Ectopic Pregnancy - Diagnosis and Management in Gynaecology and Maternal Fetal Medicine (MFM) Services

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1. Purpose of guideline

The purpose of this guideline is to provide guidance to clinicians on diagnosing and managing ectopic pregnancy.

2. Definition, incidence and aetiology

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2.1 Incidence

In most populations, one per cent of pregnancies will be extra-uterine (Department of Health, 2001). The incidence is rising in most countries, and in Auckland there is approximately one ectopic pregnancy (EP) for every 64 live births. This increased prevalence is associated with a high prevalence of chlamydial infection and pelvic inflammatory disease (PID). The use of more sensitive diagnostic techniques means that ectopic pregnancies are more often diagnosed, when previously the implantation site would have remained unknown. The more widespread use of assisted reproductive techniques (ART) such as In vitro fertilisation (IVF) also means an increase in ectopic pregnancies.

2.2 Who is at risk?

Women with a past history of:

- Pelvic infection
- Ectopic pregnancy
- Tubal surgery, pathology or sterilisation
- Sub-fertility
- Current or past Intrauterine contraceptive device (IUCD) use.

3. Clinical features, history and examination

3.1 Clinical features

Clinical features can vary according to the site of the ectopic pregnancy, and the gestational age. Many ectopic pregnancies follow a relatively chronic course, making clinical diagnosis difficult, and sudden collapse associated with tubal rupture is unusual (less than 10% of cases) (Stabile, 1996). Where the finding of a probable ectopic pregnancy is uncertain because of incongruous ultrasound and an intervention is planned, appropriate checking should occur, e.g. by repeat examination by the same clinician, or if immediate treatment is required by another clinician.
3.2 History
- Amenorrhoea: many patients deny any amenorrhoea and some vaginal bleeding around the time a period is common (probably associated with low progestogen levels).
- Unexpected or continuous light vaginal bleeding in early pregnancy may be present.
- Pelvic, abdominal, shoulder tip pain: pain is often initially mild and is usually in one or other adnexa. Exacerbations of pain which then settle for several hours or days is common.
- Past history of pelvic inflammatory disease, IUCD use or conception through assisted reproductive technology.

3.3 Signs on examination
- Minimal to severe abdominal tenderness, guarding or peritonism.
- Cervical excitation
- Adnexal mass or fullness. It is rare to be able to palpate an ectopic pregnancy on vaginal examination.
- Rarely a collapsed patient.

4. Diagnosis

Increased awareness and diagnostic advances have resulted in the earlier and more consistent diagnosis of ectopic pregnancy, contributing to a decrease in the case fatality rate by 90% over the last 20 years (Centers for Disease Control, 1995). It has also permitted a more conservative approach to the management of ectopic pregnancy. The most important advances in diagnosis have been the use of quantitative serum ß-hCG measurement and trans-vaginal ultrasound.

4.1 Serum ß-hCG
- Ninety nine percent sensitivity in detecting pregnancy (Chard, 1992).
- ß-hCG is detectable at the time of a missed menses.
- Abnormal pregnancies are associated with impaired ß-hCG production:
  - In normal early intrauterine pregnancies serum ß-hCG concentrations double every 48 hours (Pittaway, 1987).
  - An increase in serum ß-hCG concentrations of less than 66% in 48 hours is associated with either ectopic or a failing intra-uterine pregnancy in more than 85% of cases (Kadar, DeVore & Romero, 1981).
  - Serum ß-hCG measurement is the single most important test in women of reproductive age group with abdominal pain. Refer to the National Women’s Serial Serum Beta hCG form (Appendix 2).

4.1.1 Clinical application:
- Diagnosis of pregnancy
- Serial measurements can be used to assess the viability of a pregnancy
• To determine the type of sonography to be used.

Caution should be exercised in using serial measurements in very early pregnancy. For viable pregnancies, the ‘doubling-time’ for β-hCG concentrations may be occasionally greater than 48 hours. If methotrexate is being considered then a diagnosis of non-viability must be certain.

4.2 Ultrasoundography (USS)

• Used to confirm or exclude an intrauterine pregnancy. Can be combined with serum β-hCG measurement to produce the concept of a ‘discriminatory zone’, at a certain concentration of β-hCG a viable intra-uterine pregnancy will always be seen on ultrasound. An intra-uterine pregnancy will be seen at a lower β-hCG concentration with trans-vaginal ultrasound than with abdominal ultrasound scan (USS). If a patient presents with symptoms suggestive of an ectopic pregnancy, an empty uterus on trans-vaginal ultrasound and a serum β-hCG above the ‘discriminatory zone’ then an ectopic pregnancy is very likely to be present. This may be unreliable in women at high risk of a multiple pregnancy.
  ○ Discriminatory zones:
    - β-hCG of > 6000 IU/l - intrauterine gestational sac of a viable pregnancy will be seen with trans-abdominal USS (Kadar, DeVore & Romero, 1981).
    - β-hCG of > 2000 IU/l - viable intrauterine gestational sac or a viable pregnancy will be seen with trans-vaginal USS (Stovall & Ling, 1993).
    - Additional USS findings of adnexal mass or significant free fluid in pelvis is suggestive of ectopic pregnancy. If an adnexal gestation sac containing a fetal pole and a visible heartbeat is seen (about 10% of ectopic pregnancies) a diagnosis of ectopic pregnancy is certain. The finding of an adnexal mass adjacent to but separate from the ovary makes a diagnosis of ectopic pregnancy very likely. Sometimes the nature of an adnexal mass is difficult to define making diagnosis less certain. In about 70% of ectopic pregnancies, an adnexal mass will be seen on trans-vaginal ultrasound at first presentation (Atri et. al., 1996).
  ○ Caution should be exercised when using the concept of a discriminatory zone and the ultrasound finding of an empty uterus if serum β-hCG levels are close to the discriminatory zone. This is particularly true if no ectopic pregnancy is seen and medical therapy is contemplated. There may be a risk of administering methotrexate to women with a very early intra-uterine pregnancy. Consultant review prior to treatment and the use of a formally reported scan by a radiologist or sonographer at Women’s Health Ultrasound Department is essential.
  ○ Cervical, caesarean scar and corneal or interstitial ectopic pregnancies can also be diagnosed using established ultrasound criteria outlined below (Elson et. al., 2016).
  ○ Caesarean scar pregnancy:
    - An empty uterine cavity.
    - Gestational sac or solid trophoblastic mass present at level of internal os embedded at the site of previous lower uterine segment caesarean scar.
    - Evidence of prominent trophoblastic or placental circulation on Doppler.
    - Thin or absent myometrium layer between gestational sac and bladder.
    - Empty endocervical canal.
• Cornual or interstitial pregnancy:
  - Empty uterine cavity
  - Products of conception or gestational sac located laterally in the interstitial (intramural) part of the tube and surrounded by <5 mm of myometrium in all planes.
  - Interstitial line sign (presence is 80% sensitive and 98% specific for interstitial pregnancy).
  - 3D USS and Magnetic Resonance Imaging (MRI) can be used to help differentiate between interstitial pregnancy and angular implantation or early intrauterine pregnancy.

• Cervical pregnancy:
  - An empty uterine cavity
  - A barrel-shaped cervix
  - Absence of ‘sliding sign’
  - Gestational sac below level of internal os
  - Blood flow demonstrated on colour Doppler around gestational sac.

• Heterotopic pregnancy:
  - The finding of a viable intrauterine pregnancy is often thought to effectively rule out a diagnosis of ectopic pregnancy. Previously, the incidence of heterotopic pregnancy has been thought to be as low as 1:30 000 (DeVoe & Pratt, 1948). The increasing incidence of ectopic pregnancy and the use of ovulation induction drugs and ART has made heterotopic pregnancy more common and should always be considered in such women.
  - 1:100 in ART programmes.
  - 1:4000 in general population (Hahn, Bachman & McArdle, 1984).

4.3 Laparoscopy
Careful history, physical examination, ß-hCG and ultrasound will in most instances diagnose an ectopic pregnancy. A laparoscopy is usually not indicated as a diagnostic tool. A very early ectopic pregnancy may not be seen at laparoscopy (three per cent of cases) (Stable, 1996).

5. Failure to diagnose an ectopic pregnancy

5.1 Missing a diagnosis of ectopic pregnancy
On occasions, it is not possible to reach a firm diagnosis at initial presentation. The presence of trophoblast of unknown site will have been confirmed by serum ß-hCG measurement and negative ultrasound findings, but further serum ß-hCG measurement and ultrasound evaluation will be needed. This should be done no more than 48-72 hours later.

5.2 Common reasons for failing to diagnose an ectopic pregnancy are:
• A failure to consider EP as a diagnosis
• A failure to correlate scan findings and ß-hCG levels (Department of Health, 2001).

5.3 Implications of missed ectopic
• Loss of opportunity to implement more conservative management
• May require salpingectomy or laparotomy
• Blood transfusion
• Death.

5.4 Pregnancy loss counselling
• Pregnancy loss counselling is to be offered in all cases of expectant management, before administration of methotrexate, and in cases of surgical management when it is appropriate because the woman is asymptomatic.
• This may require the on call social worker to be called.
6. Treatment management guidelines

6.1 Algorithm – choosing a treatment for ectopic pregnancy

Does the patient need emergency surgery?:
- Hemodynamically unstable AND/OR
- Suspicion of impending or ongoing tubal rupture*

Is the hCG¶ >5000 mIU/mL?

Does TVUS show fetal cardiac activity◊?

Refer to Box A

Does the patient have any of the following CONTRAINDICATIONS to methotrexate treatment?
- Heterotopic pregnancy with coexisting viable intrauterine pregnancy
- Breastfeeding
- Hypersensitivity to methotrexate
- Clinically important abnormalities in baseline hematologic, renal, or hepatic laboratory values
- Immunodeficiency, active pulmonary disease, peptic ulcer disease

BOX A
EXPECTANT MANAGEMENT is an option for a small proportion of women who meet ALL of the following criteria:
- hCG <1500 mIU/mL and decreasing (we define a decreasing hCG level as a decrease of >10% across two consecutive measurements)
- TVUS with no fetal heartbeat, no gestational sac, and no extraterine mass suspicious for an ectopic pregnancy
- Able and willing to comply with close follow-up
- Prefers expectant management to methotrexate treatment

Is the patient able and willing to comply with close follow-up AND does she prefer methotrexate treatment to surgery?

Surgical Treatment

Methotrexate Therapy

Reference:
hCG: Human Chorionic gonadotropin
TVUS: Transvaginal ultrasound
* Severe or persistent lower abdominal pain and/or evidence of hematoperitoneum
¶ Serum quantitative beta-hCG

◊ Transvaginal ultrasound

hCG: Human Chorionic gonadotropin
TVUS: Transvaginal ultrasound
6.2 Expectant/conservative management
May be considered if the serum ß-hCG concentration is less than 1500 IU/l (Elson et. al., 2016).

6.3 Medical management
Intra-muscular methotrexate – see Section 8

6.4 Surgical management
- Laparotomy or laparoscopy
- Salpingotomy – opening the tube and removing trophoblast
- Salpingectomy – removing the tube
Note: The words salpingostomy and salpingotomy are used interchangeably.

7. Expectant management of ectopic pregnancy and trophoblast of unknown site

7.1 Criteria for expectant management
- Minimal symptoms
- No significant free fluid in pouch of Douglas
- ß-hCG < 1500 IU/L
- Ectopic pregnancy <30 mm on ultrasound without any fetal cardiac activity seen
- Patient able to consent, understand the need for and be able to attend follow-up.

7.2 Follow-up
Patients should have a repeat serum ß-hCG measurement and trans-vaginal ultrasound at 48-72 hours. As many as 66% of ectopic pregnancies in women with serum ß-hCG concentration less than 1500 IU/L and 90% with ß-hCG <1000 IU/L will resolve spontaneously (Elson et. al., 2016). However, some women will experience pain associated with tubal abortion and there is a risk of tubal rupture even at low ß-hCG levels. If ß-hCG levels are rising, then medical or surgical treatment should be considered.

7.3 Trophoblast of unknown site
In women with serum ß-hCG levels less than the discriminatory zone for trans-vaginal ultrasound (see section 4.2) and negative ultrasound findings a diagnosis of trophoblast of unknown site is made. Many of these women will have failing intra-uterine pregnancies and serum ß-hCG levels will usually start to fall rapidly. Others will have extra-uterine pregnancies that may eventually be detected on a subsequent ultrasound. Follow-up with trans-vaginal ultrasound and serum ß-hCG measurement at 48-72 hours is required. Once ß-hCG levels are falling rapidly, unless new symptoms develop further ultrasonography is no longer necessary.

8. Medical treatment of ectopic pregnancy

8.1 Criteria for medical therapy
- Haemodynamically and clinically stable.
- No significant free fluid in pouch of Douglas.
- Gestation sac <5 cm in adnexa, or meets USS criteria for cervical ectopic, caesarean scar ectopic or cornual/interstitial ectopic pregnancy.
- Normal blood count, liver enzymes, serum creatinine.
- β-hCG < 5000 IU/L, but ideally <1500 IU/L.
- No fetal heartbeat seen on ultrasound.
- Desire for future fertility (methotrexate is still an option for women who do not wish to have further children).
- Patient able to understand and attend for follow-up.
- Patient informed (by medical officer or consultant) regarding possible adverse actions.
- Caesarean scar pregnancy – see (Standard Operating Practice (SOP)) for Ectopic pregnancy – direct administration of methotrexate under ultrasound guidance (Appendix 1).
- Cornual pregnancy – see SOP for Ectopic pregnancy – direct administration of methotrexate under ultrasound guidance if indicated, i.e. BhCG ≥ 5000 IU/L (Appendix 1).

8.2 Drug used
Methotrexate (a single intramuscular (IM) dose initially – repeated doses may be necessary – see protocol).

8.3 Dose
50 mg/m2 of patient body-surface area (typically 75 mg to 95 mg) calculated using nomogram (see Section 9). Exact dose required must be ordered and supplied, NO part doses are permitted, i.e. if 50 mg dose is required and there is not a standard strength available, it is NOT permitted to inject a part dose using a higher strength syringe.

8.4 Side effects
- Stomatitis – common
- Gastritis, nausea – common
- Leucopenia – rare
- Thrombocytopenia – rare
- Raised liver enzymes – rare
- Alopecia – very rare.

8.5 Contra-indications
Methotrexate therapy should not be used in the presence of:
- Hepatic dysfunction: AST > 2x normal
- Abnormal renal function: Serum creatinine level elevated
- Blood dyscrasias: Leucocyte count <3X109/L  
- Platelet count < 100 X 109/L
- Active peptic ulcer disease
- Presence of fetal cardiac activity
- Any active liver or renal disease.
8.6 Protocol

<table>
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<tr>
<th>Test</th>
<th>Day</th>
<th>Therapy</th>
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<tbody>
<tr>
<td>β-hCG, FBC, AST, FBC</td>
<td>4</td>
<td>Methotrexate</td>
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<tr>
<td>β-hCG</td>
<td>7</td>
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Repeat doses of intramuscular methotrexate may be given, if β-hCG levels are rising or plateauing.

All women that have methotrexate need to have an USS the day of methotrexate administration at a Women’s Health ward by a certified sonographer or radiologist. A second opinion from another consultant or discussion with the radiologist reporting on the USS is wise if diagnosis is not certain.

All women must receive counselling (from a pregnancy loss counsellor or nurse certified as qualified) prior to methotrexate administration.

9. Administration of IM methotrexate

9.1 Before administering

- This procedure is a delegated consultant responsibility that is only undertaken by nurses who have been designated to do so following training.
- Patient must be informed of the actions, side effects, contraindications and treatment process.
- Formal pre-decision counselling must be undertaken before administration of methotrexate (counsellors are available through on call - out of hours and weekends and must be called to see patient).
- Ensure patient has the information pamphlet. Before ordering the drug, the patient’s verbal consent is to be obtained by the nurse on the register after detailed explanations. Verbal consent is to be documented.
- Ensure all blood tests are done. These must be checked by the registrar in conjunction with their supervising consultant and they must ensure that the results meet the criteria stated in this policy.
- The clinician administering the methotrexate should know the following:
  - The usual range of drug doses: the recommended dose is 50 mg/m²
  - The method of administration
  - Possible effects - both acute and delayed
  - Nursing management monitoring
  - Patient self-care skills required.
- The clinician administering the medication is to ensure that the correct follow-up has been arranged for the patient.

**Note:** Nurses or clinicians who are pregnant or trying to conceive should not administer methotrexate, as it is teratogenic.
Equipment required
- Latex gloves
- Long-sleeved cuffed gown
- Incontinence sheet
- Pre-filled injection of methotrexate – obtained from Pharmacy Aseptic Production Unit (PAPU)
- Cytotoxic information sheet
- Cytotoxic waste bin.

9.2 Process
The procedure is to be done by a clinician who is on the Limited Cytotoxic Register, or by two registered nurses, one of whom is on the Limited Cytotoxic Register.

Follow these steps to administer IM methotrexate:

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<th>Step</th>
<th>Action</th>
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<tr>
<td>1.</td>
<td>Confirm patient has consented to the procedure and received pre-decision counselling.</td>
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<tr>
<td>2.</td>
<td>Assemble equipment.</td>
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<tr>
<td>3.</td>
<td>Explain procedure to the patient.</td>
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<tr>
<td>4.</td>
<td>Identify patient at the bedside.</td>
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| 5.   | Must check patient identification:  
|      | • Verbally  
|      | • Wristband  
|      | • Cytotoxic drug sheet  
|      | • Label on syringe. |
| 6.   | Ensure patient is comfortable and relaxed. |
| 7.   | Wash hands, don gloves and gown. Place incontinence sheet under the site where the IM injection is to be given. |
| 8.   | If there are any bubbles in the syringe, leave the needle cap on while expelling the air, to prevent aerosoling. |
| 9.   | Administer methotrexate using aseptic IM technique according to recommended best practice. |
| 10.  | Dispose of equipment according to cytotoxic waste guidelines. Wash hands. |
| 11.  | Observe patient for signs of reaction. |
| 12.  | Advise patient regarding follow-up and ensure that she has the patient information leaflet. |
| 13.  | **Note:** Using single IM dose significantly reduces incidence of side effects, but patient must be told of possible side effects. |
9.3 Body surface area of adults

9.3.1 Nomogram

For determination of body surface area from height and weight, a straightedge placed from the patient’s height (left column) to her weight (right column) will give her body surface area (middle column).

Alternatively, the dose may be calculated using the following formula:

\[
(m^2) = \sqrt{\frac{Ht \text{ (cm)} \times Wt \text{ (kg)}}{3600}}
\]

**Note:** Pre-treatment counselling by the pregnancy loss counsellor or on-call social worker should be offered at all times.
A significant fall in ß-hCG may not happen on day four, but at least 15% decline in ß-hCG titre should occur between days four to seven. Otherwise, a second dose of methotrexate (50 mg/m²) should be given one week after the first dose. Patients with declining ß-hCG titres by day seven are monitored weekly until the ß-hCG titre is less than 5 IU/l. A third dose of methotrexate is given only if there is a plateauing or increase in two consecutive weekly ß-hCG titres.

Abdominal pain is common and can be expected three to seven days after the start of treatment. Women should be encouraged to return to the ward for re-assessment. A full clinical assessment including a full blood count and transvaginal scan and then discussion with the consultant on-call is essential.

Patients should refrain from alcohol, sexual intercourse and vitamin pills containing folic acid until serum ß-hCG titres are less than 5 IU/l.

Patients should be advised to use either oral contraceptives or barrier contraception for at least three months after completion of therapy.

10. Direct administration of methotrexate into the gestational sac under ultrasound guidance

This procedure is undertaken by the Maternal Fetal Medicine (MFM) specialists. Contact the MFM Senior Medical Officer on call and complete internal MFM form to make referral for consideration of procedure.

Criteria:
- Confirmed scar, cervical or cornual ectopic accessible for invasive technique
- Clinically stable
- Not suitable for or failed IM methotrexate (Note: Cannot be done within one week of IM methotrexate).
- Preference to avoid surgical management (clinician or patient) and preserve fertility
- Can be done if fetal heart beat present (potassium chloride given prior to methotrexate)
- Nil by mouth for the procedure, with on call and theatre teams aware in case of complication requiring urgent surgical management
- To remain inpatient for at least 24 hours after procedure.

10.1 Before administering
- Patient must be informed of the actions, side effects, contra-indications and treatment process.
- Formal pre-decision counselling must be undertaken before administration of methotrexate. (Counsellors are available through on-call out of hours and weekends and must be called to see patient).
- Ensure patient has the information pamphlet. Before ordering the drug, the patient’s verbal consent is to be obtained after detailed explanations. Verbal consent is to be documented.
• Ensure all blood tests are done. These must be checked by the registrar in conjunction with their supervising consultant and they must ensure that the results meet the criteria stated in this policy.
• The clinician administering the methotrexate should know the following:
  ○ The usual range of drug dose is 50mg or <50 mg/m²
  ○ The method of administration
  ○ Possible effects – both acute and delayed.

10.2 Follow-up care
Follow the same process as for Step 11, 12 and 13 of the section: IM methotrexate protocol.

11. Surgical treatment of ectopic pregnancy

11.1 Laparoscopic management of ectopic pregnancy
Laparoscopy is now increasingly used to treat as well as diagnose ectopic pregnancy. In the past, salpingotomy and salpingectomy were performed. Good evidence now suggests that there is no advantage with salpingotomy with no improvement in fertility. There is a preference for laparoscopic salpingectomy.

11.2 Criteria for laparoscopic salpingectomy
This procedure should be considered in the following groups:
• Haemodynamically stable patients
• Future fertility is not desired
• Fallopian tube is significantly damaged, particularly the fimbrial end
• Fetal heart activity is seen on the scan
• Previous tubal surgery (microsurgery/salpingotomy) on the side of the ectopic pregnancy unless the patient specifically requests tubal preservation
• Ruptured ectopic pregnancy
• Diameter of ectopic pregnancy is greater than 5 cm
• Persistent bleeding from the ectopic site at salpingotomy or if the patient has to be taken back to theatre following salpingotomy
• Patients who are unwilling or unable to return for follow-up.

There is little to be gained in terms of future fertility from saving a diseased tube if the other tube appears normal. Salpingectomy will be the most commonly performed laparoscopic procedure.

11.3 Criteria for laparoscopic salpingotomy
This procedure should be considered when all of the following criteria are met:
• Future fertility is desired
• Patient is haemodynamically stable
• Tubal pregnancy confirmed by diagnostic laparoscopy
• No fetal heartbeat seen on ultrasound
• Diameter of tubal pregnancy less than 4 cm
- Good haemostasis maintained during the procedure
- Good access to tube (no dense bowel adhesions or peritoneal adhesions)
- Tube (especially the fimbrial end) should be normal
- Serum ß-hCG levels should be less than 5000 IU/l
- Concurrent pathology requiring laparotomy is not present.

Five to 15% of patients having a laparoscopic salpingotomy will have persistent trophoblastic tissue, they may require methotrexate or further surgical management. Therefore, weekly ß-hCG measurement is essential (Yao & Tulandi, 1997).

Patients who have conservative tubal surgery have a 12 - 15% risk of an ectopic pregnancy in subsequent pregnancies (Stabile, 1996).

**Blood transfusion**
Transfusion is usually intra-operative and sometimes required post-operatively. Patients should always be counselled about the possibility of transfusion.

12. **Other issues**

12.1 **Risk of future ectopic pregnancy**
Depends on:
- Number of previous ectopic pregnancies
- State of remaining tube
- Type of treatment to present ectopic pregnancy.

First ectopic with remaining tube normal, 6-12% chance of another ectopic.

First ectopic with remaining tube abnormal, 28-50% chance of another ectopic.

More than one ectopic pregnancy, 26-40% chance of another ectopic (Stabile, 1996).

13. **Management guidelines of ruptured ectopic pregnancy**
If a referring doctor calls to admit a patient with a diagnosis of suspected ruptured ectopic pregnancy, the following advice should be given:
- If the patient is haemodynamically stable:
  - Secure good intravenous access with a large-bore needle and infuse intravenous (IV) fluids.
  - Arrange transfer by ambulance to the Women’s Assessment Unit (WAU).
  - Collect blood for full blood count, group and cross-match and ß-hCG.
  - Initial counselling for possible surgical management.
  - Arrangements to be made by Women’s Health:
    - Advise Clinical Nurse Manager and WAU of possible ruptured ectopic admission.
    - Anaesthetic and theatre staff to be informed.
    - Registrar/specialist assessment to decide on further investigations and surgical intervention, which may be laparoscopy or laparotomy.
- If ruptured ectopic and the patient is unstable:
  - Trendelenburg position.
  - Two large-bore needles for IV access and colloid infusion if available or crystalloids.
  - Oxygen to be administered.
  - Immediate ambulance transfer to WAU with the General Practitioner (GP) accompanying patient.
  - Collect blood for FBC and cross-match.

Arrangements to be made by Women’s Health:
- Specialist on call to be informed and to attend to manage patient with registrar. Registrar to notify Charge Nurse and the Clinical Nurse Manager.
- Patient should be directed to theatre from Emergency Department or WAU.
- Anaesthetic and Operating Theatre staff available for resuscitation and stabilisation of patient for immediate laparotomy.
- Uncross-match blood may be used in rare circumstances only.

14. Checklist for discharge of women with ectopic pregnancy

14.1 Process
Follow the steps below when discharging a patient who has had an ectopic pregnancy diagnosis:

<table>
<thead>
<tr>
<th>Step</th>
<th>Action</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>Discuss the surgical procedure and findings within the rest of the pelvis in detail, e.g. adhesions, ovaries, and condition of fallopian tubes.</td>
</tr>
</tbody>
</table>
| 2.   | Discuss follow-up:  
  - **Salpingectomy** either by laparoscopy or laparotomy does not require follow-up with serial β-hCG unless specifically requested.  
  - **Salpingotomy** either by laparoscopy or laparotomy requires follow-up with serial β-hCG measurement because 5-10% of these patients may require further treatment. This is generally done through WAU. |
| 3.   | **Medical management**: Ensure follow-up blood tests are arranged (refer to protocol). |
| 4.   | Full blood count if:  
  - Blood was transfused  
  - Blood loss was greater than 500 mL  
  - Requested by surgeon  
  - Clinically indicated  
  - Iron replacement may be necessary. |

14.2 Rhesus status
Currently, it is recommended that rhesus negative women are given Anti-D1:  
- Check blood group and Rhesus status  
- Anti-D given within 72 hours if indicated.
14.3 Contraception and future fertility

Patients are generally advised to delay another pregnancy until recovered both physically and emotionally from this pregnancy. The chance of another ectopic pregnancy should be explained. Patients are advised to have a pregnancy test on missing her menses and if pregnant, a transvaginal ultrasound scan should be arranged to locate the pregnancy. Patients with bilateral tubal disease or salpingectomy should be referred to discuss IVF if future fertility is desired.

The IUCD may be used with caution in women with a history of ectopic pregnancy.

The progestogen-only pill is contra-indicated in women with a history of ectopic pregnancy.

*It is important to rule out ectopic pregnancy early in all future pregnancies.*

**Note:** Unless specifically requested, most of these patients are followed by their GP or Early Pregnancy Assessment Unit (EPAU) and a copy of the histology should be available within two to four weeks.

15. Supporting evidence


16. Associated documents
• Informed Consent
• Blood Components and Blood Products Administration

Patient information
• Ectopic Pregnancy
• Methotrexate in Ectopic Pregnancy

Other
• Refer to the Medsafe website for up to date information on Methotrexate

17. Disclaimer

No guideline can cover all variations required for specific circumstances. It is the responsibility of the health care practitioners using this Auckland DHB guideline to adapt it for safe use within their own institution, recognise the need for specialist help, and call for it without delay, when an individual patient falls outside of the boundaries of this guideline.

18. Corrections and amendments

The next scheduled review of this document is as per the document classification table (page 1). However, if the reader notices any errors or believes that the document should be reviewed before the scheduled date, they should contact the owner or Document Control without delay.
19. Appendix 1: Standard operating policy: Ectopic pregnancy – direct administration of methotrexate under ultrasound guidance

Performed in fetal medicine department

Aim: Administration of potassium chloride to fetal heart and methotrexate into gestational sac under ultrasound guidance.

If there is no cardiac activity, only administration of methotrexate into the sac.

Preparation ahead of procedure

- Procedure is done in procedure room in fetal medicine clinics – should be scheduled.
- Patient needs to have a valid group and hold, be nil by mouth for at least six hours.
- Patient needs to have intravenous (IV) access (preferably green or grey).
- People that need to be notified prior to the procedure should include:
  - Theatre coordinator, 021 471 618
  - Acute on-call team or WAU consultant
  - Consider contacting interventional radiology.
- Methotrexate 50 mg, and the protective gowns and goggles needed for administration, waste bags and needle containers for disposal can be obtained from WAU.
- Spill kit should be on hand.
- Patient comes with her bed to the fetal medicine clinics so she can be transported to theatre or radiology should bleeding occur.
- Informed written consent.

Equipment

- Philips EPIC used for the procedure, with transvaginal probe and needle guide OR transabdominal probe.
- Sterile needle guide that fits on the vaginal probe and 18-gauge needle for the procedure are stored in the back room.
- Sterile condom and gel are included in the pack containing the transvaginal needle guide.
- Every operator in contact with the methotrexate needs to wear a protective gown, glasses or goggles for eye protection, a face mask to cover mouth and nose, and protective sterile gloves.
- For transvaginal approach procedure trolley should be set up with:
  - Speculum, sponge holder forceps, gauze, water-based (e.g. saline) antiseptic solution
  - Long 18-gauge needle, sterile needle guide
  - Condom for transvaginal probe and sterile gel (contained in needle guide pack)
  - Syringe with 2 mL of 20 mmol potassium chloride if required
  - Syringe with 2 mL methotrexate = 50 mg.
- For transabdominal approach procedure trolley should be set up with:
  - Standard invasive procedure equipment
  - Probe cover and gel
  - Syringe with 2 mL of 20 mmol potassium chloride if required
  - Syringe with 2 mL methotrexate = 50 mg
  - Disposable drapes (folio drapes, that are used as trolley drape).

Procedure

- Fentanyl IV (50 to 100 mg) is administered prior to the procedure.
Monitor maternal heart rate and oxygen saturations with probe.
The floor underneath the procedure area is covered with inco sheets to prevent spill of methotrexate on the floor.
The patient is positioned in lithotomy position (use lithotomy poles) for transvaginal approach.
The cervix is cleaned prior to the procedure using a speculum, spongeholder forceps and gauzes drenched in water-based antiseptic solution/sterile saline.
The biopsy guide is activated on the screen to show the needle trajectory, the vaginal probe is inserted such that the needle is in line with the fetal heart. The needle is inserted through the amniotic sac into the fetal thorax. 1 mL potassium chloride is administered. Extra potassium chloride may need to be administered until asystole achieved. The needle is then withdrawn into the amniotic or chorionic cavity and 50 mg of methotrexate (in 2 mL) is administered. The **syringe then remains attached to the needle** when the needle is withdrawn to prevent spill of methotrexate.

**After the procedure**
- A new 18-gauge needle needs to be ordered before or after the procedure because we only have two needles in stock.
- All material that has been in contact with methotrexate needs to be disposed of in special purple bag and purple sharps box. These need to go to the purple waste disposal bin located in the waste disposal room. This is found behind lift bank C on Level 9.
- Administration of Anti-D as required if the patient is Rhesus negative.

**In case of a bleeding complication**
- The patient’s own team or the acute on-call team take over to go to theatre acutely.
- Options to stop the bleeding:
  - Placement of a size 26 french 90 mL Foley catheter into the cervix, a purse string suture around the cervix may be needed to keep it in place, if this stops the bleeding observation for 24-48 hours before reducing amount of fluid in the balloon.
  - Suturing the cervix at three and nine hours just below the lateral fornix to include the cervicovaginal branches of the uterine artery.
  - Interventional radiology, or bilateral internal iliac artery ligation, bilateral uterine artery ligation.
  - If all else fails: hysterectomy.
20. Appendix 2: Serial serum beta hCG form

![Serial serum beta hCG form](image)

**hCG in normal pregnancy**

The developing placenta begins producing hCG about 3 days after implantation. The sequence of events is:

<table>
<thead>
<tr>
<th>Gestation (Weeks)</th>
<th>Mean hCG (iu/l)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1st day LMP (last monthly period)</td>
<td>0</td>
</tr>
<tr>
<td>fertilisation</td>
<td>2</td>
</tr>
<tr>
<td>implantation</td>
<td>3</td>
</tr>
<tr>
<td>placenta produces measurable hCG</td>
<td>≥ 2x</td>
</tr>
<tr>
<td>1st day of missed periods</td>
<td>4</td>
</tr>
<tr>
<td>gestational sac becomes visible on ultra-sound</td>
<td>5</td>
</tr>
<tr>
<td>peak hCG level</td>
<td>9</td>
</tr>
</tbody>
</table>

ref. The Interpretation of Laboratory Tests, Diagnostic Medlab Auckland August 2000.