs shows that caesarean section is generally more expensive than vaginal birth. This is consistent with other cost comparisons of different modes of birth (Allen et al., 2005; Druzin and El-Sayed, 2006). Furthermore, caesarean section is not without risk and it is frequently suggested that it leads to worse maternal and infant outcomes in the current pregnancy and in any subsequent pregnancies. If caesarean section typically has higher cost and worse outcomes, then caesarean section without any indicated medical risk may indicate an inefficient use of resources. In this case it might be reasonably argued that maternal request for caesarean section in the absence of medical indication shouldn’t be routinely granted in a publicly funded health care system, with the opportunity costs implied.

However, there are others who have argued that the cost-effectiveness issue is perhaps not as straightforward. First, comparisons of planned caesarean section and planned vaginal birth often do not exclude those with an obstetric indication for caesarean section. Such comparisons are therefore not done on a like-for-like basis and it is of course to be expected that the subset with indications for caesarean section are likely to experience more complications and concomitant costs, as they are inherently higher risk pregnancies. Second, unplanned caesarean section is more expensive than “elective” caesarean and this is a relatively common occurrence for planned vaginal birth but unusual when caesarean section is planned. Furthermore, unplanned caesarean section has the worst maternal and infant outcomes of all modes of birth. Normal vaginal birth is not risk free and some adverse outcomes, such as urinary incontinence, occur more frequently in women who give birth vaginally (Thom & Rortveit, 2010).

A literature search identified two primary research papers which evaluated the cost-effectiveness of maternal request caesarean section (Xu et al., 2010; Culligan et al., 2005), although there were a number of other studies comparing costs. One US study (Xu et al., 2010) used a Monte Carlo simulation decision model to compare the cost-effectiveness of maternal request caesarean section to a trial of labour (TOL) in primigravid women without a medical or obstetric indication for caesarean section. In particular the authors stated that they wanted to consider the lifetime management of pelvic floor disorders. The model was restricted to women having a single lifetime birth (accounting for 21.6% of parous women in the United States) and did not consider any impact the mode of birth for the primigravid pregnancy may have on subsequent pregnancies. In addition to pelvic floor complication, the analysis also included other maternal and perinatal mortalities. Costs were calculated using a societal perspective. This study concluded that their evaluation did not reveal a clearly preferred mode of birth based on cost-effectiveness analysis, although they also reported that when compared to a trial of labour, the probability of maternal request caesarean section being cost-effective was 82% using a societal willingness to pay for a QALY threshold of $50,000.
Another US study (Culligan et al., 2005) used a decision analysis to evaluate planned caesarean section to prevent anal incontinence and brachial plexus injuries associated with macrosomia. A population at risk of a macrosomic baby was identified using an ultrasound at a gestational age of 39 weeks. The authors argued that such a policy would be cost-effective, producing cost savings and net QALY gains.

Neither of these studies alone or in combination was deemed adequate to make recommendations for the NHS. First, one of the studies (Culligan et al., 2005) focuses on a population with an obstetric indication and in that sense is not relevant to the question being addressed. Second, both are US studies and it is well recognised that treatment costs differ, often substantially, between the UK and US. Third, whilst one study (Xu et al., 2010) was based on published literature the evidence was not clearly retrieved in a systematic view and unlike this guideline, reviewed outcomes were not based on planned mode of birth. Nor did it adopt a health service cost perspective which might have been consistent with the NHS and personal social services perspective recommended in the NICE guidelines manual (NICE, 2009).

Therefore a model was developed for this guideline, to assess the cost-effectiveness of planned vaginal birth versus planned caesarean section in primiparous women without an obstetric indication for caesarean section.

**Method**

A cost utility analysis was undertaken using a decision analytic model developed in Microsoft Excel® to compare the cost effectiveness of planned vaginal birth versus planned caesarean section in England and Wales. The population modelled was women without any obstetric indication for caesarean section and not having had a previous caesarean section. The intention is that the population does not differ systematically by mode of birth if vaginal birth is not contraindicated for women having planned caesarean section. The basic structure of the model is illustrated in Figure 13.10. As well as considering the costs of modes of birth, we also aim to evaluate the “downstream” impact on costs and health related quality of life arising from adverse events.

**Figure 13.10:** The model structure
This analysis was undertaken from the perspective of the NHS and personal social services which is in accordance with NICE guidelines methodology (NICE, 2009). Costs and benefits are compared using standard methods of incremental analysis of costs and benefits. Costs were based on 2009/10 prices. A number of sensitivity analyses were undertaken to assess the impact that changes in the base case assumptions would have on the model’s results.

The cost of method of birth

The cost of actual vaginal birth, planned caesarean section and unplanned caesarean section was calculated using NHS Reference Costs 2009/10. Data on the number of cases were used to calculate a mean weighted cost for each mode of birth. The weighted mean costs were £1,512 for an actual vaginal birth, £2,369 for an actual planned vaginal birth and £3,042 for an unplanned section as shown in Tables 13.7 – 13.9. However, the costs of a planned mode of birth will depend on the actual mode of birth, see Table 13.10. For example, it is estimated that 10% of planned vaginal births will result in an unplanned caesarean section.

A weighted mean cost for each planned mode of birth is then derived according to the relative proportion of different modes of birth occurring for each planned method, see Table 13.11. It should be noted that the costs of caesarean section will be based on all women having caesarean section, including those with an obstetric indication. This may influence both the weights applied to different types of caesarean section delivery and also their costs. However, good quality UK cost data for caesarean section performed solely on the basis of maternal request is not currently available as far as we are aware.

Table 13.7: Vaginal birth costs

<table>
<thead>
<tr>
<th>Category</th>
<th>Birth details</th>
<th>Cases</th>
<th>Cost</th>
<th>Weighted Cost</th>
</tr>
</thead>
<tbody>
<tr>
<td>Elective inpatient</td>
<td>Normal birth with complications (CC)</td>
<td>158</td>
<td>£1,558</td>
<td>£0.51</td>
</tr>
<tr>
<td>Elective inpatient</td>
<td>Normal birth without CC</td>
<td>1,101</td>
<td>£1,151</td>
<td>£2.61</td>
</tr>
<tr>
<td>Elective inpatient</td>
<td>Normal birth with epidural with CC</td>
<td>13</td>
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<td>£0.05</td>
</tr>
<tr>
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<td>Normal birth with epidural without CC</td>
<td>51</td>
<td>£1,260</td>
<td>£0.13</td>
</tr>
<tr>
<td>Elective inpatient</td>
<td>Normal delivery with induction with CC</td>
<td>740</td>
<td>£2,109</td>
<td>£3.22</td>
</tr>
<tr>
<td>Elective inpatient</td>
<td>Normal birth with induction without cc</td>
<td>2,503</td>
<td>£1,306</td>
<td>£6.74</td>
</tr>
<tr>
<td>Elective inpatient</td>
<td>Normal birth with post-partum surgical intervention</td>
<td>173</td>
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<td>Assisted birth with epidural with CC</td>
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<td>Assisted birth with epidural without CC</td>
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<td>£1.50</td>
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<td>£2,374</td>
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<tr>
<td>Non-elective (long stay)</td>
<td>Normal birth with complications (CC)</td>
<td>18,298</td>
<td>£2,138</td>
<td>£80.62</td>
</tr>
<tr>
<td>Non-elective (long stay)</td>
<td>Normal birth without CC</td>
<td>63,195</td>
<td>£1,624</td>
<td>£211.49</td>
</tr>
<tr>
<td>Category</td>
<td>Birth details</td>
<td>Cases</td>
<td>Cost</td>
<td>Weighted Cost</td>
</tr>
<tr>
<td>------------------------------</td>
<td>----------------------------------------------</td>
<td>-------</td>
<td>--------</td>
<td>---------------</td>
</tr>
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<td>Non-elective (long stay)</td>
<td>Normal birth with epidural with CC</td>
<td>2,258</td>
<td>£2,280</td>
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<td>Non-elective (long stay)</td>
<td>Normal birth with epidural without CC</td>
<td>5,307</td>
<td>£1,745</td>
<td>£19.08</td>
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<td>Non-elective (long stay)</td>
<td>Normal birth with induction with CC</td>
<td>29,101</td>
<td>£2,496</td>
<td>£149.68</td>
</tr>
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<td>Non-elective (long stay)</td>
<td>Normal birth with induction without CC</td>
<td>56,705</td>
<td>£1,831</td>
<td>£213.96</td>
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<td>Normal birth with post-partum surgical intervention</td>
<td>12,114</td>
<td>£2,272</td>
<td>£56.72</td>
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<tr>
<td>Non-elective (long stay)</td>
<td>Assisted birth with CC</td>
<td>3,781</td>
<td>£2,449</td>
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<td>Non-elective (long stay)</td>
<td>Assisted birth without CC</td>
<td>7,012</td>
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<td>Non-elective (long stay)</td>
<td>Assisted birth with epidural with CC</td>
<td>3,137</td>
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<td>Non-elective (long stay)</td>
<td>Assisted birth with epidural without CC</td>
<td>4,874</td>
<td>£2,088</td>
<td>£20.97</td>
</tr>
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<td>Non-elective (long stay)</td>
<td>Assisted birth with induction with CC</td>
<td>15,496</td>
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<td>Non-elective (long stay)</td>
<td>Assisted birth with induction without CC</td>
<td>19,401</td>
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<td>Non-elective (long stay)</td>
<td>Assisted birth with post-partum surgical intervention</td>
<td>6,202</td>
<td>£2,618</td>
<td>£33.46</td>
</tr>
<tr>
<td>Non-elective (short stay)</td>
<td>Normal birth with complications (CC)</td>
<td>14,594</td>
<td>£977</td>
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</tr>
<tr>
<td>Non-elective (short stay)</td>
<td>Normal birth without CC</td>
<td>126,917</td>
<td>£908</td>
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<td>Non-elective (short stay)</td>
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<td>942</td>
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<td>Non-elective (short stay)</td>
<td>Normal birth with epidural without CC</td>
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<td>£10.46</td>
</tr>
<tr>
<td>Non-elective (short stay)</td>
<td>Normal birth with induction with CC</td>
<td>9,503</td>
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<td>£20.95</td>
</tr>
<tr>
<td>Non-elective (short stay)</td>
<td>Normal birth with induction without cc</td>
<td>50,040</td>
<td>£989</td>
<td>£101.98</td>
</tr>
<tr>
<td>Non-elective (short stay)</td>
<td>Normal birth with post-partum surgical intervention</td>
<td>4,890</td>
<td>£1,203</td>
<td>£12.12</td>
</tr>
<tr>
<td>Non-elective (short stay)</td>
<td>Assisted birth with CC</td>
<td>1,486</td>
<td>£1,128</td>
<td>£3.45</td>
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<tr>
<td>Non-elective (short stay)</td>
<td>Assisted birth without CC</td>
<td>5,889</td>
<td>£1,060</td>
<td>£12.86</td>
</tr>
<tr>
<td>Non-elective (short stay)</td>
<td>Assisted birth with epidural with CC</td>
<td>768</td>
<td>£1,249</td>
<td>£1.98</td>
</tr>
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<td>Non-elective (short stay)</td>
<td>Assisted birth with epidural without CC</td>
<td>2,355</td>
<td>£1,159</td>
<td>£5.62</td>
</tr>
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<td>Non-elective (short stay)</td>
<td>Assisted birth with induction with CC</td>
<td>2,282</td>
<td>£1,219</td>
<td>£5.73</td>
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<td>Non-elective (short stay)</td>
<td>Assisted birth with induction without CC</td>
<td>6,609</td>
<td>£1,138</td>
<td>£15.50</td>
</tr>
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<td>Non-elective (short stay)</td>
<td>Assisted birth with post-partum surgical intervention</td>
<td>1,067</td>
<td>£1,345</td>
<td>£2.96</td>
</tr>
<tr>
<td>Day</td>
<td>Normal birth with complications (CC)</td>
<td>5</td>
<td>£1,484</td>
<td>£0.02</td>
</tr>
<tr>
<td>Day</td>
<td>Normal birth without CC</td>
<td>25</td>
<td>£980</td>
<td>£0.05</td>
</tr>
<tr>
<td>Day</td>
<td>Normal birth with induction with CC</td>
<td>3</td>
<td>£1,074</td>
<td>£0.01</td>
</tr>
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</table>
### Table 13.8: Planned caesarean section costs

<table>
<thead>
<tr>
<th>Category</th>
<th>Delivery details</th>
<th>Cases</th>
<th>Cost</th>
<th>Weighted Cost</th>
</tr>
</thead>
<tbody>
<tr>
<td>Elective</td>
<td>Planned Lower Uterine Caesarean Section</td>
<td>1,897</td>
<td>£1,822</td>
<td>£58.23</td>
</tr>
<tr>
<td>Non-elective (long stay)</td>
<td>Planned Lower Uterine Caesarean Section</td>
<td>54,206</td>
<td>£2,441</td>
<td>£2,209.09</td>
</tr>
<tr>
<td>Non-elective (short stay)</td>
<td>Planned Lower Uterine Caesarean Section</td>
<td>3,249</td>
<td>£1,488</td>
<td>£81.45</td>
</tr>
<tr>
<td>Day case</td>
<td>Planned Lower Uterine Caesarean Section</td>
<td>7</td>
<td>£2,011</td>
<td>£0.24</td>
</tr>
<tr>
<td>TOTAL</td>
<td></td>
<td>59,359</td>
<td>£2,369</td>
<td></td>
</tr>
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</table>

### Table 13.9: Unplanned caesarean section costs

<table>
<thead>
<tr>
<th>Category</th>
<th>Delivery details</th>
<th>Cases</th>
<th>Cost</th>
<th>Weighted Cost</th>
</tr>
</thead>
<tbody>
<tr>
<td>Elective</td>
<td>Emergency or Upper Uterine Caesarean Section</td>
<td>1,005</td>
<td>£2,979</td>
<td>£34.11</td>
</tr>
<tr>
<td>Non-elective (long stay)</td>
<td>Emergency or Upper Uterine Caesarean Section</td>
<td>84,286</td>
<td>£3,088</td>
<td>£2,965.19</td>
</tr>
<tr>
<td>Non-elective (short stay)</td>
<td>Emergency or Upper Uterine Caesarean Section</td>
<td>2,475</td>
<td>£1,496</td>
<td>£42.18</td>
</tr>
<tr>
<td>Day case</td>
<td>Emergency or Upper Uterine Caesarean Section</td>
<td>11</td>
<td>£2,535</td>
<td>£0.32</td>
</tr>
<tr>
<td>TOTAL</td>
<td></td>
<td>87,777</td>
<td>£3,042</td>
<td></td>
</tr>
</tbody>
</table>

### Table 13.10: Proportion of actual modes of birth for planned vaginal and caesarean section birth

<table>
<thead>
<tr>
<th>Planned Method</th>
<th>Actual Method</th>
<th>%</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vaginal</td>
<td>Vaginal</td>
<td>85</td>
<td>NHS Reference Costs 2009/10(^\text{12})</td>
</tr>
</tbody>
</table>

\(^{12}\) It is assumed that a planned vaginal birth can only result in an actual vaginal delivery or emergency caesarean section. Therefore, we assume that 59,359 births (see Table 13.18) were planned caesarean section and that these accounted for 96% of planned caesarean section births (Caesarean Section Guideline 2004), giving an estimate of 61,832 planned caesarean sections in total. From the Caesarean Section Guideline 2004 we assume that 2% of planned caesarean sections result in an
Planned Method | Actual Method | %  | Source                                
---|---|---|---
Vaginal | Unplanned caesarean | 15 | NHS Reference Costs 2009/10¹²  
Caesarean section | Vaginal | 2 | Caesarean Section Guideline 2004  
Caesarean section | Unplanned caesarean | 2 | Caesarean Section Guideline 2004  
Caesarean section | Caesarean | 96 | Caesarean Section Guideline 2004  

Table 13.11 Weighted mean cost of birth by planned mode of birth

<table>
<thead>
<tr>
<th>Planned Method</th>
<th>Weighted mean cost</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vaginal</td>
<td>£1,741</td>
<td>(0.85 x £1,512) + (0.15 x £3,042)</td>
</tr>
<tr>
<td>Caesarean section</td>
<td>£2,365</td>
<td>(0.02 x £1,512) + (0.02 x £3,042) + (0.96 x £2,369)</td>
</tr>
</tbody>
</table>

Downstream costs

If the evaluation was restricted solely to birth costs then the true opportunity cost of choosing one mode of birth over another is likely to be misreported, given that there are a number of adverse maternal and neonatal outcomes associated with birth. This analysis estimates “downstream” costs associated with adverse outcomes by utilising the clinical review of the risks of planned vaginal birth and planned caesarean section undertaken for this guideline. The outcomes are limited to those for which there was reported data in the review, which focused on outcome by planned, as opposed to actual, mode of birth. For some of these outcomes results from more than one study were presented. However, it wasn’t reasonable to pool results from these studies and in such cases the model used the risk from the largest study. Whilst this provides a consistent approach it doesn’t necessarily mean that the bigger study estimated the true risk more accurately. Sensitivity analysis could be used to test whether using estimates based on other studies made important changes to the model outcome. The cost of each adverse outcome is shown below, see Table 13.12. A weighted mean cost associated with adverse outcomes can then be calculated based on the risk of that outcome, as shown in Table 13.13. These costs are then added to the planned birth cost to give the total estimated cost of planned vaginal birth and planned caesarean section.

Table 13.12 The costs of adverse birth outcomes¹³

<table>
<thead>
<tr>
<th>Outcome</th>
<th>Cost</th>
<th>Source</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Maternal death</td>
<td>£0</td>
<td>Assumption</td>
<td>It is assumed that these costs would be included within birth costs</td>
</tr>
</tbody>
</table>
| Injury to bladder, ureter, genital tract | £504 | NHS Reference Costs 2009/10 | HRG Currency Code LB15D/E  
Bladder minor procedures |

¹³ Costs based on NHS Reference Costs are generally a weighted average of all costs given for a particular currency code. It is weighted by the cases or ‘Activity’ levels shown in the NHS Reference costs.
<table>
<thead>
<tr>
<th>Procedure</th>
<th>Cost (£)</th>
<th>Reference</th>
<th>Code</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hysterectomy</td>
<td>£2,999</td>
<td>NHS Reference Costs 2009/10</td>
<td>MA07C/D, Upper genital tract major procedures</td>
</tr>
<tr>
<td>DVT</td>
<td>£686</td>
<td>NHS Reference Costs 2009/10</td>
<td>QZ20Z</td>
</tr>
<tr>
<td>Early PPH</td>
<td>£0</td>
<td>Assumption</td>
<td>It is assumed these costs would be included within birth costs</td>
</tr>
<tr>
<td>Infection (wound and post partum)</td>
<td>£0</td>
<td>Assumption</td>
<td>It is assumed these costs would be included within birth costs</td>
</tr>
<tr>
<td>Anaesthetic complication</td>
<td>£0</td>
<td>Assumption</td>
<td>It is assumed these costs would be included within birth costs</td>
</tr>
<tr>
<td>Uterine rupture</td>
<td>£0</td>
<td>Assumption</td>
<td>It is assumed these costs would be included within birth costs</td>
</tr>
<tr>
<td>Intraoperation trauma</td>
<td>£0</td>
<td>Assumption</td>
<td>It is assumed these costs would be included within birth costs</td>
</tr>
<tr>
<td>Assisted ventilations or intubations</td>
<td>£1,962</td>
<td>NHS Reference Costs 2009/10</td>
<td>DZ27A/B/C, Respiratory Failure with Intubation</td>
</tr>
<tr>
<td>Acute renal failure</td>
<td>£3,120</td>
<td>NHS Reference Costs 2009/10</td>
<td>PA38D, Renal Disease with Renal Failure with length of stay 1 day or more</td>
</tr>
<tr>
<td>Cardiac arrest</td>
<td>£1,207</td>
<td>NHS Reference Costs 2009/10</td>
<td>EB05Z</td>
</tr>
<tr>
<td>Obstetric shock</td>
<td>£1,297</td>
<td>NHS Reference Costs 2009/10</td>
<td>EB03I, Heart failure or shock</td>
</tr>
<tr>
<td>Neonatal mortality</td>
<td>£1,150</td>
<td>NHS Reference Costs 2009/10</td>
<td>PB01Z, Major neonatal diagnoses</td>
</tr>
<tr>
<td>Outcome</td>
<td>Cost</td>
<td>Vaginal birth risk</td>
<td>Weighted Vaginal cost</td>
</tr>
<tr>
<td>----------------------------------------------</td>
<td>--------</td>
<td>--------------------</td>
<td>-----------------------</td>
</tr>
<tr>
<td>Maternal death</td>
<td>£0</td>
<td>0.00002</td>
<td>£0</td>
</tr>
<tr>
<td>Injury to bladder, ureter, genital tract</td>
<td>£504</td>
<td>0.00984</td>
<td>£4.96</td>
</tr>
<tr>
<td>Hysterectomy</td>
<td>£2,999</td>
<td>0.00016</td>
<td>£0.48</td>
</tr>
<tr>
<td>DVT</td>
<td>£686</td>
<td>0.00027</td>
<td>£0.19</td>
</tr>
<tr>
<td>Blood transfusion</td>
<td>£863</td>
<td>0.00065</td>
<td>£0.56</td>
</tr>
<tr>
<td>Early PPH</td>
<td>£0</td>
<td>0.06198</td>
<td>£0.00</td>
</tr>
<tr>
<td>Infection (wound and post partum)</td>
<td>£0</td>
<td>0.00211</td>
<td>£0.00</td>
</tr>
<tr>
<td>Anaesthetic complication</td>
<td>£0</td>
<td>0.00209</td>
<td>£0.00</td>
</tr>
<tr>
<td>Uterine rupture</td>
<td>£0</td>
<td>0.00029</td>
<td>£0.00</td>
</tr>
<tr>
<td>Intraoperation trauma</td>
<td>£0</td>
<td>0.00288</td>
<td>£0.00</td>
</tr>
<tr>
<td>Assisted ventilations or intubations</td>
<td>£1,962</td>
<td>0.00006</td>
<td>£0.12</td>
</tr>
<tr>
<td>Acute renal failure</td>
<td>£3,120</td>
<td>0.00015</td>
<td>£0.47</td>
</tr>
<tr>
<td>Cardiac arrest</td>
<td>£1,207</td>
<td>0.00039</td>
<td>£0.47</td>
</tr>
<tr>
<td>Obstetric shock</td>
<td>£1,297</td>
<td>0.00019</td>
<td>£0.25</td>
</tr>
<tr>
<td>Neonatal mortality</td>
<td>£1,150</td>
<td>0.00071</td>
<td>£0.82</td>
</tr>
<tr>
<td>HIE (CNS depression, Seizures, PH &lt; 7)</td>
<td>£1,150</td>
<td>0.00234</td>
<td>£2.69</td>
</tr>
<tr>
<td>Intracranial haemorrhage</td>
<td>£1,150</td>
<td>0.00026</td>
<td>£0.30</td>
</tr>
</tbody>
</table>

Table 13.13 The weighted cost of adverse outcomes by planned mode of birth
### Outcome Cost Vaginal birth Weighted CS risk Weighted Vaginal cost CS cost

<table>
<thead>
<tr>
<th>Outcome</th>
<th>Cost</th>
<th>Vaginal risk</th>
<th>Weighted</th>
<th>CS risk</th>
<th>Weighted</th>
<th>CS cost</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neonatal respiratory morbidity</td>
<td>£1,150</td>
<td>0.11528</td>
<td>£132.57</td>
<td>0.12046</td>
<td>£138.53</td>
<td></td>
</tr>
<tr>
<td>composite of respiratory morbidity</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>NICU admission</td>
<td>£1,087</td>
<td>0.06308</td>
<td>£68.57</td>
<td>0.13889</td>
<td>£150.97</td>
<td></td>
</tr>
<tr>
<td><strong>TOTAL</strong></td>
<td><strong>£212.45</strong></td>
<td></td>
<td><strong>£298.80</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

### QALYs

Cost effectiveness is determined by effects as well as costs and this requires any differences between planned modes of birth in terms of health related quality of life to be estimated. A health state utility for a particular outcome is estimated and combined with the duration in this state to estimate a Quality Adjusted Life Year (QALY); NICE’s preferred outcome measure for economic evaluation. All QALYs are discounted at an annual rate of 3.5% in accordance with NICE guidance.

Table 13.14 shows the estimated QALYs allocated to each adverse pregnancy outcome, using only those outcomes for which data was reported in the clinical review. Then using the clinical review undertaken for this guideline, the risks of these adverse outcomes for each mode of birth can be used to calculate a weighted QALY loss associated with each planned mode of birth as shown in Table 13.15.

No QALY was assigned to the actual mode of birth although there are studies which have done so (Xu et al., 2010; Vandenbussche et al., 1999; Turner et al., 2008). This was because duration in this birth ‘state’ is short and the concomitant QALY loss would be negligible. Similarly, the base case QALY loss for many of the adverse pregnancy outcomes are set to zero, because although there may be an important health state utility loss associated with that outcome, the duration of that loss is likely to be short in which case there will only be a very small associated QALY loss.

### Table 13.14 The QALY loss of adverse birth outcomes

<table>
<thead>
<tr>
<th>Outcome</th>
<th>QALY loss</th>
<th>Notes</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Maternal death</td>
<td>24.80</td>
<td>This is based on the 53 years remaining life expectancy of a mother (ONS, 2011) at an age at giving birth of 29.4 years, the mean maternal age at birth (ONS, 2010). It assumes remaining years are lived in full-health.</td>
<td></td>
</tr>
<tr>
<td>Injury to bladder, ureter, genital tract</td>
<td>0.00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hysterectomy</td>
<td>9.79</td>
<td>This is based on a utility loss of 0.395 (Xu et al., 2010) which is assumed to be lifelong and is therefore calculated for the remaining 53 years life expectancy of a mother in the same way as maternal death</td>
<td></td>
</tr>
<tr>
<td>DVT</td>
<td>0.00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Blood transfusion</td>
<td>0.00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Early PPH</td>
<td>0.00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Infection (wound and postpartum)</td>
<td>0.00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Anaesthetic complication</td>
<td>0.00</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
### Table 13.15 The weighted QALY loss of adverse outcomes by planned mode of birth

<table>
<thead>
<tr>
<th>Outcome</th>
<th>QALY loss</th>
<th>vaginal risk</th>
<th>Weighted vaginal QALY loss</th>
<th>CS risk</th>
<th>Weighted CS QALY loss</th>
</tr>
</thead>
<tbody>
<tr>
<td>Maternal death</td>
<td>24.80</td>
<td>0.00002</td>
<td>0.00050</td>
<td>0.00000</td>
<td>0.00000</td>
</tr>
<tr>
<td>Injury to bladder, ureter, genital tract</td>
<td>0.00</td>
<td>0.00984</td>
<td>0.00000</td>
<td>0.00000</td>
<td>0.00000</td>
</tr>
<tr>
<td>Hysterectomy</td>
<td>9.79</td>
<td>0.0016</td>
<td>0.00157</td>
<td>0.00058</td>
<td>0.00568</td>
</tr>
<tr>
<td>DVT</td>
<td>0.00</td>
<td>0.0027</td>
<td>0.00000</td>
<td>0.00060</td>
<td>0.00000</td>
</tr>
<tr>
<td>Blood transfusion</td>
<td>0.00</td>
<td>0.0065</td>
<td>0.00000</td>
<td>0.00024</td>
<td>0.00000</td>
</tr>
<tr>
<td>Early PPH</td>
<td>0.00</td>
<td>0.06198</td>
<td>0.00000</td>
<td>0.03883</td>
<td>0.00000</td>
</tr>
<tr>
<td>Infection (wound and post partum)</td>
<td>0.00</td>
<td>0.00211</td>
<td>0.00000</td>
<td>0.00601</td>
<td>0.00000</td>
</tr>
<tr>
<td>Anaesthetic complication</td>
<td>0.00</td>
<td>0.00209</td>
<td>0.00000</td>
<td>0.00528</td>
<td>0.00000</td>
</tr>
<tr>
<td>Uterine rupture</td>
<td>0.00</td>
<td>0.0029</td>
<td>0.00000</td>
<td>0.00015</td>
<td>0.00000</td>
</tr>
<tr>
<td>Intraoperation trauma</td>
<td>0.00</td>
<td>0.00288</td>
<td>0.00000</td>
<td>0.00139</td>
<td>0.00000</td>
</tr>
<tr>
<td>Assisted ventilations or intubations</td>
<td>0.00</td>
<td>0.0006</td>
<td>0.00000</td>
<td>0.00013</td>
<td>0.00000</td>
</tr>
</tbody>
</table>

This is based on a life expectancy of 80 years at birth (ONS, 2011) and assumes remaining years are lived in full-health.

- **HIE (CNS depression, 04.43 Seizures, PH < 7)**
- **Infection (wound and post partum)**

This is based on a life expectancy of 80 years at birth (ONS, 2011) and using mild cerebral palsy as a proxy to estimate health state utility loss. We assume a health state utility loss of 0.16 (Heintz et al., 2008)
Table 13.16 Combined maternal/infant QALY by planned mode of birth

<table>
<thead>
<tr>
<th>Planned mode of birth</th>
<th>QALY</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vaginal</td>
<td>51.448</td>
</tr>
<tr>
<td>Caesarean section</td>
<td>51.418</td>
</tr>
</tbody>
</table>

Results

The results are shown in Table 13.17. The base case result suggests that the birth cost of a planned vaginal birth is £800 cheaper than a planned caesarean section. For an annual birth rate of 706,000 (ONS) this might suggest that approximately £5.6 million could be saved for every one percentage point reduction in caesarean section rate, providing that the change occurred in a population similar to that used in this model.

---

14 Singleton pregnancies are assumed

15 Total lifetime QALY of healthy mother and infant: 24.80 + 27.68 = 52.48
### Table 13.17 Results

<table>
<thead>
<tr>
<th>Planned mode of birth</th>
<th>Birth cost</th>
<th>Adverse outcomes cost</th>
<th>Total cost</th>
<th>Incremental cost</th>
<th>Total QALY</th>
<th>Incremental QALY</th>
<th>ICER</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vaginal</td>
<td>£1,741</td>
<td>£212</td>
<td>£1,954</td>
<td>-</td>
<td>51.448</td>
<td>0.030</td>
<td>Dominant</td>
</tr>
<tr>
<td>Caesarean section</td>
<td>£2,365</td>
<td>£299</td>
<td>£2,664</td>
<td>£710</td>
<td>51.418</td>
<td>-</td>
<td>Dominated</td>
</tr>
</tbody>
</table>

However, beyond the immediate term it is more complicated than this. Using the base case model inputs the results show that the saving might be even greater when adverse outcomes are considered, although uncertainty surrounds the point estimates of risks on which these costs are based. Furthermore, there are other adverse outcomes not included within the clinical review which, if included, could possibly yield a different result.

Due to the relative infrequency of adverse outcomes having a major QALY loss, there is only a small difference in QALYs between the two modes of birth in the base case analysis. Here, it slightly favoured planned vaginal birth primarily because of a considerably lower neonatal mortality rate. Planned vaginal is said to be dominant, being cheaper and yielding a higher QALY.

### Sensitivity Analysis

#### Varying actual vaginal birth rate from planned vaginal birth

In this one-way sensitivity analysis shown in Figure 13.11 below, we see how the actual vaginal birth rate determines the incremental costs of maternal request caesarean section. Figure 13.14 shows that if the actual rate of vaginal birth for planned vaginal birth fell to 44% or below, then maternal request caesarean section would become the cheapest birth option when only the immediate birth costs are considered. However, the current caesarean section rate in England is 24.6% (Department of Health, 2009) and therefore, given that a large proportion of these will have a medical or obstetric indication, the lower bound of planned vaginal births which result in actual vaginal birth must be at least 75.4%.

**Figure 13.11** Incremental costs of maternal request caesarean section varying the percentage of planned vaginal births leading to actual vaginal birth
Probabilistic sensitivity analysis

The base case analysis is deterministic using point estimates for the model’s input parameters. However, it is usual practice in economic evaluation to address uncertainty in point estimate values through the use of sensitivity analysis. Where there are many input parameters probabilistic sensitivity analysis is usually recommended to address uncertainty.

In the probabilistic analysis undertaken here, 1,000 Monte Carlo simulations were run with the risks of adverse outcomes, included in the review reported for this guideline, sampled from a beta probability distribution, with the alpha parameter for each distribution given by the number of events and the beta parameter as the number of non-events. All other model inputs are fixed at their base case value, although the model allows the probabilistic analyses to be run with different values for these inputs.

The results are shown in Figure 13.12

In this analysis maternal request caesarean section had a higher incremental cost and a lower incremental QALY, suggesting that planned vaginal birth dominated maternal request caesarean section with a probability of 100%.

Figure 13.12 Probabilistic sensitivity analysis of incremental costs and incremental QALYs of maternal request caesarean section

Introducing urinary incontinence as an adverse outcome

The base case analysis and sensitivity analyses above focused solely on adverse outcomes that were included in the review, a review which focused on reported outcomes according to the planned mode of birth. However, other studies have compared adverse outcomes by actual mode of birth and this was the approach in the previous version of this guideline. In these studies there are an increased number of reported adverse outcomes particularly relating to a woman’s pelvic floor. The model has been set up to allow analyses which include the additional adverse outcomes shown in Table 13.18.

In the default case the risks, costs and QALY loss of all these adverse outcomes is set to zero but these assumptions can all be varied as part of a sensitivity analysis.

Table 13.18 Additional adverse outcomes not included in the base case analysis

<table>
<thead>
<tr>
<th>Outcome</th>
</tr>
</thead>
<tbody>
<tr>
<td>Iatrogenic surgical injury</td>
</tr>
<tr>
<td>Perineal and abdominal pain (4 months post partum)</td>
</tr>
<tr>
<td>Pulmonary embolism</td>
</tr>
</tbody>
</table>

Caesarean section: full guideline DRAFT (September 2011)
Outcome

- Faecal incontinence
- Post natal depression
- Dyspareunia
- Urinary incontinence

Here we explore what would be the impact of introducing urinary income to the analysis. Model inputs for this purpose were extrapolated from Xu et al. 2010.

Cost of stress urinary incontinence

Xu et al. assumes that only 61% of those with stress urinary incontinence seek health care. For those that do an annual cost of £375 in routine care is assumed.\(^{16,17}\) Assuming a life expectancy of 80 years and an age at birth of 29 years old, these costs would be on-going for 50 years. It is additionally assumed that there is a diagnostic cost of £150 and that 18.2% of patients with stress urinary incontinence will have surgery at 54 years of age at a cost of £6,202. Discounting future costs at an annual 3.5% discount rate as recommended by NICE, the cost of stress urinary incontinence is estimated as follows:

- **Routine care**: £9,163 x 0.61 = £5,589
- **Diagnosis**: £150 x 0.61 = £92
- **Surgery**: £6,202 x 0.182 = £1,129

**Total** = **£6,810**

QALY loss from urinary incontinence

Xu et al. report a health state utility of 0.81 for stress urinary incontinence and 0.87 after successful surgery. We assume here that this represents a health state utility loss of 0.19 and 0.13 respectively. We follow Xu et al. in assuming that the 39% of women with stress urinary incontinence who do not seek health care suffer no disutility. It is assumed that 18.2% of women with stress urinary incontinence will have surgery and that it will be successful in 81.3% of these patients. Thus, the discounted QALY loss from stress urinary incontinence is calculated as shown below:

- **Not seeking health care**: 0.39 x 0 = 0.00 QALYs
- **Never surgery**: \((0.61 - 0.182) \times \sum_{n=0}^{50} (0.19 \times 1.035^n) = 1.99\) QALYs
- **Unsuccessful surgery**: \(0.034 \times \sum_{n=0}^{50} (0.19 \times 1.035^n) = 0.16\) QALYs
- **‘Cure’**: \(0.148 \times \left( \sum_{n=0}^{24} (0.19 \times 1.035^n) + \sum_{n=25}^{50} (0.13 \times 1.035^n) \right) = 0.62\) QALYs

**Total QALY loss** = **2.77** QALYs

Urinary incontinence risk

Xu et al. in their base case analysis assume that the probability of a woman experiencing stress urinary incontinence is 19.9% for a spontaneous vaginal birth, 21.8% for an instrumental vaginal birth, 11.5% for an unplanned caesarean section and 10.0% for a caesarean section on maternal request. Therefore, for the purposes of this sensitivity analysis we calculate the urinary incontinence risk for each mode of birth as follows:

\[^{16}\] This is likely to include private expenditure which wouldn’t be counted using the NICE reference case
\[^{17}\] http://www.expedia.co.uk/pub/agent.dll - accessed 21/01/2011 exchange rate $1 = £0.629683
Planned vaginal:

Using NHS Reference Costs 2009/10 activity data normal birth accounts for 83.2% of vaginal births with the remaining 16.8% being assisted:

Weighted risk of actual vaginal birth: \(0.832 \times 0.90 \times 0.199) + (0.168 \times 0.90 \times 0.218) = 0.182\)

Weighted risk of an unplanned caesarean section: \(0.10 \times 0.115) = 0.0115\)

**Planned vaginal stress urinary incontinence risk = 0.194**

Maternal request caesarean section:

Weighted risk of actual vaginal: \(0.832 \times 0.02 \times 0.199) + (0.168 \times 0.02 \times 0.218) = 0.004\)

Weighted risk of an unplanned caesarean section: \(0.02 \times 0.115) = 0.002\)

Weighted risk of planned caesarean section: \(0.96 \times 0.10) = 0.096\)

**Maternal request caesarean section stress urinary incontinence risk = 0.102**

The results of this sensitivity analysis are shown in Table 13.19.

<table>
<thead>
<tr>
<th>Planned mode of birth</th>
<th>Birth cost</th>
<th>Adverse outcomes cost</th>
<th>Total cost</th>
<th>Incremental cost</th>
<th>Total QALY</th>
<th>Incremental QALY</th>
<th>ICER per QALY</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vaginal</td>
<td>£1,741</td>
<td>£1,534</td>
<td>£3,275</td>
<td>-</td>
<td>51.911</td>
<td>-</td>
<td></td>
</tr>
<tr>
<td>Caesarean section</td>
<td>£2,365</td>
<td>£993</td>
<td>£3,359</td>
<td>£84</td>
<td>52.135</td>
<td>0.224</td>
<td>£373</td>
</tr>
</tbody>
</table>

Including urinary incontinence greatly reduces the incremental costs of a maternal request caesarean section, because the “downstream” costs of a planned vaginal birth increase more due to the higher risk of stress urinary incontinence with vaginal birth. Similarly, the greater reduction in health related quality of life arising in women having a planned vaginal birth from stress urinary incontinence now leads to caesarean section on maternal request having a higher QALY. An incremental cost-effectiveness ratio of £373 per QALY would suggest that a maternal request caesarean section could be considered a cost-effective alternative to planned vaginal birth.

Discussion

The results presented in this analysis do not definitively determine the relative cost-effectiveness of maternal request caesarean section as opposed to planned vaginal birth. In terms of the immediate costs of birth, this model, using the most comprehensive and detailed NHS Reference Costs yet produced for modes of birth, does suggest that a planned vaginal birth is cheaper. The base case analysis suggests that the NHS could save in the region of £5.6 million in birth costs for every percentage point reduction in caesarean section, at least if the reduction occurred in women with similar obstetric and medical characteristics to the model population. It is possible that the differential of £700 is over-stated as the cost data for caesarean section does also include caesarean section where there was a medical or obstetric indication. As was shown in the sensitivity analysis varying the proportion of planned vaginal births leading to actual births, the immediate costs of planned vaginal birth are likely to remain cheaper even if the base case proportion of actual vaginal births was overstated.

The base case and probabilistic sensitivity analysis both suggested that planned vaginal birth was more cost-effective and cheaper than maternal request caesarean section. In a publicly funded health service such a result can be used to justify a decision not to make caesarean section available purely on grounds of maternal request. In these analyses planned vaginal birth was also cheaper in terms of the “downstream” costs associated with adverse outcomes. Here the adverse outcomes were limited to those included in the review produced for this guideline, which in turn was based on studies.
reporting adverse outcomes based on planned mode of birth rather than actual mode of birth, although the former, if studied sufficiently, should implicitly capture the effects of actual birth. In addition to the cheaper costs of planned vaginal birth, the cost-effectiveness was also driven by a smaller loss of QALYs with planned vaginal birth. The drivers of this are the increased relative risk of hysterectomy and neonatal mortality with maternal request caesarean section. In the base case model, a planned vaginal birth had a higher maternal mortality than a maternal request caesarean section, although higher maternal mortality for caesarean section generally is often reported (Harper et al., 2003). Furthermore, one of the studies included in the clinical review undertaken for this guideline (Deneux-Tharaux, 2006a) reported higher maternal mortality for planned caesarean section. We did not use the data from this study as our method where outcomes were reported in more than one study was to use the largest study. However, if higher maternal mortality with planned caesarean section is assumed this simply strengthens the cost-effectiveness conclusion of the base case analysis.

However, the risk could also have been evaluated by actual mode of birth rather than the planned mode of birth as was the case with the previous version of this guideline. In such studies women are often reported at being at greater risk of urinary incontinence following vaginal birth, as was also reported in the previous version of this guideline. To address this issue a sensitivity analysis was undertaken in which urinary incontinence was introduced as an adverse outcome. The model inputs for this adverse outcome were taken from or extrapolated from a study by Xu et al. (2010) which suggested that if society was willing to pay $50,000 per QALY then there was an 82% probability that maternal request caesarean section was cost-effective for a primigravid woman without medical or obstetric indication and having only one childbirth in her lifetime. In this sensitivity analysis, the model suggested that maternal request caesarean section would be cost-effective even if remaining slightly more expensive as a result of the lower QALY loss arising from reduced rates of stress urinary incontinence. Clearly, there are other adverse outcomes besides urinary incontinence which weren’t reported in the clinical review for this guideline which may also have a bearing on the cost-effectiveness of the different modes of birth.

However, the Xu study may also have its limitations especially in an England and Wales context. Costs differ substantially between the US and England and Wales. The costs they report, which include productivity losses, are different than that would be used by an evaluation employing NICE methodology. Furthermore, whilst their parameter estimates were obtained from the published literature there is no indication to suggest that these were retrieved in a systematic way. There are a number of studies which have contested the extent to which caesarean section is protective against urinary stress incontinence especially across the entire childbearing population (Nygaard, 2006). Others have acknowledged an increased short term occurrence with vaginal birth but claim that severe symptoms do not differ by mode of birth (Press et al., 2007). To fully model the effect of stress urinary incompetence is incredibly complicated if it is accepted that there are differences by mode of birth. It essentially involves having a model for a complete disease pathway broken down by disease severity. Conservative and medical treatment alternatives exist for stress urinary incontinence and these are not included within the Xu model.

**Conclusion**

This model suggests that the immediate birth costs are lower for planned vaginal delivery than they are for maternal request caesarean section. However, the model does not conclusively demonstrate the cost-effectiveness of one mode of birth over the other. Using the adverse outcomes data only included in the review produced for this guideline, planned vaginal birth does appear more cost-effective but its cost-effectiveness relative to maternal request caesarean section is likely to be reduced to some extent if adverse outcomes such as urinary incontinence are included within the model.

Given these results, there is no strong health economic evidence which would lead to a revision of previously issued NICE guidance.
13.4 Cost effectiveness of VBAC – one previous CS with no plans for further children

Introduction

The current guidance of the American College of Obstetricians and Gynaecologists is that, "Attempting a vaginal birth after cesarean (VBAC) is a safe and appropriate choice for most women who have had a prior cesarean delivery, including for some women who have had two previous cesareans." (ACOG, 2010).

In women having had a previous caesarean section there are risks associated with both a trial of labour and with repeat caesarean section. The greatest risk of adverse maternal outcomes occurs in a failed trial of labour. However, successful VBAC has the fewest complications and therefore the failure rate for trial of labour is likely to be an important determinant of the overall comparative risks of a trial of labour and repeat caesarean section.

The previous caesarean section guideline concluded that the cost-effectiveness of a trial of labour compared to a planned caesarean section couldn’t be “categorically determined” with the results sensitive to the rates of adverse events.

Method

A cost utility analysis was undertaken using a decision analytic model developed in Microsoft Excel® to compare the cost effectiveness of VBAC (vaginal birth after caesarean section) versus planned caesarean section in England and Wales in woman having one previous caesarean section and with no plans for further children. As well as considering the costs of birth, we also aim to evaluate the “downstream” impact on costs and health related quality of life arising from adverse events. This analysis was undertaken from the perspective of the NHS and personal social services which is in accordance with NICE guidelines methodology (NICE, 2009). Costs and benefits are compared using standard methods of incremental analysis of costs and benefits. Costs were based on 2009/10 prices. A number of sensitivity analyses were undertaken to assess the impact that changes in the base case assumptions would have on the model’s results.

Using the Excel model

For full model functionality macros need to be enabled. Click the ‘Options’ button and then select the ‘Enable this content’ in the Security Alert Dialog Box.

The model is navigated using on-screen buttons. On the data entry screens, data is either entered directly into cells or assigned using the slider bars. Where slider bar data entry is possible, data can also be entered directly but the slider will then not work until the ‘default’ button is pressed. Pressing the ‘default’ button returns all data to its original value.

The cost of birth

The costs are the same as used in the model comparing the cost-effectiveness of planned vaginal birth versus maternal request caesarean section (see section 13.3). As in the maternal request model, the actual cost of a planned birth depends on the actual mode of birth. For a planned VBAC the values are taken from the National Sentinel Caesarean Section Audit (Thomas J and Paranjothy S, 2001) which indicated that a trial of labour after caesarean section had a 64% success rate. For a planned caesarean section the rates for the different actual modes of birth are the same as used in the maternal request model. The default values for the actual birth method by planned method are shown in table 13.20.

Table 13.20 Proportion of actual modes of birth for planned vaginal and caesarean section birth

<table>
<thead>
<tr>
<th>Planned Method</th>
<th>Actual Method</th>
<th>%</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vaginal</td>
<td>Vaginal</td>
<td>64</td>
<td>NSCSA (2001)</td>
</tr>
<tr>
<td>Vaginal</td>
<td>Unplanned caesarean</td>
<td>36</td>
<td>NSCSA (2001)</td>
</tr>
<tr>
<td>Caesarean section</td>
<td>Vaginal</td>
<td>2</td>
<td>As maternal request</td>
</tr>
</tbody>
</table>
Caesarean section  Unplanned caesarean  2  
As maternal request model  
(see section 13.2)

Caesarean section  Caesarean  96  
As maternal request model  
(see section 13.2)

Downstream costs
In addition to the costs of birth this analysis estimates "downstream" costs associated with adverse outcomes by utilising the clinical review of VBAC undertaken for this guideline. The outcomes are limited to those for which there was reported data in the review, which focused on outcome by planned, as opposed to actual, mode of birth. Most of the costs for these adverse outcomes were reported in section 13.2. The additional cost for this analysis is shown in Table 13.21. The GDG said that transient tachypnea would typically involve an admission to a neonatal intensive care unit and it was costed on this basis. The clinical review also reported bag and mask ventilation as an outcome but the GDG considered that the costs of this would be negligible. A weighted mean cost associated with adverse outcomes can then be calculated based on the risk of that outcome, as shown in Table 13.22. These costs are then added to the planned birth cost to give the total estimated cost of planned vaginal birth and planned caesarean section.

Table 13.21 The costs of adverse birth outcomes

<table>
<thead>
<tr>
<th>Outcome</th>
<th>Cost</th>
<th>Source</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Transient tachypnea</td>
<td>£1,087</td>
<td>NHS Reference Costs 2009/10</td>
<td>HRG XA01Z Neonatal critical care intensive care</td>
</tr>
</tbody>
</table>

Table 13.22 The weighted cost of adverse outcomes by planned birth type

<table>
<thead>
<tr>
<th>Outcome</th>
<th>Cost</th>
<th>Vaginal birth risk</th>
<th>Weighted Vaginal cost</th>
<th>CS risk</th>
<th>Weighted CS cost</th>
</tr>
</thead>
<tbody>
<tr>
<td>Maternal death</td>
<td>£0</td>
<td>0.00002</td>
<td>£0</td>
<td>0.00008</td>
<td>£0.00</td>
</tr>
<tr>
<td>Hysterectomy</td>
<td>£2,999</td>
<td>0.00117</td>
<td>£3.51</td>
<td>0.00119</td>
<td>£3.57</td>
</tr>
<tr>
<td>Blood transfusion</td>
<td>£863</td>
<td>0.00383</td>
<td>£3.31</td>
<td>0.00304</td>
<td>£2.62</td>
</tr>
<tr>
<td>Infection (wound and post partum)</td>
<td>£0</td>
<td>0.04600</td>
<td>£0.00</td>
<td>0.03200</td>
<td>£0.00</td>
</tr>
<tr>
<td>Uterine rupture</td>
<td>£0</td>
<td>0.00714</td>
<td>£0.00</td>
<td>0.00023</td>
<td>£0.00</td>
</tr>
<tr>
<td>Neonatal mortality</td>
<td>£1,150</td>
<td>0.00115</td>
<td>£1.32</td>
<td>0.00063</td>
<td>£0.72</td>
</tr>
<tr>
<td>Transient</td>
<td>£1,087</td>
<td>0.12373</td>
<td>£134.49</td>
<td>0.12873</td>
<td>£139.93</td>
</tr>
</tbody>
</table>

18 Costs based on NHS Reference Costs are generally a weighted average of all costs given for a particular currency code. It is weighted by the cases or 'Activity' levels shown in the NHS Reference costs.
QALYs

A QALY loss was estimated for maternal death, neonatal mortality and hysterectomy. The values for these outcomes are shown in section 13.2. All QALYs are discounted at an annual rate of 3.5% in accordance with NICE guidance.

Using the clinical review undertaken for this guideline, the risks of these adverse outcomes for each birth type were used to calculate a weighted QALY loss associated with each planned birth type as shown in Table 13.23.

Table 13.23 The weighted QALY loss of adverse outcomes by planned birth type

<table>
<thead>
<tr>
<th>Outcome</th>
<th>Cost</th>
<th>Vaginal birth risk</th>
<th>Weighted Vaginal cost</th>
<th>CS risk</th>
<th>Weighted CS cost</th>
</tr>
</thead>
<tbody>
<tr>
<td>Maternal death</td>
<td>24.80</td>
<td>0.00002</td>
<td>0.00050</td>
<td>0.00008</td>
<td>0.00198</td>
</tr>
<tr>
<td>Hysterectomy</td>
<td>9.59</td>
<td>0.00117</td>
<td>0.01122</td>
<td>0.00119</td>
<td>0.01141</td>
</tr>
<tr>
<td>Blood transfusion</td>
<td>0.00</td>
<td>0.00383</td>
<td>0.00000</td>
<td>0.00304</td>
<td>0.00000</td>
</tr>
<tr>
<td>Infection (wound and post partum)</td>
<td>0.00</td>
<td>0.04600</td>
<td>0.00000</td>
<td>0.03200</td>
<td>0.00000</td>
</tr>
<tr>
<td>Uterine rupture</td>
<td>0.00</td>
<td>0.00714</td>
<td>0.00000</td>
<td>0.00023</td>
<td>0.00000</td>
</tr>
<tr>
<td>Neonatal mortality</td>
<td>27.68</td>
<td>0.00115</td>
<td>0.03183</td>
<td>0.00063</td>
<td>0.01744</td>
</tr>
<tr>
<td>Transient tachypnea</td>
<td>0.00</td>
<td>0.12373</td>
<td>0.00000</td>
<td>0.12873</td>
<td>0.00000</td>
</tr>
<tr>
<td><strong>TOTAL</strong></td>
<td><strong>0.04355</strong></td>
<td></td>
<td><strong>0.03083</strong></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Conceptually it might be considered easier to compare incremental QALY gains rather than QALY losses. Therefore, the weighted QALY losses from adverse maternal and neonatal outcomes are subtracted from the lifetime QALY of mother and infant\(^{19}\) in the absence of any adverse outcomes\(^{20}\). These are based on the 53 years remaining life expectancy of the mother and the 80 years life expectancy at birth of the infant (ONS, 2011). It is assumed that remaining life years are lived in full health and that QALYs are discounted using an annual discount rate of 3.5% (NICE, 2009). The QALYs associated with planned vaginal birth and planned caesarean section is given in Table 13.24.

Table 13.24 Combined maternal/infant QALY by planned mode of birth

<table>
<thead>
<tr>
<th>Planned mode of birth</th>
<th>QALY</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vaginal</td>
<td>52.437</td>
</tr>
<tr>
<td>Caesarean section</td>
<td>52.449</td>
</tr>
</tbody>
</table>

---

\(^{19}\) Singleton pregnancies are assumed

\(^{20}\) Total lifetime QALY of healthy mother and infant: 23.70 + 27.68 = 51.38
Results
The results are shown in Table 13.25. The base case result suggests that the birth cost of a planned VBAC is £307 cheaper than a planned caesarean section. However, the base case also suggests that planned caesarean has a higher QALY and the corresponding ICER of £24,141 indicates that planned caesarean can be considered borderline cost-effective relative to VBAC using the NICE advisory threshold of £20,000 to £30,000 per QALY, especially given the level of uncertainty surrounding long-term downstream costs and QALYs.

Table 13.25 Results

<table>
<thead>
<tr>
<th>Planned mode of birth</th>
<th>Birth cost</th>
<th>Adverse outcomes cost</th>
<th>Total cost</th>
<th>Incremental cost</th>
<th>Total QALY</th>
<th>Incremental QALY</th>
<th>ICER</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vaginal</td>
<td>£2,063</td>
<td>£143</td>
<td>£2,205</td>
<td>-</td>
<td>52.437</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Caesarean section</td>
<td>£2,365</td>
<td>£147</td>
<td>£2,512</td>
<td>£307</td>
<td>52.449</td>
<td>0.012</td>
<td>£24,141</td>
</tr>
</tbody>
</table>

Sensitivity analysis
i. Including data from review of maternal request for caesarean section

In this analysis, data for other adverse outcomes reported as part of the review for maternal request caesarean section, but not the VBAC review, were included. This suggested that a planned caesarean section had an ICER of £30,513 relative to VBAC, which would only still be considered to be borderline cost-effective if the upper limit of the NICE willingness to pay threshold of £30,000 per QALY was being used.

ii. Varying actual vaginal birth rate from planned VBAC (including data from maternal request caesarean section review)

In this one-way sensitivity analysis shown in Figure 13.13 below, we see how the actual vaginal birth rate determines the incremental costs of a planned caesarean section relative to VBAC. Figure 13.13 shows that if the actual rate of vaginal birth for planned vaginal birth fell to approximately 45% or below then planned caesarean section would become the cheapest birth option when only the immediate birth costs are considered.

Figure 13.13 Incremental costs of maternal request caesarean section varying the percentage of planned vaginal births leading to actual vaginal birth
iii. Probabilistic sensitivity analysis- VBAC review data

The base case analysis is deterministic using point estimates for the model's input parameters. However, it is usual practice in economic evaluation to address uncertainty in point estimate values through the use of sensitivity analysis. Where there are many input parameters probabilistic sensitivity analysis is usually recommended to address uncertainty.

In the probabilistic analysis undertaken here, 1,000 Monte Carlo simulations were run with the risks of adverse outcomes sampled from a beta probability distribution, with the alpha parameter for each distribution given by the number of events and the beta parameter as the number of non-events. The risks were taken from those reported for the VBAC review undertaken for this guideline. All other model inputs are fixed at their base case value, although the model allows the probabilistic analyses to be run with different values for these inputs. The results are shown in Figure 13.14.

Figure 13.14 Probabilistic sensitivity analysis of incremental costs and incremental QALYs of planned caesarean section relative to VBAC

The red line indicates a £20,000 per QALY threshold. Points under this line are considered to be cost-effective and points above are considered not to be cost-effective. More simulations occur above the threshold suggesting that there a better than a 50% chance that VBAC is cost-effective relative to a planned caesarean section. However, considerable uncertainty remains as a substantial minority of simulations show a planned caesarean section as the more cost-effective option.

iv. Probabilistic Sensitivity analysis – VBAC and maternal request review data

This probabilistic sensitivity analysis additionally includes the risk reported in the maternal request model. The results are shown in Figure 13.15.
Figure 13.15 Probabilistic sensitivity analysis of incremental costs and incremental QALYs of planned caesarean section relative to VBAC

In this case there is a greater probability that VBAC is cost-effective relative to a planned caesarean section.

Discussion

The results presented in this cost-effectiveness analysis do not provide strong evidence either way on the cost-effectiveness of VBAC relative to caesarean section in women who have had one previous birth. As with the maternal request model not all important risks have necessarily been included within the model as inputs were limited to those reported in the clinical reviews undertaken for this guideline. Important adverse outcomes not included are those relating to the pelvic floor and to subsequent pregnancies, e.g. placenta praevia. This just adds a further level of uncertainty to the equivocal cost-effectiveness conclusions of this model. In addition in the probabilistic sensitivity analysis that used data only reported in the maternal request review, it is likely that these risks will be higher in this population and the nature of this increased risk is likely to vary according to the planned mode of birth.

Conclusion

This model suggests that either VBAC or a planned caesarean section can still be supported on cost-effectiveness grounds for a woman’s 2nd birth. VBAC as a non-intervention is the woman’s right should she so choose but on the other hand any additional costs of a planned caesarean section are relatively small and can plausibly be justified in terms of additional benefit. Therefore, this model would support women being able to choose their preferred mode of birth in consultation with the health care professionals responsible for her care. Considerations about any future pregnancies may be an important factor in the decisions made.
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Thomas, J., Paranjothy, S., James, D. 2004. National cross sectional survey to determine whether the decision to delivery interval is critical in emergency caesarean section. BMJ; 7441:665-


Thomas, J., Paranjothy, S., Royal College of Obstetricians and Gynaecologists: Clinical Effectiveness Support Unit. 2001. The National Sentinel Caesarean Section Audit Report

Thomas, J., Paranjothy, S., James, D. 2004. National cross sectional survey to determine whether the decision to delivery interval is critical in emergency caesarean section. BMJ; 7441:665-
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### Abbreviations

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>APH</td>
<td>antepartum haemorrhage</td>
</tr>
<tr>
<td>ART</td>
<td>antiretroviral therapy</td>
</tr>
<tr>
<td>CEMD</td>
<td>Confidential Enquiry into Maternal Deaths</td>
</tr>
<tr>
<td>CFM</td>
<td>colour flow mapping</td>
</tr>
<tr>
<td>CI</td>
<td>confidence interval</td>
</tr>
<tr>
<td>CS</td>
<td>caesarean section</td>
</tr>
<tr>
<td>CSR</td>
<td>caesarean section rate</td>
</tr>
<tr>
<td>CTG</td>
<td>cardiotocograph</td>
</tr>
<tr>
<td>DDI</td>
<td>decision-to-delivery interval</td>
</tr>
<tr>
<td>DGH</td>
<td>district general hospital (non-teaching hospital)</td>
</tr>
<tr>
<td>DIC</td>
<td>disseminated intravascular coagulopathy</td>
</tr>
<tr>
<td>EFM</td>
<td>electronic fetal monitoring</td>
</tr>
<tr>
<td>ECV</td>
<td>external cephalic version</td>
</tr>
<tr>
<td>EL</td>
<td>evidence level</td>
</tr>
<tr>
<td>ERCS</td>
<td>elective repeat caesarean section</td>
</tr>
<tr>
<td>FBS</td>
<td>fetal blood sampling</td>
</tr>
<tr>
<td>FHR</td>
<td>fetal heart rate</td>
</tr>
<tr>
<td>FIGO</td>
<td>International Federation of Gynecology and Obstetrics</td>
</tr>
<tr>
<td>FTP</td>
<td>“failure to progress” (in labour)</td>
</tr>
<tr>
<td>GDG</td>
<td>Guideline Development Group</td>
</tr>
<tr>
<td>HAART</td>
<td>highly active antiretroviral therapy</td>
</tr>
<tr>
<td>HCV</td>
<td>Hepatitis C virus</td>
</tr>
<tr>
<td>HDU</td>
<td>high dependency unit</td>
</tr>
<tr>
<td>HIE</td>
<td>hypoxic-ischemic encephalopathy</td>
</tr>
<tr>
<td>HIV</td>
<td>human immunodeficiency virus</td>
</tr>
<tr>
<td>HSV</td>
<td>Herpes Simplex virus</td>
</tr>
<tr>
<td>HTA</td>
<td>Health Technology Assessment</td>
</tr>
<tr>
<td>ICU</td>
<td>intensive care unit</td>
</tr>
<tr>
<td>ITU</td>
<td>Intensive therapy unit</td>
</tr>
<tr>
<td>IV</td>
<td>intravenous</td>
</tr>
</tbody>
</table>
15.2 Glossary

Absolute risk

Measures the probability of an event or outcome occurring (e.g. an adverse reaction to the drug being tested) in the group of people under study. Studies that compare two or more groups of patients may report results in terms of the Absolute Risk Reduction.

Absolute risk reduction (ARR)

The ARR is the difference in the risk of an event occurring between two groups of patients in a study – for example if 6% of patients die after receiving a new experimental drug and 10% of patients die after having the old drug treatment then the ARR is 10 – 6% = 4%. Thus by using the new drug instead of the old drug 4% of patients can be prevented from dying. Here the ARR measures the risk reduction associated with a new treatment.
Allied health professionals
Healthcare professionals, other than doctors, midwives and nurse/midwife, directly involved in the provision of healthcare. Includes several groups such as physiotherapists, occupational therapists, dieticians, etc. (Formerly known as professions allied to medicine or PAMs.)

Applicability
The extent to which the results of a study or review can be applied to the target population for a clinical guideline.

Appraisal of evidence
Formal assessment of the quality of research evidence and its relevance to the clinical question or guideline under consideration, according to predetermined criteria.

Best available evidence
The strongest research evidence available to support a particular guideline recommendation.

Bias
Influences on a study that can lead to invalid conclusions about a treatment or intervention. Bias in research can make a treatment look better or worse than it really is. Bias can even make it look as if the treatment works when it actually doesn’t. Bias can occur by chance or as a result of systematic errors in the design and execution of a study. Bias can occur at different stages in the research process, e.g. in the collection, analysis, interpretation, publication or review of research data. For examples see Selection bias, Performance bias, Information bias, Confounding, Publication bias.

Blinding or masking
The practice of keeping the investigators or subjects of a study ignorant of the group to which a subject has been assigned. For example, a clinical trial in which the participating patients or their doctors are unaware of whether they (the patients) are taking the experimental drug or a placebo (dummy treatment). The purpose of ‘blinding’ or ‘masking’ is to protect against bias. See also Double blind study, Single blind study, Triple blind study.

Case-control study
A study that starts with the identification of a group of individuals sharing the same characteristics (e.g. people with a particular disease) and a suitable comparison (control) group (e.g. people without the disease). All subjects are then assessed with respect to things that happened to them in the past, e.g. things that might be related to getting the disease under investigation. Such studies are also called retrospective as they look back in time from the outcome to the possible causes.

Case series
Description of several cases of a given disease, usually covering the course of the disease and the response to treatment. There is no comparison (control) group of patients.

Causal relationship
Describes the relationship between two variables whenever it can be established that one causes the other. For example there is a causal relationship between a treatment and a disease if it can be shown that the treatment changes the course or outcome of the disease. Usually randomised controlled trials are needed to ascertain causality. Proving cause and effect is much more difficult than just showing an association between two variables. For example, if it happened that everyone who had eaten a particular food became sick, and everyone who avoided that food remained well, then the food would clearly be associated with the sickness. However, even if leftovers were found to be contaminated, it could not be proved that the food caused the sickness – unless all other possible causes (e.g. environmental factors) had been ruled out.

Clinical audit
A systematic process for setting and monitoring standards of clinical care. Whereas ‘guidelines’ define what the best clinical practice should be, ‘audit’ investigates whether best practice is being carried out. Clinical audit can be described as a cycle or spiral. Within the cycle there are stages that follow a systematic process of establishing best practice, measuring care against
specific criteria, taking action to improve care, and monitoring to sustain improvement. The spiral suggests that as the process continues, each cycle aspires to a higher level of quality.

**Clinical effectiveness**

The extent to which a specific treatment or intervention, when used under usual or everyday conditions, has a beneficial effect on the course or outcome of disease compared to no treatment or other routine care. (Clinical trials that assess effectiveness are sometimes called management trials.) Clinical ‘effectiveness’ is not the same as efficacy.

**Clinical governance**

A framework through which NHS organisations are accountable for both continuously improving the quality of their services and safeguarding high standards of care by creating an environment in which excellence in clinical care will flourish.

**Clinical impact**

The effect that a guideline recommendation is likely to have on a treatment, or treatment outcomes, of the target population.

**Clinical question**

This term is sometimes used in guideline development work to refer to the questions about treatment and care that are formulated in order to guide the search for research evidence. When a clinical question is formulated in a precise way, it is called a focused question.

**Clinician**

A health care professional providing patient care, e.g. doctor, nurse/midwife, physiotherapist.

**Cochrane Collaboration**

An international organisation in which people find, appraise and review specific types of studies called randomised controlled trials. The Cochrane Database of Systematic Reviews contains regularly updated reviews on a variety of health issues and is available electronically as part of the Cochrane Library.

**Cochrane Library**

The Cochrane Library consists of a regularly updated collection of evidence-based medicine databases including the Cochrane Database of Systematic Reviews (reviews of randomised controlled trials prepared by the Cochrane Collaboration). The Cochrane Library is available on CD-ROM and the Internet.

**Cohort study**

An observational study that takes a group (cohort) of patients and follows their progress over time in order to measure outcomes such as disease or mortality rates and make comparisons according to the treatments or interventions that patients received. Thus within the study group, subgroups of patients are identified (from information collected about patients) and these groups are compared with respect to outcome, e.g. comparing mortality between one group that received a specific treatment and one group which did not (or between two groups that received different levels of treatment). Cohorts can be assembled in the present and followed into the future (a ‘concurrent’ or ‘prospective’ cohort study) or identified from past records and followed forward from that time up to the present (a ‘historical’ or ‘retrospective’ cohort study). Because patients are not randomly allocated to subgroups, these subgroups may be quite different in their characteristics and some adjustment must be made when analysing the results to ensure that the comparison between groups is as fair as possible.

**Co-morbidity**

Co-existence of a disease or diseases in the people being studied in addition to the health problem that is the subject of the study.

**Confidence interval**

A way of expressing certainty about the findings from a study or group of studies, using statistical techniques. A confidence interval describes a range of possible effects (of a treatment or intervention) that are consistent with the results of a study or group of studies. A wide confidence interval indicates a lack of certainty or precision about the true size of the clinical effect and is seen in studies with too few patients. Where confidence intervals are narrow
they indicate more precise estimates of effects and a larger sample of patients studied. It is usual to interpret a ‘95%’ confidence interval as the range of effects within which we are 95% confident that the true effect lies.

**Confounding factor**

Something that influences a study and can contribute to misleading findings if it is not understood or appropriately dealt with. For example, if a group of people exercising regularly and a group of people who do not exercise have an important age difference then any difference found in outcomes about heart disease could well be due to one group being older than the other rather than due to the exercising. Age is the confounding factor here and the effect of exercising on heart disease cannot be assessed without adjusting for age differences in some way.

**Consensus methods**

A variety of techniques that aim to reach an agreement on a particular issue. Formal consensus methods include Delphi and nominal group techniques, and consensus development conferences. In the development of clinical guidelines, consensus methods may be used where there is a lack of strong research evidence on a particular topic.

**Consensus statement**

A statement of the advised course of action in relation to a particular clinical topic, based on the collective views of a body of experts.

**Considered judgement**

The application of the collective knowledge of a guideline development group to a body of evidence, to assess its applicability to the target population and the strength of any recommendation that it would support. Consistency The extent to which the conclusions of a collection of studies used to support a guideline recommendation are in agreement with each other.

**Control group**

A group of patients recruited into a study that receives no treatment, a treatment of known effect, or a placebo (dummy treatment) - in order to provide a comparison for a group receiving an experimental treatment, such as a new drug.

**Cost benefit analysis**

A type of economic evaluation where both costs and benefits of health care treatment are measured in the same monetary units. If benefits exceed costs, the evaluation would recommend providing the treatment.

**Cost effectiveness**

A type of economic evaluation that assesses the additional costs and benefits of doing something different. In cost effectiveness analysis, the costs and benefits of different treatments are compared. When a new treatment is compared with current care, its additional costs divided by its additional benefits is called the cost effectiveness ratio. Benefits are measured in natural units, for example, cost per additional heart attack prevented.

**Cost utility analysis**

A special form of cost effectiveness analysis where benefit is measured in quality adjusted life years. A treatment is assessed in terms of its ability to extend or improve the quality of life.

**Cross-sectional study**

The observation of a defined set of people at a single point in time or time period – a snapshot. (This type of study contrasts with a longitudinal study which follows a set of people over a period of time.)

**Declaration of interest**

A process by which members of a working group or committee ‘declare’ any personal or professional involvement with a company (or related to a technology) that might affect their objectivity e.g. if their position or department is funded by a pharmaceutical company.

**Double blind study**

A study in which neither the subject (patient) nor the observer (investigator/clinician) is aware of which treatment or intervention the subject is receiving. The purpose of blinding is to protect against bias.

**Economic evaluation**

Comparative analysis of alternative courses of action in terms of both their costs and consequences.
<table>
<thead>
<tr>
<th>Term</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>Efficacy</td>
<td>The extent to which a specific treatment or intervention, under ideally controlled conditions (e.g. in a laboratory), has a beneficial effect on the course or outcome of disease compared to no treatment or other routine care.</td>
</tr>
<tr>
<td>Elective</td>
<td>Name for clinical procedures that are regarded as advantageous to the patient but not urgent.</td>
</tr>
<tr>
<td>Epidemiology</td>
<td>Study of diseases within a population, covering the causes and means of prevention.</td>
</tr>
<tr>
<td>Evidence based</td>
<td>The process of systematically finding, appraising, and using research findings as the basis for clinical decisions.</td>
</tr>
<tr>
<td>Evidence-based clinical practice</td>
<td>Evidence-based clinical practice involves making decisions about the care of individual patients based on the best research evidence available rather than basing decisions on personal opinions or common practice (which may not always be evidence based). Evidence-based clinical practice therefore involves integrating individual clinical expertise and patient preferences with the best available evidence from research.</td>
</tr>
<tr>
<td>Evidence table</td>
<td>A table summarising the results of a collection of studies which, taken together, represent the evidence supporting a particular recommendation or series of recommendations in a guideline.</td>
</tr>
<tr>
<td>External validity</td>
<td>The degree to which the results of a study hold true in non-study situations, e.g. in routine clinical practice. May also be referred to as the generalisability of study results to non-study patients or populations.</td>
</tr>
<tr>
<td>Extrapolation</td>
<td>The application of research evidence based on studies of a specific population to another population with similar characteristics.</td>
</tr>
<tr>
<td>Forest plot</td>
<td>A graphical display of results from individual studies on a common scale, allowing visual comparison of results and examination of the degree of heterogeneity between studies.</td>
</tr>
<tr>
<td>Generalisability</td>
<td>The extent to which the results of a study hold true for a population of patients beyond those who participated in the research. See also External validity.</td>
</tr>
<tr>
<td>Gold standard</td>
<td>A method, procedure or measurement that is widely accepted as being the best available.</td>
</tr>
<tr>
<td>Good practice point</td>
<td>Recommended good practice based on the expert experience of the guideline development group (and possibly incorporating the expertise of a wider reference group). A guideline development group may produce a ‘Good practice point’ (rather than an evidence based recommendation) on an important topic when there is a lack of research evidence.</td>
</tr>
<tr>
<td>Grade of recommendation</td>
<td>A code (e.g. A,B,C,D) linked to a guideline recommendation, indicating the strength of the evidence supporting that recommendation.</td>
</tr>
<tr>
<td>Grey literature</td>
<td>Reports that are unpublished or have limited distribution, and are not included in bibliographic retrieval systems.</td>
</tr>
<tr>
<td>Guideline</td>
<td>A systematically developed tool which describes aspects of a patient’s condition and the care to be given. A good guideline makes recommendations about treatment and care, based on the best research available, rather than opinion. It is used to assist clinician and patient decision-making about appropriate health care for specific clinical conditions.</td>
</tr>
<tr>
<td>Guideline recommendation</td>
<td>Course of action advised by the guideline development group on the basis of their assessment of the supporting evidence.</td>
</tr>
<tr>
<td>Health economics</td>
<td>A field of conventional economics which examines the benefits of health care.</td>
</tr>
</tbody>
</table>
interventions (e.g. medicines) compared with their financial costs.

Health technology
Health technologies include medicines, medical devices such as artificial hip joints, diagnostic techniques, surgical procedures, health promotion activities (e.g. the role of diet versus medicines in disease management) and other therapeutic interventions.

Health Technology Appraisal (HTA)
A health technology appraisal, as undertaken by NICE, is the process of determining the clinical and cost effectiveness of a health technology. NICE health technology appraisals are designed to provide patients, health professionals and managers with an authoritative source of advice on new and existing health technologies.

Heterogeneity
Or lack of homogeneity. The term is used in meta-analyses and systematic reviews when the results or estimates of effects of treatment from separate studies seem to be very different – in terms of the size of treatment effects or even to the extent that some indicate beneficial and others suggest adverse treatment effects. Such results may occur as a result of differences between studies in terms of the patient populations, outcome measures, definition of variables or duration of follow-up.

Hierarchy of evidence
An established hierarchy of study types, based on the degree of certainty that can be attributed to the conclusions that can be drawn from a well conducted study. Well-conducted randomised controlled trials (RCTs) are at the top of this hierarchy. (Several large statistically significant RCTs which are in agreement represent stronger evidence than say one small RCT.) Well-conducted studies of patients' views and experiences would appear at a lower level in the hierarchy of evidence.

Homogeneity
This means that the results of studies included in a systematic review or meta analysis are similar and there is no evidence of heterogeneity. Results are usually regarded as homogeneous when differences between studies could reasonably be expected to occur by chance. See also Consistency.

Information bias
Pertinent to all types of study and can be caused by inadequate questionnaires (e.g. difficult or biased questions), observer or interviewer errors (e.g. lack of blinding), response errors (e.g. lack of blinding if patients are aware of the treatment they receive) and measurement error (e.g. a faulty machine).

Intention to treat analysis
An analysis of a clinical trial where patients are analysed according to the group to which they were initially randomly allocated, regardless of whether or not they had dropped out, fully complied with the treatment, or crossed over and received the alternative treatment. Intention-to-treat analyses are favoured in assessments of clinical effectiveness as they mirror the non-compliance and treatment changes that are likely to occur when the treatment is used in practice.

Internal validity
Refers to the integrity of the study design.

Intervention
Healthcare action intended to benefit the patient, e.g. drug treatment, surgical procedure, psychological therapy, etc.

Level of evidence
A code (e.g. 1a, 1b) linked to an individual study, indicating where it fits into the hierarchy of evidence and how well it has adhered to recognised research principles.

Literature review
A process of collecting, reading and assessing the quality of published (and unpublished) articles on a given topic.

Meta analysis
Results from a collection of independent studies (investigating the same treatment) are pooled, using statistical techniques to synthesise their findings into a single estimate of a treatment effect. Where studies are not compatible e.g. because of differences in the study populations or in the outcomes
measured, it may be inappropriate or even misleading to statistically pool results in this way. See also Systematic review and Heterogeneity.

<table>
<thead>
<tr>
<th>Methodological quality</th>
<th>The extent to which a study has conformed to recognised good practice in the design and execution of its research methods.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Multicentre study</td>
<td>A study where subjects were selected from different locations or populations, e.g. a co-operative study between different hospitals; an international collaboration involving patients from more than one country.</td>
</tr>
<tr>
<td>Necrotising enterocolitis</td>
<td>A condition in which sections of the intestine become inflamed and undergo necrosis (death of tissue). This can lead to perforation of the intestine</td>
</tr>
<tr>
<td>Non-experimental study</td>
<td>A study based on subjects selected on the basis of their availability, with no attempt having been made to avoid problems of bias.</td>
</tr>
</tbody>
</table>

**Number needed to treat (NNT)**
This measures the impact of a treatment or intervention. It states how many patients need to be treated with the treatment in question in order to prevent an event which would otherwise occur. E.g. if the NNT = 4, then 4 patients would have to be treated to prevent one bad outcome. The closer the NNT is to 1, the better the treatment is. Analogous to the NNT is the Number Needed to Harm (NNH), which is the number of patients that would need to receive a treatment to cause one additional adverse event. e.g. if the NNH = 4, then 4 patients would have to be treated for one bad outcome to occur.

**Objective measure**
A measurement that follows a standardised procedure which is less open to subjective interpretation by potentially biased observers and study participants.

**Observational study**
In research about diseases or treatments, this refers to a study in which nature is allowed to take its course. Changes or differences in one characteristic (e.g. whether or not people received a specific treatment or intervention) are studied in relation to changes or differences in other(s) (e.g. whether or not they died), without the intervention of the investigator. There is a greater risk of selection bias than in experimental studies.

**Odds ratio**
Odds are a way of representing probability, especially familiar for betting. In recent years odds ratios have become widely used in reports of clinical studies. They provide an estimate (usually with a confidence interval) for the effect of a treatment. Odds are used to convey the idea of ‘risk’ and an odds ratio of 1 between two treatment groups would imply that the risks of an adverse outcome were the same in each group. For rare events the odds ratio and the relative risk (which uses actual risks and not odds) will be very similar. See also Relative risk, Risk ratio.

**Outcome**
The end result of care and treatment and/ or rehabilitation. In other words, the change in health, functional ability, symptoms or situation of a person, which can be used to measure the effectiveness of care/ treatment/ rehabilitation. Researchers should decide what outcomes to measure before a study begins; outcomes are then assessed at the end of the study.

**Peer review**
Review of a study, service or recommendations by those with similar interests and expertise to the people who produced the study findings or recommendations. Peer reviewers can include professional and/ or patient/ carer representatives.

**Planned CS**
A CS that is scheduled before the onset of labour.

**Prognostic factor**
Patient or disease characteristics, e.g. age or co-morbidity, which influence the course of the disease under study. In a randomised trial to compare two treatments, chance imbalances in variables (prognostic factors) that influence patient outcome are possible, especially if the size of the study is fairly small. In terms of analysis these prognostic factors become confounding factors.
Prospective study

A study in which people are entered into the research and then followed up over a period of time with future events recorded as they happen. This contrasts with studies that are retrospective.

P value

If a study is done to compare two treatments then the P value is the probability of obtaining the results of that study, or something more extreme, if there really was no difference between treatments. (The assumption that there really is no difference between treatments is called the ‘null hypothesis’.) Suppose the p-value was $p = 0.03$. What this means is that if there really was no difference between treatments then there would only be a 3% chance of getting the kind of results obtained. Since this chance seems quite low we should question the validity of the assumption that there really is no difference between treatments. We would conclude that there probably is a difference between treatments. By convention, where the value of $p$ is below 0.05 (i.e. less than 5%) the result is seen as statistically significant. Where the value of $p$ is 0.001 or less, the result is seen as highly significant.

P values just tell us whether an effect can be regarded as statistically significant or not. In no way do they relate to how big the effect might be, for which we need the confidence interval.

Qualitative research

Qualitative research is used to explore and understand people’s beliefs, experiences, attitudes, behaviour and interactions. It generates non-numerical data, e.g. a patient’s description of their pain rather than a measure of pain. Qualitative research techniques such as focus groups and in depth interviews have been used in one-off projects commissioned by guideline development groups to find out more about the views and experiences of patients and carers.

Quantitative research

Research that generates numerical data or data that can be converted into numbers, for example clinical trials.

Random allocation or randomisation

A method that uses the play of chance to assign participants to comparison groups in a research study, for example, by using a random numbers table or a computer-generated random sequence. Random allocation implies that each individual (or each unit in the case of cluster randomisation) being entered into a study has the same chance of receiving each of the possible interventions.

Randomised controlled trial

A study to test a specific drug or other treatment in which people are randomly assigned to two (or more) groups: one (the experimental group) receiving the treatment that is being tested, and the other (the comparison or control group) receiving an alternative treatment, a placebo (dummy treatment) or no treatment. The two groups are followed up to compare differences in outcomes to see how effective the experimental treatment was. (Through randomisation, the groups should be similar in all aspects apart from the treatment they receive during the study.)

Relative risk

A summary measure which represents the ratio of the risk of a given event or outcome (e.g. an adverse reaction to the drug being tested) in one group of subjects compared to another group. When the ‘risk’ of the event is the same in the two groups the relative risk is 1. In a study comparing two treatments, a relative risk of 2 would indicate that patients receiving one of the treatments had twice the risk of an undesirable outcome than those receiving the other treatment. Relative risk is sometimes used as a synonym for risk ratio.

Reliability

Reliability refers to a method of measurement that consistently gives the same results. For example someone who has a high score on one occasion tends to have a high score if measured on another occasion very soon afterwards. With physical assessments it is possible for different clinicians to make independent assessments in quick succession – and if their assessments tend to agree then the method of assessment is said to be
<table>
<thead>
<tr>
<th>Term</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>Retrospective study</td>
<td>A retrospective study deals with the present/past and does not involve studying future events. This contrasts with studies that are prospective. Review Summary of the main points and trends in the research literature on a specified topic. A review is considered non-systematic unless an extensive literature search has been carried out to ensure that all aspects of the topic are covered and an objective appraisal made of the quality of the studies.</td>
</tr>
<tr>
<td>Risk ratio</td>
<td>Ratio of the risk of an undesirable event or outcome occurring in a group of patients receiving experimental treatment compared with a comparison (control) group. The term relative risk is sometimes used as a synonym of risk ratio.</td>
</tr>
<tr>
<td>Sample</td>
<td>A part of the study’s target population from which the subjects of the study will be recruited. If subjects are drawn in an unbiased way from a particular population, the results can be generalised from the sample to the population as a whole. Sampling refers to the way participants are selected for inclusion in a study.</td>
</tr>
<tr>
<td>Selection bias</td>
<td>Selection bias has occurred if:</td>
</tr>
<tr>
<td></td>
<td>a) the characteristics of the sample differ from those of the wider population from which the sample has been drawn OR</td>
</tr>
<tr>
<td></td>
<td>b) there are systematic differences between comparison groups of patients in a study in terms of prognosis or responsiveness to treatment.</td>
</tr>
<tr>
<td>Selection criteria</td>
<td>Explicit standards used by guideline development groups to decide which studies should be included and excluded from consideration as potential sources of evidence.</td>
</tr>
<tr>
<td>Semi-structured interview</td>
<td>Structured interviews involve asking people pre-set questions. A semi-structured interview allows more flexibility than a structured interview. The interviewer asks a number of open-ended questions, following up areas of interest in response to the information given by the respondent.</td>
</tr>
<tr>
<td>Statistical power</td>
<td>The ability of a study to demonstrate an association or causal relationship between two variables, given that an association exists. For example, 80% power in a clinical trial means that the study has a 80% chance of ending up with a p value of less than 5% in a statistical test (i.e. a statistically significant treatment effect) if there really was an important difference (e.g. 10% versus 5% mortality) between treatments. If the statistical power of a study is low, the study results will be questionable (the study might have been too small to detect any differences). By convention, 80% is an acceptable level of power. See also p value.</td>
</tr>
<tr>
<td>Structured interview</td>
<td>A research technique where the interviewer controls the interview by adhering strictly to a questionnaire or interview schedule with pre-set questions.</td>
</tr>
<tr>
<td>Study population</td>
<td>People who have been identified as the subjects of a study.</td>
</tr>
<tr>
<td>Survey</td>
<td>A study in which information is systematically collected from people (usually from a sample within a defined population).</td>
</tr>
<tr>
<td>Systematic review</td>
<td>A review in which evidence from scientific studies has been identified, appraised and synthesised in a methodical way according to predetermined criteria. May or may not include a meta-analysis.</td>
</tr>
</tbody>
</table>
| Target population             | The people to whom guideline recommendations are intended to apply. Recommendations may be less valid if applied to a population with different characteristics from the participants in the research study – e.g. in terms of reliable.
age, disease state, social background.

Validity
Assessment of how well a tool or instrument measures what it is intended to measure.

1
2 Appendices A–K are in a separate file.