1 JULY 2012 – 30 JUNE 2013

QUEENSLAND PERINATAL DATA COLLECTION (PDC)

Manual of Instructions
for the completion and notification of births to
the Perinatal Data Collection

DATA COLLECTIONS UNIT (DCU)
QUEENSLAND HEALTH
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<td>Health Statistics Centre</td>
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<td>ICD-10-AM</td>
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1 THE MANUAL

1.1 PURPOSE

This Instruction Manual describes the data items that are collected as part of the Queensland Perinatal Data Collection (PDC). It is intended to be a reference for all public hospitals, private hospitals, and private midwifery or medical practitioners who deliver babies outside hospitals, as well as Health Service Districts and Division personnel who are involved in the collection and use of perinatal data.

1.2 PAPER FORMS VS ELECTRONIC EXTRACT

All data providers should use this manual, whether using the paper forms (MR63D), the Perinatal Online application (PNO) or providing an electronic extract.

For specific instructions on how to complete the PNO, refer to the Perinatal Online User manual or the Perinatal Online Administration manual.

Where differences occur between the electronic system used and Queensland Health’s Data Collections Unit (DCU) requirements, the data extracted should be mapped or grouped to meet the DCU file format and requirements.

1.3 MAINTENANCE OF THE MANUAL

It is important that the information in this Manual is updated with any changes forwarded by the Data Collections Unit so that the Manual remains a relevant and up-to-date reference for contributors to and managers of the Collection, and for users of the data.

Amendments to the Collection form (MR63D) may need to be made to reflect changes in legislation, standards and policies, and therefore the Instruction Manual will also need to be updated accordingly. Any such changes are likely to occur each financial year.

If you have any queries or questions relating to this document or to the Perinatal Data Collection, please contact the Data Collection Coordinator (details below).

If you require any further copies of this Manual, also contact the Data Collection Coordinator.

Data Collection Liaison Officer
Perinatal Data Collection
Data Collections Unit
Health Statistics Centre
Queensland Health
GPO Box 48
Brisbane Qld 4001

Telephone: (07) 3235 4359
Facsimile: (07) 3234 0279

Email: perimail@health.qld.gov.au
1.4 ACKNOWLEDGMENTS

Definitions have been taken from the Queensland Health Data Dictionary (QHDD) and the National Health Data Dictionary (NHDD) as prepared by Queensland Health and the Australian Institute of Health and Welfare (AIHW) where applicable to this Collection.

We would like to thank all the midwives and medical practitioners who complete the Perinatal Data Collection (MR63D) form as well as input and submit data electronically using Perinatal Online Form or via other electronic systems.
2 INTRODUCTION

2.1 BACKGROUND

The Health Act 1937–1988 was replaced by the Public Health Act 2005. Chapter 6, Part 1 - Perinatal Statistics includes a requirement that perinatal data be provided to the Chief Executive of Queensland Health for every baby born in Queensland. The Queensland Perinatal Data Collection commenced in November 1986.

2.2 REQUIREMENTS

The Perinatal Data Collection Form (MR63D) is required to be completed (or in the case of hospitals providing electronic extracts, an extract is required) by all public hospitals, private hospitals, and private midwifery or medical practitioners who deliver babies outside hospitals, for all births occurring in Queensland. The scope of the Collection includes all live births, and stillbirths of at least 20 weeks gestation and/or at least 400 grams in weight. Information relating to neonatal morbidity is collected up until the baby is discharged from the birth admission or up until the baby reaches 28 days of age.

The quality of information produced from the PDC depends on the accurate, consistent and timely completion of the forms. Completed forms and electronic extracts are validated and queries relating to missing, contradictory or ambiguous data are directed back to the hospital or independent practitioner.

2.3 AIMS OF THE PERINATAL DATA COLLECTION

The aims of the PDC are to monitor patterns of obstetric and neonatal practice in the State and to provide statistical information on specific topics within these fields to assist with the planning of Queensland Health services. It is also intended to be a basic source of information for research in obstetric and neonatal care and to be used in the education of students of midwifery and medicine.

In addition to information collected via the perinatal data forms and via electronic extracts, details from Certificates of Perinatal Death, Histopathology reports and post mortem reports supplement the Collection.

2.4 CONFIDENTIALITY OF DATA

All unit record information collected by Data Collections Unit is treated as strictly confidential. All information collected is used for statistical purposes only.

Data Collections Unit adheres to Information Standard IS42A which requires personal information to be managed in accordance with National Privacy Principals.

2.5 PERINATAL STATISTICS AND PUBLICATIONS

The Health Statistics Centre (HSC) releases an annual report presenting summary statistics based on the data collected via the PDC. This report is available on QHEPS:


or via the following website:

- [http://www.health.qld.gov.au](http://www.health.qld.gov.au) - use the search engine and the terms “health statistics centre” and follow the prompts to publications and then perinatal.
Through the National Perinatal Epidemiology and Statistics Unit (NPESU) of the AIHW, Queensland data is used in the compilation of Australia-wide figures and can be compared with perinatal statistics from other States and Territories.

Data is also available via request, on an ad hoc or regular basis, from the Statistical Output Unit (SOU) within HSC. The release of data is governed by patient confidentiality legislation in the Public Health Act 2005. Requests for data should be made via e-mail to HlthStat@health.qld.gov.au or by phoning (07) 3234 1875. (Note that in some instances charges may apply – contact SOU for further details).

2.6 THE FORM

The form is designed to be an integral part of the obstetric record, both to reduce duplication of recording and to ensure optimum accuracy of data. The hospital copies can be used as a summary for the patient’s chart and this includes some items which are not essential for the PDC but may be useful in hospitals. Items not needed specifically for the PDC but included for hospitals’ use are not highlighted white on the hospital copies and have been marked with an asterisk (*) in this Manual.

Currently, there is no distinction in PNO between mandatory and non-mandatory fields.

2.6.1 PERINATAL DATA COLLECTION FORM (MR63D) (SEE APPENDIX B)

From 1 July 2007, the MR63D form is supplied as an A3 size sheet which will fold in half to A4 size for placement within the medical record. The MR66 Congenital Anomaly form has been subsumed into the MR63D form. This form consists of three sheets – an original and two duplicates:

- The original (green) must be retained for your own hospital records and should be referred to when clarifying or confirming queries.
- The first duplicate (green) may be placed in the baby’s chart or forwarded to the private medical practitioner or Child Health Nurse. This is left to the discretion of individual hospitals.
- The second duplicate (white) is to be returned to Data Collections Unit within 35 days of the baby’s birth.

2.7 DISPATCH OF FORMS

Instructions for the dispatch of the Data Collections Unit copies of the MR63D forms are included in Appendix A. These forms should be forwarded to the Data Collections Unit within 35 days of the birth of a baby. Hospitals should dispatch the returns on a fortnightly or monthly basis, with an accompanying Dispatch Cover Note (see Appendix A).

Facilities using PNO may refer to the Perinatal Online Administration manual for details on the extraction of data.

2.8 ELECTRONIC TRANSFER OF DATA

For facilities providing data via electronic extract, please contact PDC to obtain the most current file format required (see Appendix C for example file format, current at time of publication). Prior to providing an electronic extract of data to PDC, individual facilities should contact the Principal Data Collection Officer, Joanne Bunney, phone (07) 3237
1464 or via e-mail, joanne.bunney@health.qld.gov.au. Extracts are required within 35 days of the birth of a baby.
3 GENERAL INSTRUCTIONS

3.1 COMPLETING THE FORMS

- Please PRINT clearly using a ballpoint pen (not a felt pen) and press firmly.

- The paper has been carbonised so please take care not to write on paper placed over these forms, or place undue sharp pressure on the original.

- If an error is made on the form, it is preferable to cross through the incorrect response and rewrite the answer, rather than overwriting the original answer, as this is easier to read, and reduces errors in interpretation.

- Please enter the appropriate information in the areas provided, or tick the appropriate boxes. If the boxes do not provide the appropriate alternative, please specify details under ‘Other’ in the space provided.

- Using a question mark (?) on the form to indicate that a condition is suspected will always generate a query to confirm the suspected condition. Wherever possible please confirm prior to reporting. If the diagnosis can not be confirmed, indicate this also on the form by writing beside the condition ‘unable to be confirmed’.

- The forms should be as complete as possible. Do not leave any fields blank. If any details are unknown the best estimate should be used, or ‘not known’ written beside the missing item.

- In the case of multiple births, a separate form should be completed for each baby. For example, in the case of twins, two forms are to be completed, identifying each twin as Twin I and Twin II. The Data Collections Unit copies should be pinned together so that common information need not be completed on the second form. Details in the LABOUR AND DELIVERY, BABY, POSTNATAL and BABY DISCHARGE DETAILS sections are required for each baby.

- If the baby is transferred to another hospital after birth, please complete the form and document the transfer destination so that, if necessary, further enquiries can be made about congenital anomalies.

- The items marked with an asterisk (*) are for hospital use only and do not form part of the information processed for the PDC. These items are not highlighted white on the hospital copies of the form.

- Currently, there is no distinction in PNO between mandatory and non-mandatory fields.
4 MOTHER’S DETAILS

All items contained in this section of the form must be completed clearly. Wherever possible, it is preferred that printed labels be used to provide maternal details and to identify the MR63D forms, however this is not mandatory.

If used on the original and duplicate copies, labels should be placed in the upper right hand corner, ensuring that no other information is obscured. If an identification label is used only on the hospital copies (and not the duplicates), DO NOT FORGET to complete MOTHER’S USUAL RESIDENCE, DATE OF BIRTH, NAMES and UR NUMBER on the second duplicate (i.e. the Data Collections Unit copy).

4.1 PLACE OF DELIVERY

PLACE OF DELIVERY

Enter the name of the hospital where the birth occurred. Where both public and private facilities exist please specify (eg Mater Mothers Public or Mater Mothers Private).

For births notified by a hospital but not delivered in the hospital (eg Born before arrival (BBA) or home birth), enter the name of the hospital completing the form. If a home birth is notified by the accoucheur, write ‘Home’ and complete the details on the reverse side of the Data Collections Unit copy.

This field allows the Data Collections Unit to follow up queries concerning missing or inconsistent data. It also enables individual hospitals to receive feedback on the data they record on the form.

4.2 DATE OF ADMISSION

Enter the day, month and year of the date of admission of the mother for delivery using all boxes, eg 1 November 2012 should be entered as:

DATE OF ADMISSION

(0112012)

For this Collection, record the date of admission for the delivery to the facility where the delivery takes place. For planned home births where the baby is not admitted to a hospital, this field is not required.

4.3 MOTHER’S COUNTRY OF BIRTH

MOTHER’S COUNTRY OF BIRTH

Enter the country of birth of the mother. Be as specific as possible, eg. enter Zimbabwe rather than Africa.

Ethnicity is an important concept, both in the study of disease patterns and the need for and provision of services. Country of birth is the most easily collected and consistently
reported of possible ethnicity data items. It is recognised that country of birth is one of a number of surrogate measures for ethnicity.

4.4 INDIGENOUS STATUS

Tick the box (one box only) that corresponds to the Indigenous Status of the mother.

Note that a mother’s indigenous status cannot be determined simply by observation and therefore this question must be asked of all mothers. For further information regarding determining Indigenous status, please refer to the ‘Are you of Aboriginal or Torres Strait Islander origin?’ pamphlet. If you require copies of this publication, please contact the National Centre for Aboriginal and Torres Strait Islander Statistics (Australian Bureau of Statistics) on the free call number 1800 633 216.

Definitions:
An Aboriginal or Torres Strait Islander is a person of Aboriginal or Torres Strait Islander descent who identifies as an Aboriginal or Torres Strait Islander and is accepted as such by the community in which she lives.

- **Aboriginal**
  Aboriginal but not Torres Strait Islander origin.

- **Torres Strait Islander**
  Torres Strait Islander but not Aboriginal origin.

- **Aboriginal and Torres Strait Islander**
  Both Aboriginal and Torres Strait Islander origin.

- **Neither Aboriginal nor Torres Strait Islander**
  Neither Aboriginal nor Torres Strait Islander origin.

Given the gross inequalities in health status between Indigenous and Non-indigenous peoples in Australia, the size of the Aboriginal and Torres Strait Islander populations and their historical and political context, there is a strong case for ensuring that information on Indigenous status is collected for planning and service delivery purposes and for monitoring Aboriginal and Torres Strait Islander health.
4.5 MARITAL STATUS

Tick the box (one box only) that corresponds to the marital status of the mother.

Marital status is a core data element in a wide range of social, labour and demographic statistics. Its main purpose is to establish the living arrangements of individuals, to facilitate analysis of the association of marital status with the need for and use of services and for epidemiological analysis.

4.6 ACCOMMODATION STATUS OF MOTHER

Tick the box (one box only) that corresponds to the type of ward accommodation the mother has elected to be accommodated in regardless of the method of payment for the hospital admission. This item does not indicate the insurance status of the mother.

For home births where the baby is not admitted to a hospital, this field is not required.

Definitions:

- **Public**
  A public patient is a person, eligible for Medicare, who, on admission to a recognised hospital or soon after:
  - receives a public hospital service free of charge; or
  - elects to be a public patient; or
  - whose treatment is contracted to a private hospital.

- **Private**
  A private patient is a person who, on admission to a recognised hospital or soon after:
  - elects to be a private patient treated by a medical practitioner of her own choice; or
  - elects to occupy a bed in a single room (where such an election is made, the patient is responsible for meeting certain hospital charges as well as the professional charges raised by any treating medical practitioner); or
  - a person, eligible for Medicare, who chooses to be admitted to a private hospital (where such a choice is made, the patient is responsible for meeting all hospital charges as well as the professional charges raised by any treating medical practitioner).
Note that ineligible and compensable patients who are chargeable but use public hospital doctors are classified as public. Those who use private doctors are to be classified as private.

4.7 SEROLOGY*

This field is not mandatory, however if results reported in this field affect the management of the pregnancy, please report the associated condition in Medical Conditions (see 6.5) or Pregnancy Complications (see 6.6).

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<th>SEROLOGY</th>
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<tr>
<td>RPR</td>
<td>IgG</td>
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<tr>
<td>Rubella</td>
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<td>Blood group</td>
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</tr>
<tr>
<td>Rh</td>
<td></td>
</tr>
<tr>
<td>Antibodies</td>
<td>Yes □ No □</td>
</tr>
<tr>
<td>Other</td>
<td></td>
</tr>
</tbody>
</table>

RPR…...IgG…. Enter ‘Pos’ or ‘Neg’ in both fields to show RPR and IgG status
Rubella Enter immune or not immune
Blood group Enter blood group, eg ‘O’, ‘A’, ‘B’ or ‘AB’
Rh Enter the Rhesus factor (+ or -)
Antibodies Tick the appropriate box for ‘Yes’ or ‘No’

4.8 MOTHER FAMILY NAME

The mother’s full family name should be recorded.
If family name is not known or cannot be established, record UNKNOWN.

Some people do not have a family name and a given name and they have only one name by which they are known. If the mother has only one name, record it as the family name.

<table>
<thead>
<tr>
<th>FAMILY NAME</th>
</tr>
</thead>
<tbody>
<tr>
<td>1ST GIVEN NAME</td>
</tr>
<tr>
<td>2ND GIVEN NAME</td>
</tr>
</tbody>
</table>

The use of hospital labels is the preferred method to identify forms, as long as they contain all of the relevant information, as it reduces errors in transcription of written information (such as UR numbers and Date of Birth).

4.9 GIVEN NAMES

A mother may have more than one given name.

A mother’s given name(s) should be recorded. Where applicable it is essential that the given names are recorded for the first two recorded given names of a mother.

If given name is not known or cannot be established, record UNKNOWN.
Some people do not have a family name and a given name and they have only one name by which they are known. If the mother has only one name, record it as the family name.

The use of hospital labels is the preferred method to identify forms, as long as they contain all of the relevant information, as it reduces errors in transcription of written information (such as UR numbers and Date of Birth).

### 4.10 UR NUMBER

Enter the Unit Record (UR) number assigned to the mother (if applicable).

| UR No. | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |

For home births where the baby is not admitted to a hospital, this field is not required, however, if the private midwifery practitioner assigns a record number for administrative purposes it can be included.

Confidentiality of data is maintained through the storage of this data in a separate table by PDC, with limited access. PDC adhere to Queensland Health’s confidentiality of data standards including IS42A.

### 4.11 DATE OF BIRTH (MOTHER)

Record the date of birth of the mother using the full date (ie. ddmmyyyy) and leading zeros where necessary. Example: 10 January 1985 should be entered as:

| DOB | 1 0 0 1 1 9 8 5 |

If the day of birth is unknown, use 15.
If the month of birth is unknown, use 06.
If the year of birth is unknown, estimate the year from the age of the mother.
If the age of the mother is unknown and it is not possible to estimate an age and hence a year of birth (eg. for unconscious mothers, use the year 1900)

Example: If a mother is admitted in 2012 and does not know her exact date of birth but knows that she is 30 years of age, record the date of birth as follows:

| DOB | 1 5 0 6 1 9 8 0 |

Although provision is made for recording an unknown date of birth (using 15/06/1900), every effort should be made during the course of the admission to determine (and record) the mother’s actual date of birth. The mother’s date of birth is an important requirement for the correct identification of the individual.
4.12 ESTIMATED DATE OF BIRTH FLAG (MOTHER)

The Estimated Date of Birth box indicates whether the mother’s date of birth has been estimated.

If an estimate has been used in place of either the day or the month or the year, then Estimated Date of Birth box must be ticked.

<table>
<thead>
<tr>
<th>DOB</th>
<th>1</th>
<th>5</th>
<th>0</th>
<th>6</th>
<th>1</th>
<th>9</th>
<th>8</th>
<th>0</th>
</tr>
</thead>
</table>

Estimated Date of Birth

4.13 ADDRESS OF USUAL RESIDENCE

The collection of the address details of a mother is critical for patient follow up and as a means of reporting information about the geographic location of the residence of a mother. A mother may have one address or many addresses. The last known address should be recorded.

Enter the street number, street name, suburb/town and postcode where the mother usually resides (not postal address). For interstate mothers, enter the address and name of the State of the mother’s usual residence.

<table>
<thead>
<tr>
<th>USUAL RESIDENCE</th>
<th>POSTCODE</th>
<th>STATE</th>
<th>SLA</th>
</tr>
</thead>
</table>

If the mother is not a resident of Australia or an Australian External Territory, or has no fixed address, use one of the following supplementary codes as the postcode of usual residence.

<table>
<thead>
<tr>
<th>Code</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>9301</td>
<td>Papua New Guinea</td>
</tr>
<tr>
<td>9302</td>
<td>New Zealand</td>
</tr>
<tr>
<td>9399</td>
<td>Overseas other (not PNG or NZ)</td>
</tr>
<tr>
<td>9799</td>
<td>At Sea</td>
</tr>
<tr>
<td>9989</td>
<td>No Fixed Address</td>
</tr>
<tr>
<td>0989</td>
<td>Not stated or unknown</td>
</tr>
</tbody>
</table>

Please note that it is particularly important to record the country of residence accurately for patients from Papua New Guinea and New Zealand.

For Australian External Territory addresses, the actual postcode and State ID is to be used from 1 July 2005, rather than a supplementary postcode and State ID. Australian External Territories include the following: Christmas Island, Cocos (Keeling) Islands, and Norfolk Island.

This information is used to determine the Statistical Local Area (SLA) of usual residence, enabling the comparison of the use of services by persons residing in different geographical areas, the characterisation of catchment areas and populations for facilities for planning purposes and the documentation of the provision of services to residents of States or Territories other than Queensland.
For those hospitals sending data electronically, please contact the CRDS Administrator on (07) 3836 0598 or via e-mail CRDS@health.qld.gov.au for a complete list of valid SLA codes.

4.14 ANTENATAL TRANSFER

**ANTENATAL TRANSFER**

(Include transfers from planned home birth to hospital, from birthing centre to acute areas etc.)

<table>
<thead>
<tr>
<th>Reason for transfer</th>
<th>Time of Transfer</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>prior to onset of labour</td>
</tr>
</tbody>
</table>

Transferred from ________________

Tick ‘Yes’ or ‘No’ to indicate whether the mother has been transferred from a different location. This includes transfers from home births to hospital, from birthing centre to acute care area.

4.14.1 REASON FOR TRANSFER

Enter the reason for the transfer of the mother from the initial location, eg ‘unavailability of medical services’, ‘premature rupture of membranes’.

Reason for transfer ________________

4.14.2 TRANSFERRED FROM

Enter the initial place of treatment that the mother has been transferred from. Enter the full name of the facility, including whether public or private where applicable, or where transferred from a home birth, enter ‘Home’.

Transferred from ________________

4.14.3 TIME OF TRANSFER

Tick whether the mother was transferred ‘prior to onset of labour’ or ‘during labour’.

Time of Transfer

- prior to onset of labour
- during labour
5  **PREVIOUS PREGNANCIES**

*Note:* This section refers to all previous pregnancies and therefore excludes the current pregnancy.

### 5.1 PREVIOUS PREGNANCIES

<table>
<thead>
<tr>
<th>PREVIOUS PREGNANCIES</th>
</tr>
</thead>
<tbody>
<tr>
<td>None (go to next section)</td>
</tr>
</tbody>
</table>

If the mother has had no previous pregnancies, tick ‘None’ and go to the next section **PRESENT PREGNANCY**. **DO NOT** complete the remaining fields in this section.

If the mother has had previous pregnancies, complete all sections in Previous Pregnancies field (5.2 – 5.4).

### 5.2 NUMBER OF PREVIOUS PREGNANCIES

<table>
<thead>
<tr>
<th>Number of previous pregnancies resulting in:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Only livebirths</td>
</tr>
<tr>
<td>Only stillbirths</td>
</tr>
<tr>
<td>Only abortions/miscarriages/ectopic/hydiform mole</td>
</tr>
<tr>
<td>Livebirth &amp; stillbirth</td>
</tr>
<tr>
<td>Livebirth &amp; abortion/miscarriage/ectopic/hydiform mole</td>
</tr>
<tr>
<td>Stillbirth &amp; abortion/miscarriage/ectopic/hydiform mole</td>
</tr>
<tr>
<td>Livebirth, stillbirth &amp; abortion/miscarriage/ectopic/hydiform mole</td>
</tr>
</tbody>
</table>

**TOTAL NUMBER of previous pregnancies**

Enter the number of **previous pregnancies** (not number of previous babies) resulting in each of:

- Only livebirths (Number of previous pregnancies resulting in livebirths only);
- Only stillbirths (Number of previous pregnancies resulting in stillbirths only);
- Only abortions/miscarriage/ectopic/hydiform mole (Number of previous pregnancies resulting in abortion/miscarriage/ectopic/hydiform mole only);
- Livebirth & stillbirth (Number of previous pregnancies resulting in an outcome of livebirth and stillbirth in the same pregnancy);
- Livebirth & abortion/miscarriage/ectopic/hydiform mole (Number of previous pregnancies resulting in an outcome of livebirth and abortion/miscarriage/ectopic/hydiform mole in the same pregnancy);
- Stillbirth & abortion/miscarriage/ectopic/hydiform mole (Number of previous pregnancies resulting in an outcome of stillbirth and abortion/miscarriage/ectopic/hydiform mole in the same pregnancy);
- Livebirth, stillbirth & abortion/miscarriage/ectopic/hydiform mole (Number of previous pregnancies resulting in an outcome of livebirth and stillbirth and abortion/miscarriage/ectopic/hydiform mole in the same pregnancy).

A tick or cross is not sufficient; the actual number of pregnancies must be recorded, even if that number is zero.

*Note:* This field refers to the **number of pregnancies, not the number of babies born**. Consequently, a pregnancy resulting in multiple births should be counted as only one pregnancy.
The total number of previous pregnancies should be entered at the bottom of the list of outcomes in the field provided. Note that the total number entered should be equal to the combined numbers entered as outcomes.

**Definitions:**

- **Live birth**
  The complete expulsion or extraction from its mother of a product of conception, irrespective of the duration of pregnancy, which, after such separation, breathes or shows any other evidence of life, such as beating of the heart, pulsation of the umbilical cord or definite movement of voluntary muscles, whether or not the umbilical cord has been cut or the placenta is attached.

- **Stillbirth**
  A fetal death prior to the complete expulsion or extraction from its mother of a product or conception of 20 or more completed weeks of gestation and/or of 400 grams or more birthweight; the death is indicated by the fact that after such separation the foetus does not breathe or show any other evidence of life, such as beating of the heart, pulsation of the umbilical cord, or definite movement of voluntary muscles.

- **Abortion/Miscarriage/Ectopic/Hydatiform mole**
  Includes spontaneous abortion (less than 20 weeks gestation and less than 400 grams birthweight); induced abortion (termination of pregnancy before 20 weeks gestation); ectopic pregnancy; or molar pregnancy.

Note, that in the case of medical abortion or termination of pregnancy where gestation is 20 weeks or greater and/or birthweight 400g or greater, the pregnancy should be recorded as determined by the outcome (i.e. live birth or stillbirth).

**5.3 METHOD OF DELIVERY OF LAST BIRTH**

<table>
<thead>
<tr>
<th>METHOD OF DELIVERY OF LAST BIRTH</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vaginal non-instrumental</td>
</tr>
<tr>
<td>Forceps</td>
</tr>
<tr>
<td>Vacuum extractor</td>
</tr>
<tr>
<td>LSCS</td>
</tr>
<tr>
<td>Classical CS</td>
</tr>
<tr>
<td>Other (specify)</td>
</tr>
</tbody>
</table>

Tick the box(es) that correspond to the method of delivery of the last birth. If a previous multiple pregnancy resulted in two or more different outcomes (e.g. vaginal non-instrumental and LSCS), tick both boxes. This should be further clarified by noting in this section that a multiple pregnancy occurred.

Note: This relates to the last birth, and therefore not necessarily the last pregnancy. For example, if the mother has had two previous pregnancies and the last pregnancy resulted in a spontaneous abortion while the pregnancy before that resulted in a lower segment caesarean birth then tick ‘LSCS’.
Method of delivery should only be provided for abortion/miscarriage when gestation is 20 weeks or greater and/or birthweight 400g or more.

(See Section 7.10 for definitions of Methods of Birth).

5.4 NUMBER OF PREVIOUS CAESAREANS

Enter the number of previous caesarean sections the mother has had. Enter zero if the mother has had no previous caesarean sections.
PRESENT PREGNANCY

6.1 LMP

Enter the day, month and year of the first day of the mother’s last menstrual period (LMP) using all boxes. For example, a LMP of 1 November 2011 should be entered as:

LMP 0;1;1;2;0;1;1

If the exact day is unknown, enter month and year as show below:

LMP ?;?;1;1;2;0;1;1

If the date of the LMP is unknown, enter ‘99 99 99’ as shown below. This may occur in cases where there is a history of abnormal or irregular periods, or a delay of ovulation has occurred following the use of the contraceptive pill.

LMP 9;9;9;9;9;9;9;9

In the case of hospitals reporting this information electronically, if only month and year are known, the day is entered as 01, 15 or 28 for early, mid or late in the month. The LMP Estimation Flag must be completed as an E for estimated. If the date is unknown, leave the field blank.

6.2 EDC

Enter the day, month and year of the best-estimated date of confinement (EDC) for this pregnancy using all boxes. For example, an EDC of 1 November 2012 should be entered as:

EDC 0;1;1;2;0;1;2

If the exact day is unknown, enter month and year as shown below:

EDC ?;?;1;1;2;0;1;2

Assessment

EDC ?;?;1;1;2;0;1;2
By US scan/dates/clinical assessment

Indicate how the EDC was determined by circling US scan, dates or clinical assessment.

If more than one EDC is available, (either by US scan, dates or clinical assessment), then record the one that has been deemed to be clinically the most reliable (i.e. the date used by the clinician, on which clinical decisions regarding the management of the pregnancy have been based).

In the case of hospitals reporting this information electronically, if only month and year are known, the day is entered as 01, 15 or 28 for early, mid or late in the month. The EDC
Estimation Flag must be completed as an E for estimated. If the date is unknown, leave the field blank.

6.3 HEIGHT

<table>
<thead>
<tr>
<th>HEIGHT</th>
<th>cm</th>
</tr>
</thead>
</table>

Record the mother’s height in total centimetres. This can either be measured or self-reported. Height will be used in conjunction with self-reported weight for Body Mass Index (BMI) assessment to assist in identifying pregnancies at risk.

6.4 WEIGHT

<table>
<thead>
<tr>
<th>WEIGHT</th>
<th>kg</th>
</tr>
</thead>
</table>

(self-reported at conception)

Record the mother’s weight in total kilograms. This will be the self-reported weight of the mother in the four to six weeks prior to or at conception. Weight will be used in conjunction with height for Body Mass Index (BMI) assessment to assist in identifying pregnancies at risk.

6.5 ANTENATAL CARE

<table>
<thead>
<tr>
<th>ANTENATAL CARE</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>(You may tick more than one box)</td>
<td></td>
</tr>
<tr>
<td>No antenatal care</td>
<td></td>
</tr>
<tr>
<td>Public hospital/clinic midwifery practitioner</td>
<td></td>
</tr>
<tr>
<td>Public hospital/clinic medical practitioner</td>
<td></td>
</tr>
<tr>
<td>General practitioner</td>
<td></td>
</tr>
<tr>
<td>Private medical practitioner</td>
<td></td>
</tr>
<tr>
<td>Private midwife practitioner</td>
<td></td>
</tr>
</tbody>
</table>

Tick the box(es) that correspond to the antenatal care received for the current pregnancy. More than one box may be ticked. If the mother received no antenatal care, tick ‘No antenatal care’.

Definitions are listed on the next page.
6.6 DID THE SAME MIDWIFE(S) WHO PROVIDED ANTENATAL CARE ALSO PROVIDE THE WOMAN’S INTRAPARTUM AND POST-DISCHARGE CARE? (PERINATAL ONLINE HOSPITALS ONLY)

Did the same Midwife(s) who provided antenatal care also provide the woman’s intrapartum and post-discharge care?

No [ ] Yes [ ]

Definition:
- **Continuity of Midwife Carer**
  Where the woman has a ‘primary’ or ‘named’ midwife(s), providing the majority of pregnancy, birth and post birth care

Did the same Midwife(s) who provided antenatal care also provide the woman’s intrapartum and post-discharge care?

Answer ‘yes’ if all of the following occurred:

a) The woman received antenatal care on a number of occasions by the named midwife or small group of 2 – 4 midwives to the extent that they could be considered ‘known’ by the woman;
b) The woman was cared for in labour/birth by the named or one of the small group of midwives; and
c) It is embedded in the maternity service that the named or one of the small group of midwives will provide the postnatal care to the woman after discharge.

If you require further assistance with this data field please phone the Maternity, Primary Community & Extended Care Branch on (07) 3234 0691.
### 6.7 TOTAL NUMBER OF VISITS

<table>
<thead>
<tr>
<th>TOTAL NUMBER OF VISITS</th>
<th></th>
</tr>
</thead>
</table>

Enter the total number of antenatal visits for the current pregnancy. This information can be obtained from the case notes (hospital clinic patients) or by asking the mother. The question is designed to measure the amount of supervision in the current pregnancy.

Note that if more than one type of antenatal care has been provided please report the total number of visits for the pregnancy, not just those provided at the reporting facility.
### 6.8 CURRENT MEDICAL CONDITIONS

**CURRENT MEDICAL CONDITIONS**

You may tick more than one box

- None
- Essential hypertension
- Pre-existing diabetes mellitus
  - insulin treated
  - oral hypoglycaemic therapy
  - other
- Asthma (treated during this pregnancy)
- Epilepsy
- Genital herpes (active during this pregnancy)
- Anaemia
- Renal condition (specify) ________________
- Cardiac condition (specify) ________________
- Hepatitis B Active
- Hepatitis B Carrier
- Hepatitis C Active
- Hepatitis C Carrier
- Other (specify) ________________

Tick the box(es) that correspond to any medical conditions the mother has which may significantly affect the current pregnancy or its management, or document the condition(s) in the space provided (see Appendix D for examples). If the mother has no current medical conditions, tick ‘None’. Where ‘Renal condition’, ‘Cardiac condition’ or ‘Other’ is ticked, please provide as much detail as possible to allow an appropriate morbidity code to be assigned.

**Definition:**

- **Current medical conditions**
  Includes pre-existing maternal conditions, hypertension or diabetes, and other diseases, illnesses or conditions arising during the current pregnancy, that are not directly attributable to pregnancy but may significantly affect care during the current pregnancy and/or pregnancy outcome.

- **Pre-existing diabetes mellitus**
  Diabetes pre-existing prior to pregnancy. Indicate whether insulin treated, oral hypoglycaemic therapy treated or other (includes diet, exercise, lifestyle management).
6.9  GESTATION AT FIRST ANTENATAL VISIT

<table>
<thead>
<tr>
<th>GESTATION AT FIRST ANTENATAL VISIT</th>
<th></th>
<th>Weeks</th>
</tr>
</thead>
</table>

Record the number of completed weeks of the current pregnancy when the mother had her first contact for antenatal care. The first contact for antenatal care is the first contact with a doctor or nurse where actual pre-birth maternity care was provided. It does not include a contact if it was to confirm the pregnancy only or those contacts that occurred during the pregnancy that related to other non pregnancy related issues.

6.10  PREGNANCY COMPLICATIONS

<table>
<thead>
<tr>
<th>PREGNANCY COMPLICATIONS</th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>None</td>
<td></td>
<td></td>
</tr>
<tr>
<td>APH (&lt;20 weeks)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>APH (20 weeks or later) due to</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• abruption</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• placenta praevia</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• other</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Gestational diabetes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• insulin treated</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• oral hypoglycaemic therapy</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• other</td>
<td></td>
<td></td>
</tr>
<tr>
<td>PIH/PE</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• mild</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• moderate</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• severe</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Other (specify)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Tick the box(es) that correspond to any complications of the current pregnancy. If there are complications other than those listed, tick ‘Other’ and specify the complication(s) in the space provided (see Appendix D for examples). If there are no pregnancy complications, tick ‘None’.

Definitions are listed on the next page.
6.11 SMOKING

SMOKING

During the first 20 weeks of pregnancy
Did the mother smoke? No ☐ Yes ☐
If yes, how many cigarettes per day? ☐ ☐ ☐
Was smoking cessation advice offered by a health care provider? No ☐ Yes ☐

After 20 weeks of pregnancy
Did the mother smoke? No ☐ Yes ☐
If yes, how many cigarettes per day? ☐ ☐ ☐
Was smoking cessation advice offered by a health care provider? No ☐ Yes ☐

Smoking during the first 20 weeks of pregnancy

Tick the box that corresponds to the mother’s smoking status during the first 20 weeks of pregnancy.

If the mother smoked at all during the first 20 weeks of pregnancy, record the number of cigarettes smoked per day.

Next, tick the box that indicates whether the mother was offered smoking cessation advice by a health care provider at any time during the first 20 weeks of pregnancy.

Definitions:
- Pregnancy complications
  Complications of pregnancy arising up to the period immediately preceding labour and delivery that are directly attributable to the pregnancy and may significantly affect care during the current pregnancy and/or the outcome.

- APH (Antepartum haemorrhage)
  - Abruptio
    Abruptio placenta. An antepartum haemorrhage resulting from the placenta becoming totally or partially detached from the uterine wall whilst the foetus is still in utero.
  - Placenta praevia
    An antepartum haemorrhage resulting from the placenta being located over or very near to the internal os.
  - Other
    Any other antepartum haemorrhage, or cause unknown.

- Gestational diabetes
  Diabetes specifically occurring during pregnancy. Indicate whether insulin treated, oral hypoglycaemic therapy treated or other (includes diet, exercise, lifestyle management).

- PIH/PE
  Pregnancy Induced Hypertension/Pre-Eclampsia. Indicate whether mild, moderate or severe.
Smoking cessation advice can include anything from a stop smoking pamphlet included in an antenatal package/visit, through to a full stop smoking program.

**Smoking after 20 weeks of pregnancy**

Tick the box that corresponds to the mother’s smoking status after 20 weeks of pregnancy.

If the mother smoked at all after 20 weeks of pregnancy, record the number of cigarettes smoked per day.

Next, tick the box that indicates whether the mother was offered smoking cessation advice by a health care provider after the first 20 weeks of pregnancy. Smoking cessation advice can include anything from a stop smoking pamphlet included in an antenatal package/visit, through to a full stop smoking program.

Cigarette smoking is the most important modifiable risk factor for preterm birth, which is the strongest predictor of perinatal death and disability.

### 6.12 PROCEDURES AND OPERATIONS

<table>
<thead>
<tr>
<th>PROCEDURES AND OPERATIONS (during pregnancy, labour and delivery)</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>None</td>
<td>☐</td>
</tr>
<tr>
<td>Chorionic Villus Sampling</td>
<td>☐</td>
</tr>
<tr>
<td>Amniocentesis (diagnostic)</td>
<td>☐</td>
</tr>
<tr>
<td>Cordocentesis</td>
<td>☐</td>
</tr>
<tr>
<td>Cervical suture (for cervical incompetence)</td>
<td>☐</td>
</tr>
<tr>
<td>Other (specify)</td>
<td>☐</td>
</tr>
</tbody>
</table>

Tick the boxes that correspond to any medical or surgical procedures and/or operations that were performed on the mother or foetus during the current pregnancy. Please also include those performed during labour and delivery. If a procedure and/or operation was performed other than those listed, tick ‘Other’ and specify in the space provided (see Appendix D for examples). If no procedures or operations were performed during this pregnancy, tick ‘None’. Where procedures are reported that may be performed via different approaches please provide as many details as possible.

For example: cholecystectomy, which may be open or via laparoscope please report as either ‘open cholecystectomy’ or ‘laparoscopic cholecystectomy’. 
6.13 NUMBER OF ULTRASOUND SCANS

<table>
<thead>
<tr>
<th>ULTRASOUNDS</th>
<th>Number of scans</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nuchal translucency ultrasound</td>
<td>No</td>
</tr>
<tr>
<td>Morphology ultrasound scan</td>
<td>No</td>
</tr>
<tr>
<td>Assessment for chorionicity scan</td>
<td>No</td>
</tr>
</tbody>
</table>

Enter the number of ultrasound scans performed during the current pregnancy. Enter zero if no ultrasound scans were performed.

This number indicates the total number of obstetric ultrasound scans performed during the current pregnancy. This will therefore include those performed by a radiographer in a recognised medical imaging unit and/or those performed by a health care professional(s) (e.g., Doctor or Midwife) in a variety of health care settings including hospital wards, community clinics or the premises of private practitioners.

Note that it does not include other non-obstetric ultrasounds (e.g., maternal renal, or gallbladder scan) and may necessitate asking the mother for confirmation of the number, as not all ultrasounds performed will have a written report.

6.14 TYPES OF ULTRASOUND SCANS

<table>
<thead>
<tr>
<th>ULTRASOUNDS</th>
<th>Number of scans</th>
</tr>
</thead>
<tbody>
<tr>
<td>Were any of the following performed?</td>
<td></td>
</tr>
<tr>
<td>Nuchal translucency ultrasound</td>
<td>No</td>
</tr>
<tr>
<td>Morphology ultrasound scan</td>
<td>No</td>
</tr>
<tr>
<td>Assessment for chorionicity scan</td>
<td>No</td>
</tr>
</tbody>
</table>

Indicate if a nuchal translucency scan was performed.
Indicate if a morphology ultrasound scan was performed.
Indicate if an assessment for chorionicity scan was performed.

Definitions:

**Nuchal translucency:**
An ultrasound to assess for major chromosomal abnormalities.

**Morphology:**
An ultrasound to allow the early diagnosis of morphologic abnormalities.

**Chorionicity:**
An ultrasound to distinguish between twins who share a membrane. This will identify those multiples who share a chorion and are at risk of twin to twin transfusion syndrome.
6.15 **ASSISTED CONCEPTION**

Was this pregnancy the result of assisted conception?

<p>| | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>No</td>
<td>Yes</td>
</tr>
</tbody>
</table>

If yes, indicate method(s) used

- AIH/AID
- Ovulation induction
- IVF
- GIFT
- ICSI (intracytoplasmic sperm injection)
- Other (specify)

Tick ‘Yes’ or ‘No’ to indicate whether this pregnancy was achieved via assisted conception. If ‘Yes’, tick the box(es) that correspond to the method(s) used to successfully assist conception for this pregnancy.

**Definitions:**

- **AIH/AID**
  Artificial insemination using either the husband or male partner’s sperm or donor sperm.

- **Ovulation induction**
  Ovulation is induced by pharmacological therapy such as Clomid.

- **IVF**
  In Vitro Fertilisation: Co-incubation of sperm and oocyte outside the body of the woman.

- **GIFT**
  Gamete Intra Fallopian Transfer: A medical procedure of transferring an egg(s) and sperm to the body of the woman.

- **ICSI**
  Intracytoplasmic Sperm Injection: Involves the injection of a single sperm directly into the ovum, combined with IVF.

- **Other**
  Indicate the type of method used, eg Assisted hatching, Blastocyst culture.
7 LABOUR AND DELIVERY

7.1 INTENDED PLACE OF BIRTH AT ONSET OF LABOUR

<table>
<thead>
<tr>
<th>INTENDED PLACE OF BIRTH AT ONSET OF LABOUR</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hospital</td>
</tr>
<tr>
<td>Birthing Centre</td>
</tr>
<tr>
<td>Home</td>
</tr>
<tr>
<td>Other</td>
</tr>
</tbody>
</table>

Tick the box (one box only) that corresponds to the intended place of birth at onset of labour. If intended place of birth was other than those listed, tick ‘Other’ and specify in the space provided.

Definitions:

- **Hospital**
  A health care facility established under Commonwealth, State or Territory legislation as a hospital or a free-standing day procedure unit and authorised to provide treatment and/or care to patients.

- **Birthing centre**
  A facility where women are able to birth in an environment which:
  (a) is free-standing or physically separate from a labour ward but has access to emergency medical facilities for both mother and child if required; and
  (b) has home-like atmosphere; and
  (c) focuses on a model of care (eg Midwifery model) which ensures continuity of care/caregiver; a family-centred approach; and informed client participation in choices related to the management of care.

- **Home**
  Home may be the mother’s own home or where the baby is born in a home environment where “home” may actually be that of a midwifery practitioner or any other person and attended by a midwifery practitioner.

Mothers who plan to give birth in birthing centres or at home usually have different risk factors compared to those who plan to give birth in hospital.

7.2 ACTUAL PLACE OF BIRTH OF BABY

<table>
<thead>
<tr>
<th>ACTUAL PLACE OF BIRTH OF BABY</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hospital</td>
</tr>
<tr>
<td>Birthing Centre</td>
</tr>
<tr>
<td>Home</td>
</tr>
<tr>
<td>Other (BBA)</td>
</tr>
</tbody>
</table>
Tick the box (one box only) that corresponds to the actual place where the birth of the baby occurred (see Section 7.1 for definitions). If the actual place of birth of the baby was other than those listed, tick ‘Other’ and specify in the space provided, eg hospital car park, on the way to hospital in an ambulance, etc.

Note that if the mother at the onset of labour intended to have her baby in a hospital but actually delivered at home, this should be reported as ‘Other (BBA)’ in this field.

This field is used in conjunction with the ‘Intended Place of Birth at Onset of Labour’ field. It identifies mothers who intend to deliver at hospital but deliver at home, compared to those mothers who intend to deliver at home and do so.

This information is used to analyse the risk factors and outcomes by place of birth. While most deliveries occur within hospitals an increasing number of births now occur in other settings. It is important to monitor the births occurring outside hospitals and to ascertain whether or not the actual place of birth was planned.

### 7.3 ONSET OF LABOUR

<table>
<thead>
<tr>
<th>ONSET OF LABOUR</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tick one box only</td>
</tr>
<tr>
<td>Spontaneous</td>
</tr>
<tr>
<td>Induced</td>
</tr>
<tr>
<td>No labour</td>
</tr>
<tr>
<td>(caesarean section)</td>
</tr>
</tbody>
</table>

Tick the box (one box only) that corresponds to how labour commenced. ‘No labour’ can only be associated with a caesarean section.

**Definitions:**

- **Spontaneous**  
  Labour commences at the onset of regular uterine contractions, which act to produce progressive cervical dilatation, and is distinct from spurious labour or spontaneous pre-labour rupture of membranes.

- **Induced**  
  Medical and/or surgical procedure performed for the purpose of stimulating and establishing labour in a woman who has not commenced labour spontaneously.

- **No labour (caesarean section)**  
  Indicates the absence of labour, as in a caesarean section performed before the onset of labour or a failed induction.

Note that when a failed induction of labour results in a caesarean, ‘No labour (caesarean section)’ should be ticked, and the reason for caesarean should be reported as failed induction of labour.

The onset of labour is closely associated with type of delivery and maternal and neonatal morbidity. Induction rates vary for maternal risk factors and obstetric complications and are indicators of obstetric intervention.
7.4 METHODS USED TO INDUCE LABOUR OR AUGMENT LABOUR?

Methods used to *induce labour* or *augment labour*?
(You may tick more than one box)

- Artificial rupture of membranes (ARM)
- Oxytocin
- Prostaglandins
- Other (specify)

If the labour was induced or spontaneous in onset but subsequently augmented, tick the box(es) that correspond to the method used. If a method was used other than those listed, tick ‘Other’ and specify in the space provided, eg Foley’s catheter.

7.5 REASON FOR INDUCTION

If labour induced
Reason for induction

If labour was induced, specify the reason for induction in the space provided, eg rupture of membranes > 24 hours before delivery, post-term, etc. If the reason for induction was a social reason, specify the actual reason(s) rather than writing ‘social reasons’.

Note that ‘failure to progress’, or any other conditions that pertain to labour, are not valid reasons for induction as labour has not yet commenced. Also note that ‘augmentation’ is not a valid reason for induction as augmentation is any medical or surgical intervention that assists with the continuation of a labour that has had a spontaneous or induced onset, eg ARM, administration of oxytocin.

Where a failed induction of labour has occurred, ensure that ‘No labour (caesarean section)’ has been ticked. The reason the induction was attempted should be reported in the appropriate field (eg medical conditions or pregnancy complications).

7.6 MEMBRANES RUPTURED

MEMBRANES RUPTURED

____days  ____hours  ____mins

before delivery

Enter the number of days, hours and minutes before delivery the membranes were ruptured. If membranes ruptured at delivery, then record ‘at delivery’ or enter 0. If a ‘no labour’ caesarean section occurs, it cannot be assumed that the membranes ruptured at delivery so record the actual time or write ‘at delivery’ or enter ‘0’ as above.
7.7 LENGTH OF 1ST AND 2ND STAGE OF LABOUR

LENGTH OF LABOUR

<table>
<thead>
<tr>
<th></th>
<th>1&lt;sup&gt;st&lt;/sup&gt; stage</th>
<th>2&lt;sup&gt;nd&lt;/sup&gt; stage</th>
</tr>
</thead>
<tbody>
<tr>
<td>hours</td>
<td></td>
<td></td>
</tr>
<tr>
<td>minutes</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Enter in the length of each of 1<sup>st</sup> stage and 2<sup>nd</sup> stage of labour in hours and minutes.

Definitions:

- **Stage 1**
  
  Begins with the onset of regular uterine contractions and is complete when the cervix is fully dilated (10cm).

- **Stage 2**
  
  Begins when the cervix is fully dilated (10cm) and is complete with the birth of the baby.

Where the labour is interrupted (eg by caesarean section) and therefore either stage one or two are interrupted, complete as follows:

- If stage one is complete, and stage two interrupted, then report total length of stage one in hours and minutes, and enter ‘not completed’ for stage two.
- If neither stage is complete, then indicate by writing ‘not completed’ in both sections of the field.

Please note that if quantitative measurement has not been performed, then clinical judgement based on subjective observation is appropriate (i.e. vaginal examination to confirm dilation is not mandatory). Use of other clinical observations used to manage labour are appropriate indicators of stages of labour.

Where length of stages is unknown please write ‘unknown’.

7.8 PRESENTATION AT BIRTH

Tick one box only

- Vertex
- Breech
- Face
- Brow
- Transverse/shoulder
- Other (specify)

Tick the box (one box only) that corresponds to the presentation of the fetus at birth. If the presentation at birth is other than those listed, tick ‘Other’ and specify the presentation in the space provided.

If the presentation is unknown, for example, due to extreme prematurity or macerated fetus, document this in the space provided.
Presentation types other than vertex are associated with higher rates of caesarean section, instrumental delivery, perinatal mortality and neonatal morbidity.

7.9 METHOD OF BIRTH

Tick the box (one box only) that corresponds to the method of birth of the baby, i.e. the method of complete expulsion or extraction from its mother of a product of conception. If the method of birth was other than those listed, tick ‘Other’ and specify the method in the space provided.

Note that a vaginal breech with forceps applied to the after coming head should be recorded as ‘Forceps’. Forceps used to assist delivery at caesarean should be reported as a caesarean.
7.10 WATER BIRTH

Definitions:
- **Vaginal non-instrumental**
  A birth which is achieved solely by the mother’s expulsive efforts requiring no mechanical or surgical assistance.

- **Forceps**
  Where forceps are applied to assist the delivery process, including rotation forceps, liftout, etc.

- **Vacuum Extractor**
  An assisted birth using a suction cap applied to the baby’s head, including rotation vacuum, also known as Ventousse Extractor.

- **LSCS**
  Lower segment caesarean section.

- **Classical CS**
  Classical caesarean section.

- **Other**
  Includes birth methods not classified above, eg Hysterotomy or extraction at post mortem.

Tick the box to indicate if this birth was a water birth.

If the birth was a water birth, tick the box to indicate if it was an unplanned or a planned water birth.

For a birth to be considered a water birth, the baby’s head must remain submerged under water until after the body is born.

7.11 REASON FOR FORCEPS/VACUUM

If forceps or vacuum were used as the method of birth, specify the reason in the space provided, eg ‘prolonged active 2nd stage’, ‘Direct OP’.
7.12 REASON FOR CAESAREAN

If caesarean section was performed as the method of birth, specify the reason in the space provided, eg 'repeat caeser', 'fetal distress', 'prolonged labour', etc.

Where a caesarean occurs as a result of a failed forceps/vacuum, then reason for caesarean should be reported as 'failed forceps/vacuum' and the original indication for the trial of forceps/vacuum (eg prolonged active 2nd stage) should be reported as a labour and delivery complication.

7.13 CERVICAL DILATATION PRIOR TO CAESAREAN

Cervical dilatation prior to caesarean
3cm or less
More than 3cm
Not measured

If a caesarean was performed, tick the box (one box only) that corresponds to the level of dilatation of the cervix prior to the caesarean. If the cervical dilatation was not measured, tick ‘Not measured’.

Note this field is mandatory when the method of birth is a caesarean, including no labour caesarean. It is not necessary to complete for any other method of birth.

7.14 ANTIBIOTICS AT TIME OF CAESAREAN

ANTIBIOTICS RECEIVED AT TIME OF CAESAREAN
Tick one box only

None
Prophylactic antibiotics received
Antibiotics already received

When the method of birth is either a lower segment caesarean section or a classical caesarean section, tick the box (one box only) that corresponds to the administration of antibiotics to the mother in relation to the caesarean.

If antibiotics were not received at the time of LSCS or classical caesarean section, tick the ‘None’ box.

If antibiotics have been received for prophylaxis of infection specifically associated with the caesarean, tick the ‘Prophylactic antibiotics received’ box.

If antibiotics have been received for a known condition (eg, chorioamnionitis, pneumonia, etc) at the time of LSCS or classical caesarean, tick the ‘Antibiotics already received’ box. This does not include antibiotic prophylaxis.
This information is used to assist the identification of adverse outcomes in relation to maternal health and wellbeing.

### 7.15 PLACENTA/CORD*

<table>
<thead>
<tr>
<th>PLACENTA / CORD</th>
</tr>
</thead>
</table>

Indicate whether the placenta was complete or other and/or whether the cord had 3 vessels or other at delivery in the space provided. Report any malformations noted, e.g. circumvallate placenta, velamentous cord insertion, true knot in cord.

### 7.16 PRINCIPAL ACCOUCHEUR

<table>
<thead>
<tr>
<th>PRINCIPAL ACCOUCHEUR</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tick one box only</td>
</tr>
<tr>
<td>Obstetrician</td>
</tr>
<tr>
<td>Other medical officer</td>
</tr>
<tr>
<td>Midwife</td>
</tr>
<tr>
<td>Midwifery Student</td>
</tr>
<tr>
<td>Medical student</td>
</tr>
<tr>
<td>Other (specify)</td>
</tr>
</tbody>
</table>

Tick the box (one box only) that corresponds to the principal person who assisted the mother in the birth of the baby. If the principal accoucheur is other than those listed, tick “Other” and specify the accoucheur in the space provided.

**Definitions:**
- **Obstetrician**
  A medical doctor who is qualified in the field of obstetrics.

- **Other medical officer**
  Includes registrar, junior house officer, resident, general practitioner, etc.

- **Midwife**
  A registered nurse who is qualified in the field of midwifery.

- **Midwifery Student**
  A registered nurse training to obtain qualifications in the field of midwifery.

- **Medical student**
  A student training to obtain qualifications to become a medical doctor.

- **Other**
  Includes a registered nurse without midwifery qualifications, doulas, ambulance officer, self, husband, other patient, etc.
7.17 PERINEUM

Please tick the most severe

<table>
<thead>
<tr>
<th>Intact</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Grazes</td>
<td></td>
</tr>
<tr>
<td>Lacerated</td>
<td></td>
</tr>
<tr>
<td>- 1st degree</td>
<td></td>
</tr>
<tr>
<td>- 2nd degree</td>
<td></td>
</tr>
<tr>
<td>- 3rd degree</td>
<td></td>
</tr>
<tr>
<td>- 4th degree</td>
<td></td>
</tr>
</tbody>
</table>

Episiotomy? No [ ] Yes [ ]

Tick the box that corresponds to the condition of perineum following delivery. Tick ‘Yes’ or ‘No’ to indicate whether or not an episiotomy was performed.

Note that if an episiotomy has been performed, the perineum can not be intact and this box should be left blank along with the laceration boxes.

If both a 2nd degree tear and an episiotomy occurred, please note which occurred first.

If an episiotomy is extended to a 3rd or 4th degree tear, tick both corresponding boxes (i.e. episiotomy as well as either 3rd or 4th degree tear).

Definitions:

- **Intact**
  The perineum is intact following delivery.

- **Graze**
  A slight abrasion of the skin following delivery.

- **Lacerated**
  If the perineum is lacerated following delivery, indicate the degree of laceration.
  - **1st Degree**
    Tear or laceration involving one of the fourchette, hymen, labia, skin, vagina or vulva.
  - **2nd Degree**
    Tear or laceration involving the pelvic floor or perineal muscles or vaginal muscles.
  - **3rd Degree**
    Tear or laceration involving the anal sphincter or recto vaginal septum.
  - **4th Degree**
    Third degree tear or laceration also involving the anal mucosa or rectal mucosa.

- **Episiotomy**
  Surgical incision into the perineum and vagina to assist delivery.
Perineal laceration (tear) may cause significant maternal morbidity in the postnatal period. Episiotomy is an indicator of management during labour and, to some extent intervention rates.

7.18 OTHER GENITAL TRAUMA

Specify any other genital trauma experienced by the mother in the space provided, including high vaginal tears where the perineum is intact, cervical tears, urethral tears, etc.

7.19 SURGICAL REPAIR OF THE VAGINA OR PERINEUM

Tick ‘Yes’ or ‘No’ to indicate whether the vagina or perineum was surgically repaired. Note that if an episiotomy has been performed, then corresponding surgical repair would be expected.

7.20 NON-PHARMACOLOGICAL ANALGESIA DURING LABOUR/DELIVERY

Tick the box(es) under the Non-Pharmacological Analgesia during Labour/Delivery heading that correspond to the non-pharmacological analgesia administered to the mother during labour and delivery. If non-pharmacological analgesia used was other than those listed, tick ‘Other’ and specify the non-pharmacological analgesia in the space provided. If no non-pharmacological analgesia was administered, tick ‘None’.
### Definitions:
- **Heat Pack**: Includes the use of electronic heat pads, heat wheat packs and gel packs.
- **Water Immersion**: The labouring woman places her body into water or other liquid so that it is completely covered by the liquid.
- **TENS**: an electronic device that delivers small electrical impulses to the body via electrodes placed on the skin.
- **Other**: Includes the use of medication, visualisation and hypnotherapy.

---

#### 7.21 PHARMACOLOGICAL ANALGESIA DURING LABOUR/DELIVERY

| PHARMACOLOGICAL ANALGESIA DURING LABOUR/DELIVERY |  
|---|---  
| None |  
| Nitrous oxide |  
| Systemic opioid (incl. narcotic (IV/IM)) |  
| Epidural |  
| Spinal |  
| Combined Spinal-Epidural |  
| Caudal |  
| Other (specify) |  

Tick the box(es) under the Pharmacological Analgesia heading that correspond to the pharmacological analgesia administered to the mother during labour and delivery. If a pharmacological analgesia other than those listed was used, tick ‘Other’ and specify the pharmacological analgesia in the space provided. If no pharmacological analgesia was administered, tick ‘None’.

Definitions are listed on the next page.
**Definitions:**

- **Analgesia**
  Agents administered to the mother by injection or inhalation to relieve pain during labour and delivery.

- **Nitrous Oxide**
  Gas providing light anaesthesia delivered in various concentrations with oxygen.

- **Systemic Opioid (incl. narcotic (IM/IV))**
  Opioid analgesics that acts on the patient’s central nervous system. This includes drugs which have an agonist action at the opioid receptor on the cell.

- **Epidural**
  Injection of a local anaesthetic into the epidural space of the spinal column.

- **Spinal**
  Injection of an analgesic drug or anaesthetic drug into the subarachnoid space of the spinal cord, also called the Subarachnoid Block Anaesthesia.

- **Combined Spinal-Epidural**
  Needle-through-needle injection of an analgesic drug or anaesthetic drug into both the epidural space and the subarachnoid space of the spinal column.

- **Caudal**
  Injection of a local anaesthetic agent into the caudal portion of the spinal canal through the sacrum.

### 7.22 LABOUR AND DELIVERY COMPLICATIONS

<table>
<thead>
<tr>
<th>LABOUR AND DELIVERY COMPLICATIONS</th>
</tr>
</thead>
<tbody>
<tr>
<td>You may tick more than one box</td>
</tr>
<tr>
<td>None</td>
</tr>
<tr>
<td>Meconium liquor</td>
</tr>
<tr>
<td>Fetal distress</td>
</tr>
<tr>
<td>Cord prolapse</td>
</tr>
<tr>
<td>Cord entanglement with compression</td>
</tr>
<tr>
<td>Failure to progress</td>
</tr>
<tr>
<td>Prolonged second stage (active)</td>
</tr>
<tr>
<td>Precipitate labour/delivery</td>
</tr>
<tr>
<td>Retained placenta with manual removal</td>
</tr>
<tr>
<td>with haemorrhage</td>
</tr>
<tr>
<td>without haemorrhage</td>
</tr>
<tr>
<td>Primary PPH (500-999ml)</td>
</tr>
<tr>
<td>Primary PPH (=&gt;1000ml)</td>
</tr>
<tr>
<td>Other (specify)</td>
</tr>
</tbody>
</table>
Tick the box(es) that correspond to any complications that arose during labour and delivery. If complications arose other than those listed, tick ‘Other’ and specify the complication(s) in the space provided (see Appendix D for examples). If no complications were experienced, tick ‘None’.

**Definition:**
- **Labour and delivery complications**
  Medical and obstetric complications (necessitating intervention) arising after the onset of labour and before the completed delivery of the baby and placenta.

Complications of labour and delivery may cause maternal morbidity and may affect the health status of the baby at birth.

### 7.23 ANAESTHESIA FOR DELIVERY

<table>
<thead>
<tr>
<th>ANAESTHESIA FOR DELIVERY</th>
</tr>
</thead>
<tbody>
<tr>
<td>None</td>
</tr>
<tr>
<td>Epidural</td>
</tr>
<tr>
<td>Spinal</td>
</tr>
<tr>
<td>Combined Spinal-Epidural</td>
</tr>
<tr>
<td>General anaesthetic</td>
</tr>
<tr>
<td>Local to perineum</td>
</tr>
<tr>
<td>Pudendal</td>
</tr>
<tr>
<td>Caudal</td>
</tr>
<tr>
<td>Other (specify)</td>
</tr>
</tbody>
</table>

Tick the box(es) under the Anaesthesia heading that correspond to the anaesthesia administered to the mother for delivery. If the anaesthesia used was other than those listed, tick ‘Other’ and specify the anaesthesia used in the space provided. If no anaesthesia was administered, tick ‘None’.

Please note that a response is required in non-pharmacological analgesia, pharmacological analgesia and anaesthesia fields, eg if delivery is by elective caesarean section, and no non-pharmacological or pharmacological analgesia are used, then ‘None’ should be ticked in both fields.

Note also that local to the perineum for the sole purpose of repair of tear or episiotomy is not considered anaesthetic for delivery, and therefore should not be included.

Definitions are listed on the next page.
Definitions:

- **Anaesthesia**
  Agents administered to the mother for the operative/instrumental delivery of the baby (caesarean section, forceps or vacuum delivery).

- **Epidural**
  Injection of a local anaesthetic into the epidural space of the spinal column.

- **Spinal**
  Injection of an analgesic drug or anaesthetic drug into the subarachnoid space of the spinal cord. Also called the Subarachnoid Block Anaesthesia.

- **Combined Spinal-Epidural**
  Needle-through-needle injection of an analgesic drug or anaesthetic drug into both the epidural space and the subarachnoid space of the spinal column.

- **General Anaesthetic**
  Various anaesthetic agents given primarily by inhalation or intravenous injection.

- **Local to Perineum**
  Infiltrating the perineum with local anaesthetic.

- **Pudendal**
  Injection of local anaesthetic to the pudendal nerves.

- **Caudal**
  Injection of a local anaesthetic agent into the caudal portion of the spinal canal through the sacrum.

### 7.24 CTG IN LABOUR

| CTG in labour? | No □ | Yes □ |

Tick ‘Yes’ or ‘No’ to indicate whether Cardiotocography (CTG) monitoring was performed during labour. Any external trace (including ‘routine baseline’ traces) recorded during labour, regardless of the duration of recording (i.e. continuous or intermittent) should be reported. A baseline trace recorded prior to labour commencing should not be included.

### 7.25 FSE IN LABOUR

| FSE in labour? | No □ | Yes □ |

Tick ‘Yes’ or ‘No’ to indicate whether Fetal Scalp Electrode (FSE) monitoring was performed during labour.
7.26  **FETAL SCALP pH AND RESULT**

<table>
<thead>
<tr>
<th></th>
<th>No</th>
<th>Yes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fetal Scalp pH?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fetal Scalp pH result</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Indicate whether fetal scalp pH was measured or not.

If the fetal scalp pH was taken then record the fetal scalp pH result.

7.27  **LACTATE AND RESULT**

<table>
<thead>
<tr>
<th></th>
<th>No</th>
<th>Yes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lactate?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lactate result</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Tick ‘Yes’ or ‘No’ to indicate whether fetal scalp lactate was measured.

If the fetal scalp lactate was taken, record the fetal scalp lactate result.
8 BABY

Identification labels may be attached to the back of the original and duplicate copy of the MR63D form. If an identification label is used only on the hospital copies DO NOT FORGET to complete BABY’S UR NUMBER and DATE OF BIRTH on the Data Collections Unit copy. If a label is used on the duplicate copies, then identifying information that is not required by Data Collections Unit can be crossed through using a felt tipped pen (as ball point will affect the clarity of information on the form due to the carbonisation of the paper).

Note: In the case of multiple births, a separate MR63D must be completed for each baby. If the forms are pinned together prior to dispatch, the common information need not be repeated. Details in the LABOUR AND DELIVERY, BABY, POSTNATAL and BABY DISCHARGE DETAILS must be completed for each baby.

8.1 BABY’S UR NUMBER

Enter the Unit Record (UR) number assigned to the baby (if applicable), eg:

| BABY’S UR No. | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |

For home births where the baby is not admitted to a hospital, this field is not required, however if the private midwifery practitioner assigns a record number for administrative purposes it can be included.

8.2 INDIGENOUS STATUS – BABY

Tick the box (one box only) that corresponds to the Indigenous Status of the baby.

Definitions:
An Aboriginal or Torres Strait Islander is a person of Aboriginal or Torres Strait Islander descent who identifies as an Aboriginal or Torres Strait Islander and is accepted as such by the community in which she lives.

- **Aboriginal**
  Aboriginal but not Torres Strait Islander origin.

- **Torres Strait Islander**
  Torres Strait Islander but not Aboriginal origin.

- **Aboriginal and Torres Strait Islander**
  Both Aboriginal and Torres Strait Islander origin.

- **Neither Aboriginal nor Torres Strait Islander**
  Neither Aboriginal nor Torres Strait Islander origin.
Note that a baby’s indigenous status cannot be determined simply by observation and therefore this question must be asked of all mothers. For further information regarding determining Indigenous status, please refer to the ‘Are you of Aboriginal or Torres Strait Islander origin?’ pamphlet. If you require copies of this publication, please contact the National Centre for Aboriginal and Torres Strait Islander Statistics (Australian Bureau of Statistics) on the free call number 1800 633 216.

8.3 DATE OF BIRTH

Enter the day, month and year of the baby’s date of birth using all boxes, eg 1 July 2010 should be entered as:

DOB 0;1;0;7;2;0;1;0

8.4 TIME OF BIRTH

Enter the time of birth of the baby using the 24 hour clock, eg 2.30pm should be entered as 14:30 hours. If the time of birth of the baby is midnight, this should be recorded as 00:00 hours to indicate the start of the day.

TIME OF BIRTH [ ] [ ] [ ] hours

8.5 BIRTHWEIGHT

Enter the first weight of the fetus or baby obtained after birth in grams, eg 3500 grams.

BIRTHWEIGHT [ ] [ ] [ ] grams

8.6 GESTATION

Enter the estimated gestational age of the baby in completed weeks and days, as determined by clinical assessment after birth. Do not use ‘T’ for term, or ‘K’.

Gestational age is a key outcome of pregnancy and an important risk factor for neonatal outcomes.

8.7 HEAD CIRCUMFERENCE AT BIRTH

Enter the head circumference of the baby at birth in centimetres, to the nearest one decimal place.
8.8 LENGTH AT BIRTH

LENGTH AT BIRTH

Enter the length of the baby at birth in centimetres, to the nearest one decimal place.
8.9 PLURALITY

<table>
<thead>
<tr>
<th>PLURALITY</th>
</tr>
</thead>
<tbody>
<tr>
<td>Single</td>
</tr>
<tr>
<td>Twin I</td>
</tr>
<tr>
<td>Twin II</td>
</tr>
<tr>
<td>Other (Specify)</td>
</tr>
</tbody>
</table>

Tick one box only to indicate whether this pregnancy has resulted in a ‘Single’ birth, or for a multiple birth, tick the box for which baby the form is being completed. For example, if the form relates to the second twin, tick ‘Twin II’.

For the first baby of triplets or higher, tick ‘Other’ and write, for example, ‘Triplet I’ in the space provided.

Note: The plurality refers to the total number of births resulting from this pregnancy. If the pregnancy commences as a twin pregnancy but one foetus is miscarried/aborted before 20 weeks and/or 400 grams, the plurality would be single.

8.10 SEX

<table>
<thead>
<tr>
<th>SEX</th>
</tr>
</thead>
<tbody>
<tr>
<td>Male</td>
</tr>
<tr>
<td>Female</td>
</tr>
<tr>
<td>Indeterm.</td>
</tr>
</tbody>
</table>

Tick the box (one box only) that corresponds to the sex of the baby. If the sex of the baby cannot be determined, tick ‘Indeterm’.

8.11 BIRTH STATUS

<table>
<thead>
<tr>
<th>BIRTH STATUS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Born alive</td>
</tr>
<tr>
<td>Stillborn - macerated</td>
</tr>
<tr>
<td>No     Yes</td>
</tr>
</tbody>
</table>

Tick the box that corresponds to the result of the birth. If the baby was born alive, tick ‘Born alive’. If the baby was not born alive, tick ‘Stillborn’.

If the baby was stillborn, indicate whether the baby was macerated by ticking ‘Yes’ or ‘No’.

Note that maceration status should only be completed in the case of stillbirths, and should not be used to indicate ‘peeling skin’ associated with a post term infant.

Definitions are listed on next page.
**Definitions:**

- **Live birth**
  The complete expulsion or extraction from its mother of a product of conception, irrespective of the duration of pregnancy, which, after such separation, breathes or shows any other evidence of life, such as beating of the heart, pulsation of the umbilical cord or definite movement of voluntary muscles, whether or not the umbilical cord has been cut or the placenta is attached.

- **Stillbirth**
  A fetal death prior to the complete expulsion or extraction from its mother of a product of conception of 20 or more completed weeks of gestation or of 400 grams or more birthweight; the death is indicated by the fact that after such separation the foetus does not breathe or show any other evidence of life, such as beating of the heart, pulsation of the umbilical cord, or definite movement of voluntary muscles.

- **Macerated**
  Softening and breaking down of skin caused by prolonged exposure to amniotic fluid in a deceased foetus.

### 8.12 APGAR SCORE

<table>
<thead>
<tr>
<th>APGAR SCORE</th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Heart rate</td>
<td>1 min</td>
<td>5 min</td>
</tr>
<tr>
<td>Respiratory effort</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Muscle tone</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Reflex irritability</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Colour</td>
<td></td>
<td></td>
</tr>
<tr>
<td>TOTAL</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Enter the 1 minute and 5 minute Apgar scores in the boxes for each of the conditions listed (refer to table below).

<table>
<thead>
<tr>
<th>Sign</th>
<th>Scores 0</th>
<th>Scores 1</th>
<th>Scores 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Heart rate</td>
<td>Absent</td>
<td>&lt;100 beats/min</td>
<td>&gt;100 beats/min</td>
</tr>
<tr>
<td>Respiratory rate</td>
<td>Absent</td>
<td>Slow, irregular</td>
<td>Good lusty cry</td>
</tr>
<tr>
<td>Muscle Tone</td>
<td>Flaccid, limp</td>
<td>Flexion of extremities</td>
<td>Active flexion</td>
</tr>
<tr>
<td>Reflex Irritability</td>
<td>No response</td>
<td>Grimace, some motion</td>
<td>Cry, cough</td>
</tr>
<tr>
<td>Colour</td>
<td>Cyanotic, pale</td>
<td>Pink body, acrocyanosis</td>
<td>Pink body/extremities</td>
</tr>
</tbody>
</table>


Enter the total Apgar scores in the boxes provided.

The Apgar score is a numerical score to evaluate the baby’s condition at 1 minute and 5 minutes after birth. It is an indicator of the health of the baby, particularly after complications of pregnancy and/or labour and birth. It is useful in deciding the need for and adequacy of resuscitation.
8.13 REGULAR RESPIRATION

Enter, to the nearest minute, the time the baby took to establish regular, spontaneous breathing. If respirations were established 30 to 59 seconds after birth, record as 1 minute.

If the baby established respirations spontaneously tick the 'at birth box'; if the baby was ventilated, tick the 'intubated/ventilated' box; if respirations were never established, tick the 'respirations not established' box.

8.14 RESUSCITATION

Tick the box(es) that correspond to the method of resuscitation used. If resuscitation methods were used other than those listed, tick 'Other' and specify the method(s) used in the space provided, eg tactile stimulation. Include other drugs used for resuscitation, eg adrenalin, etc. If no methods were used, tick 'None'.

Definitions are listed on the next page.
This information is required to analyse the need for resuscitation after complications of labour and delivery and to evaluate level of services required for different birth settings.

8.15 CORD Ph AND VALUE

<table>
<thead>
<tr>
<th>Cord pH?</th>
<th>Yes</th>
<th>No</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cord pH value</td>
<td></td>
<td></td>
</tr>
<tr>
<td>BE</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Tick ‘Yes’ or ‘No’ to indicate whether pH of the umbilical cord was measured.

If the Cord pH was measured provide the cord pH value.

Record the Base Excess (BE) level if measured.

Note: this is not a mandatory field on the form and subsequently no information is stored by the PDC from this field.
8.16 VITAMIN K (FIRST DOSE)

Tick the box (one box only) that corresponds to the method of administration for first dose of Vitamin K was administered. If no Vitamin K was administered, tick 'None'.

8.17 HEPATITIS B VACCINATION (BIRTH DOSE)

Tick the box (one box only) that corresponds to whether or not the birth dose Hepatitis B vaccination was given. Note that this is not exclusive to doses given immediately after birth or whilst still within the delivery room, and therefore includes doses given prior to discharge. This field does not refer to administration of Hepatitis B immunoglobulin.

8.18 HEPATITIS B IMMUNOGLOBULIN

Tick the box (one box only) that corresponds to whether or not Hepatitis B immunoglobulin was given. Note that this is not exclusive to dose given immediately after birth or whilst still within the delivery room, and therefore includes any dose given prior to discharge. This field does not refer to administration of Hepatitis B Vaccination.
9 POSTNATAL DETAILS

9.1 NEONATAL MORBIDITY

<table>
<thead>
<tr>
<th>BABY</th>
<th>NEONATAL MORBIDITY</th>
<th></th>
<th>Diagnosis</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>None</td>
<td>Jaundice</td>
<td></td>
<td>Diagnosis</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Respiratory distress</td>
<td></td>
<td>Diagnosis</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Hypo/Hyperglycaemia or Normal</td>
<td></td>
<td>Results</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Neonatal abstinence syndrome</td>
<td></td>
<td>Drug name</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Infection</td>
<td></td>
<td>Diagnosis</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Other (specify)</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Tick the box(es) that correspond to the conditions/diseases/illnesses/birth traumas experienced by the baby up to the time of discharge or when the baby reaches 28 days of age. Document the diagnosis in the space provided. If a condition is present other than those listed, tick ‘Other’ and specify the condition(s) in the space provided. If there is no neonatal morbidity, tick ‘None’ (See Appendix D for examples of neonatal morbidity).

Examples of diagnoses include:

- **Jaundice**
  Physiological, ABO incompatibility, etc.
  (Indicate whether phototherapy was used to treat the jaundice.)

- **Respiratory distress**
  Transient tachypnoea of the newborn, respiratory distress syndrome, etc.

- **Hypo/Hyperglycaemia or Normal**
  When blood glucose monitoring has been reported, please supply the outcome of the observation (hypoglycaemia, hyperglycaemia or normal).

- **Neonatal Abstinence Syndrome**
  Please specify the name of the drug used by mother.

- **Infection**
  Cytomegalovirus, septicaemia, eye infection, etc and also specify the name of the bacteria where applicable.
9.2 NEONATAL TREATMENT

<table>
<thead>
<tr>
<th>NEONATAL TREATMENT</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>None</td>
<td></td>
</tr>
<tr>
<td>Oxygen for &gt;4 hours</td>
<td></td>
</tr>
<tr>
<td>Phototherapy</td>
<td></td>
</tr>
<tr>
<td>IV/IM antibiotics</td>
<td></td>
</tr>
<tr>
<td>IV fluid</td>
<td></td>
</tr>
<tr>
<td>Mechanical ventilation</td>
<td></td>
</tr>
<tr>
<td>Blood glucose monitoring</td>
<td></td>
</tr>
<tr>
<td>CPAP</td>
<td></td>
</tr>
<tr>
<td>Oro / naso gastric feeding</td>
<td></td>
</tr>
<tr>
<td>Other treatment</td>
<td></td>
</tr>
</tbody>
</table>

Tick the box(es) that correspond to any neonatal treatments given up to the time of discharge or when the baby reached 28 days of age. If a treatment is used other than those listed, tick ‘Other’ and specify the treatment(s) in the space provided. If no treatments were used, tick ‘None’.

Note that if a treatment has been specified, ensure that a corresponding morbidity has also been specified (e.g., if phototherapy is ticked, jaundice should also be ticked in morbidities). If blood glucose monitoring is indicated, then the reason for the monitoring and the outcome of the monitoring should be specified (see 9.1).

9.3 ADMITTED TO ICN/SCN

Hospital nurseries are approved for neo-natal facilities for the treatment of newly born children, under the Health Insurance Act 1973. Hospitals with facilities which meet the criteria (outlined in the Act) may apply for approval under Section 3(2) of the Act to:

The Director,
Insurance and Hospitals Services Section (MDP86),
Australian Department of Health and Aged Care,
GPO Box 9848,
Canberra, ACT 2601.

Approvals will be renewed every 3 years. (See appendix E for list of facilities with approved Level 2 and 3 nurseries at the time of publication).

Was baby admitted to ICN/SCN?
  No [   ]  Yes [   ]

If yes, how many days was baby admitted to:
  • ICN (days) [   ]
  • SCN (days) [   ]

Tick ‘Yes’ or ‘No’ to indicate if the baby was admitted to Intensive Care Nursery (ICN) or Special Care Nursery (SCN).

Specify the type of nursery the baby was admitted to by entering the number of days the baby was admitted to ICN and/or SCN, including 0 if the baby was not admitted. Reporting in this field is only required for those facilities where approval is current. Note
that admissions to a neonatal service level 1 (mature infant nursery) **should not be reported**.

### Definitions:

- **Neonatal Service Level 1 - Mature Infant Nursery (MIN)**
  Neonatal service level 1 primarily cares for healthy infants of 37 weeks gestation or later, and their mothers, postnatally. Requires a secure area for nursing/supervising infants (See Appendix E for specific criteria).

- **Neonatal Service Level 2 - Special Care Nursery (SCN)**
  Neonatal service level 2 provides services at a higher level than a level 1 neonatal service (neonates of 32 weeks gestation or later) and may be used in a ‘step down’ capacity by level 3 neonatal services. This practice usually aims to stabilise the baby on ventilation, in consultation with the Neonatologist from a level 3 neonatal service, before transfer to a higher level service (preferably within 6 hours), (See appendix E for specific criteria).

- **Neonatal Services Level 3 - Intensive Care Nursery (NICU)**
  Neonatal service level 3 provides the highest level of life support including medium to long term ventilation of neonates. Services provided from these units include infant follow-up programs with paediatrician(s) experienced in the follow-up of very premature neonates and access to allied health professionals including a paediatric dietician and social worker (See appendix E for specific criteria).

**SOURCE:** Queensland Health Clinical Services Capability Framework V2.0 (2005)

### 9.4 MAIN REASON FOR ADMISSION TO ICN/SCN

<table>
<thead>
<tr>
<th>Main reason for admission to ICN/SCN</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
</tr>
</tbody>
</table>

If the baby was admitted to either an ICN or SCN, enter one main reason for admission in the space provided. The reason should be a condition, not a treatment, eg ‘prematurity’ rather than ‘tube feeding’, or ‘respiratory distress’ rather than ‘oxygen therapy or observation’. The treatment should be included in the Neonatal Treatments field (see 9.2).
9.5 **CONGENITAL ANOMALY**

**CONGENITAL ANOMALY**

<table>
<thead>
<tr>
<th>No</th>
<th>Yes</th>
<th>Suspected</th>
</tr>
</thead>
</table>

If yes or suspected enter details below or in the Congenital Anomaly section

Tick ‘Yes’, ‘No’ or ‘Suspected’ to indicate whether a congenital anomaly is present or suspected. Congenital anomalies are abnormalities (including deformities) that were present at birth and detected prior to separation from care (See Appendix D for examples of congenital anomalies).

In the case of a diagnosed or suspected anomaly, enter a brief description in the space provided then ensure that the Additional Congenital Anomaly Data section of the form is completed. The medical practitioner responsible for the baby should complete the Congenital Anomaly section, which can be updated up to 28 days after the birth.

Perinatal Data Collection will be reporting against each congenital anomaly whether or not the congenital anomaly was diagnosed prior to birth.

**Hospitals supplying Perinatal data by MR63D**

When entering the description of each congenital anomaly, enter ‘No’ or ‘Yes’ alongside each congenital anomaly to indicate whether the congenital anomaly was diagnosed prior to birth or not.

**Hospitals supplying Perinatal data electronically**

Supply ‘1’ (no) or ‘2’ (yes) to indicate whether the congenital anomaly was diagnosed prior to birth or not as per the file format.

Validations will be generated if this data item has not been supplied.
10 DISCHARGE DETAILS

10.1 DISCHARGE DETAILS OF THE MOTHER

10.1.1 Puerperium Complications

Tick the box(es) that correspond to the puerperium complications experienced by the mother. If a complication is experienced other than those listed, tick ‘Other’ and specify the complication(s) in the space provided (see Appendix D for examples). If no complications are experienced, tick ‘None’.

This field should reflect conditions, not treatments or procedures. For example, a spinal headache would be reported in this field, but if it required intervention such as a blood patch, the treatment would be reported in the puerperium procedures and operations field.

<table>
<thead>
<tr>
<th>Complication</th>
<th>Box</th>
</tr>
</thead>
<tbody>
<tr>
<td>None</td>
<td></td>
</tr>
<tr>
<td>Haemorrhoids</td>
<td></td>
</tr>
<tr>
<td>Wound Infection</td>
<td></td>
</tr>
<tr>
<td>Anaemia</td>
<td></td>
</tr>
<tr>
<td>Dehiscence/disruption of wound</td>
<td></td>
</tr>
<tr>
<td>Febrile</td>
<td></td>
</tr>
<tr>
<td>UTI</td>
<td></td>
</tr>
<tr>
<td>Spinal headache</td>
<td></td>
</tr>
<tr>
<td>Secondary PPH</td>
<td></td>
</tr>
<tr>
<td>Other (specify)</td>
<td></td>
</tr>
</tbody>
</table>

**Definition:**

- **Puerperium complications**
  Medical and obstetric complications of the mother occurring during the postnatal period up to the time of separation from care.

Complications of the puerperal period may cause maternal morbidity, and occasionally death, and may be an important factor in prolonging the duration of hospitalisation after childbirth.
10.1.2 THROMBOPROPHYLAXIS FOLLOWING CAESAREAN

<table>
<thead>
<tr>
<th>THROMBOPROPHYLAXIS FOLLOWING CAESARIAN</th>
</tr>
</thead>
<tbody>
<tr>
<td>You may tick more than one box</td>
</tr>
<tr>
<td>None</td>
</tr>
<tr>
<td>Pharmacological Thromboprophylaxis</td>
</tr>
<tr>
<td>Intermittent Calf Compression</td>
</tr>
<tr>
<td>TED Stocking</td>
</tr>
<tr>
<td>Other (specify)</td>
</tr>
</tbody>
</table>

When the method of birth is either a lower segment caesarean section or a classical caesarean section, tick the box(es) that correspond to any puerperium thromboprophylaxis administered following caesarean section.

If Thromboprophylaxis following LSCS or classical caesarean section was not administered, tick the ‘None’ box.

If Thromboprophylaxis following LSCS or classical caesarean was via pharmacological thromboprophylaxis methods, tick the ‘Pharmacological Thromboprophylaxis’ box.

If Thromboprophylaxis following LSCS or classical caesarean was via intermittent calf compression, tick the ‘Intermittent Calf Compression’ box.

If Thromboprophylaxis following LSCS or classical caesarean was via TED Stocking, tick the ‘TED Stocking’ box.

If Thromboprophylaxis following LSCS or classical caesarean was via other thromboprophylaxis methods, tick the ‘Other (specify)’ box and record the method(s) used in the space provided.

This information is used to assist the identification of adverse outcomes in relation to maternal health and wellbeing.

10.1.3 PUERPERIUM PROCEDURES AND OPERATIONS

<table>
<thead>
<tr>
<th>Puerperium Procedures and Operations</th>
</tr>
</thead>
<tbody>
<tr>
<td>You may tick more than one box</td>
</tr>
<tr>
<td>None</td>
</tr>
<tr>
<td>Blood Transfusion</td>
</tr>
<tr>
<td>Blood Patch</td>
</tr>
<tr>
<td>D &amp; C</td>
</tr>
<tr>
<td>Other (specify)</td>
</tr>
</tbody>
</table>

Tick the box(es) that correspond to any medical or surgical procedures and/or operations that were performed on the mother during the puerperium. If a procedure and/or operation were performed other than those listed, tick ‘Other’ and specify in the space provided (see Appendix D for examples). If no procedures or operations were performed during the puerperium, tick ‘None’. Where procedures are reported that may be performed via different approaches please provide as many details as possible. For example: ligation of fallopian tubes, which may be via laparotomy or laparoscopy, please report as either ‘open abdominal ligation’ or ‘laparoscopic ligation’.
10.1.4 DISCHARGE DETAILS

<table>
<thead>
<tr>
<th>Discharged</th>
<th>Transferred</th>
<th>Died</th>
<th>Remaining in</th>
<th>Place of transfer</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Tick the box (one box only) that corresponds to whether the mother was discharged, transferred to another facility, remaining in hospital or died during the current admission. If the mother was transferred to another facility, enter full name of the other facility in the space provided. In cases such as Mater Mother’s Hospital indicate whether the transfer was to the public or private facility. For PDC purposes, a patient transferred from unit to unit within the same facility (eg maternity to intensive care) is not considered a transfer or discharge.

Enter the day, month and year the mother was discharged, transferred or died using all boxes. If the mother is remaining in after 28 days tick the remaining in box and provide the discharge date when available.

Note that if the baby had an extended stay in hospital and the mother was registered as a boarder so that she could be near her baby, enter the date she was formally discharged as an admitted patient, i.e. the day she changed from an admitted patient to a boarder.

Do not complete the discharge details field when a planned homebirth occurred unless the baby was transferred to a facility following delivery.

10.1.5 EARLY DISCHARGE PROGRAM

Early Discharge Program
No ☐ Yes ☐

Tick the ‘Yes’ box if the mother was released from hospital to an Early Discharge or other similar program. Note there is currently no standard definition available that constitutes an early discharge program. Please report whatever individual facilities regard as an early discharge program.

10.2 DISCHARGE DETAILS OF THE BABY

10.2.1 NEONATAL SCREENING

Enter the day, month and year when the neonatal screening was performed using all boxes, eg if the neonatal screening was performed on 1 November 2010, enter:

<table>
<thead>
<tr>
<th>BABY</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neonatal</td>
</tr>
<tr>
<td>Screening</td>
</tr>
<tr>
<td>0 : 1 1 : 1 2 : 0 1 : 0</td>
</tr>
</tbody>
</table>

Note that this is not a mandatory field on the form, and subsequently no information is stored by PDC from this field.
For enquiries regarding neonatal screening tests please contact the Neonatal Screening Unit on 3636 7171 or 3636 7051.

10.2.2 DISCHARGE WEIGHT

Enter the weight* of the baby on discharge in grams.

<table>
<thead>
<tr>
<th>Discharge weight</th>
<th>grams</th>
</tr>
</thead>
</table>

Note that this is not a mandatory field on the form, and subsequently no information is stored by PDC from this field.

10.2.3 DISCHARGE DETAILS

Tick the box (one box only) that corresponds to whether the baby was discharged, transferred to another facility, remaining in hospital or died during the admission. If the baby was transferred to another facility, enter the full name of the other facility in the space provided. In cases such as Mater Mother’s Hospital indicate whether the transfer was to the public or private facility. For PDC purposes, a baby transferred from unit to unit within the same facility (e.g., Level 3 Nursery to Level 2 Nursery) is not considered a transfer or discharge.

Enter the day, month and year the baby was discharged, transferred or died using all boxes. If the baby is remaining in after 28 days tick the ‘Remaining in’ box and provide the discharge date when available.

Do not complete the discharge details field when a planned homebirth occurred unless the baby was transferred to a facility following delivery.

10.2.4 FLUID BABY RECEIVED AT ANY TIME FROM BIRTH TO DISCHARGE

<table>
<thead>
<tr>
<th>TYPES OF FLUID BABY RECEIVED AT ANY TIME FROM BIRTH TO DISCHARGE:</th>
</tr>
</thead>
<tbody>
<tr>
<td>You may tick more than one box</td>
</tr>
<tr>
<td>Breast milk/colostrum</td>
</tr>
<tr>
<td>Infant formula</td>
</tr>
<tr>
<td>Water, fruit juice or</td>
</tr>
<tr>
<td>water-based products</td>
</tr>
<tr>
<td>Nil by mouth</td>
</tr>
</tbody>
</table>

Tick the box that applies to the type of fluid the baby received at any time from birth to discharge. More than one box may be ticked. This field may be used as an indicator for the Baby Friendly Health Initiative.
10.2.5 FLUID BABY RECEIVED IN THE 24 HOURS PRIOR TO DISCHARGE

**TYPES OF FLUID BABY RECEIVED IN THE 24 HOURS PRIOR TO DISCHARGE:**

You may tick more than one box

- Breast milk/colostrum
- Infant formula
- Water, fruit juice or water-based products
- Nil by mouth

Tick the box that applies to the type of fluid(s) the baby received in the 24 hours prior to discharge (or part thereof). More than one box may be ticked. This field may be used as an indicator for the Baby Friendly Health Initiative.

**NOTE:** If the baby has received a type of fluid in the 24 hours prior to discharge, the type of fluid must also be selected in the types of fluid the baby received at any time from birth to discharge. See section 10.2.5.

**Definitions:**

- **Breast milk/colostrum:**
  Includes breast milk/colostrum received directly from the breast as well as expressed breast milk/colostrum received by but not limited to syringe, cup or enteral tube.

- **Infant formula:**
  Refers to commercially prepared formulas that adequately meet the nutritional needs of the newborn.

- **Water, fruit juice or water-based products:**
  Other types of fluid include but is not limited to water, fruit juice, herbal tea or flavoured water.

10.2.6 ALTERNATE FEEDING METHOD

**ALTERNATE FEEDING METHOD**

You may tick more than one box

- Has the baby been fed by?
  - Bottle
  - Cup
  - Syringe
  - Other (specify)

Tick the box(es) that apply to the type of alternate methods used to feed the baby from birth to discharge (or part thereof). More than one box may be ticked.

This includes babies who are fed expressed breast milk/colostrum via an alternate feeding method. This will enable a broader understanding of bottle usage by reducing association with infant formula and consideration of other liquids such as expressed breast milk. This may be an indicator for the Baby Friendly Health Initiative.
11 ADDITIONAL CONGENITAL ANOMALY DATA

11.1 INDICATE BY SHADING OR MARKING THE APPROPRIATE DIAGRAM(S)

See Appendix B for the diagrams included in Section B of the MR63D form.

In the case of congenital anomaly (ies) with apparent physical defects, indicate by shading or marking the anatomical site(s) affected on the appropriate diagram(s).

11.2 ADDITIONAL CONGENITAL ANOMALY DESCRIPTION OR DETAILS

<table>
<thead>
<tr>
<th>Additional Congenital Anomaly description or details</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
</tr>
</tbody>
</table>

Extra space is provided for a more detailed description of any congenital anomaly which does not fit in the space provided in the postnatal details section of the form.

11.3 MEDICAL PRACTITIONER’S SIGNATURE

Medical Practitioner’s Signature

This form should be signed by the medical practitioner in charge of the neonatal care of the baby.

11.4 SURNAME

Surname (BLOCK LETTERS) ____________________________

Enter the surname of the medical practitioner as it may be necessary to elicit further details at a later date.

11.5 DESIGNATION

Designation ________________________________________

Enter the position/designation of the medical practitioner.

11.6 DATE

Date / /

Enter the date the medical practitioner signed the form.
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