Staging Classifications and Clinical Practice
Guidelines for Gynaecological Cancers

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I would like to thank L Zigliani
for her contribution to this handbook.
Preface

The third edition of the Good Practice Guidelines on Gynaecologic Cancers was a collaborative effort between FIGO and the International Gynecologic Cancer Society. Both organisations put women’s health and combat female cancers as their priority objectives. New advances were added to this new edition. The guidelines are based on strongest available evidence which varies from topic to topic. The evidence has been graded as follows:

- A – randomised controlled trial
- B – prospective (cohort) study with a comparison group
- C – retrospective follow-up study
- D – cross sectional study

However, guidelines on breast cancer have not been revised. It is envisaged that further revision will be needed in future. In the mean time, based on current evidence, these guidelines could serve as a good reference where adaptation to local use could be based on.

Hextan YS Ngan
Chairman of FIGO Committee on Gynecologic Oncology

The International Gynecologic Cancer Society (IGCS) is pleased to be able to continue the collaboration with FIGO to develop treatment guidelines for gynaecologic malignancies. The guidelines for all primary gynaecological sites have been updated, with significant recent advances included in the text and in the bibliography. The final product is the result of input from many specialists in both organisations.

These guidelines are intended to represent a reasonable synthesis of the current literature for each disease site, though they are not intended to be absolutely prescriptive. Patient management always needs to be individualised, taking into account a patient’s general medical condition, her cultural environment, and the available expertise and technology.

Neville F Hacker
Chairman of the IGCS Guidelines Committee
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Principles of Cancer Staging

The major tasks faced by a clinician having made a diagnosis of cancer are to determine the most effective therapy and to formulate a prognosis for the patient. In order to optimally manage a cancer, both the extent of the disease and knowledge of its biology are essential. The extent of the disease is generally expressed in terms of its stage. The major purpose of staging that has been agreed upon internationally is to offer a classification of a cancer’s extent so as to provide a method of conveying one’s clinical experience to others for the comparison of treatment methods without confusion or ambiguity.

Cancer staging is central to the modern management of cancer patients. Cancer is also a biologic continuum and a dynamic process, which is artificially compartmentalised by staging systems. It is clear, however, that the phases or substages must have clinical relevance. Cancer staging systems should also be evidence-based and they should be user friendly. Staging systems need to be based on the best available knowledge at hand and this implies that the changes will occur over time based upon the development or the acquisition of new knowledge.

It also follows that the acquisition of this knowledge is facilitated by the use of the staging system insofar as staging will help with knowledge creation by facilitating clinical research, producing new data on similar groups of patients, and integrating these new data about similar patients from diverse sources. Staging also helps knowledge dissemination by providing a common international language for information sharing, and facilitates the teaching of both new and established health care workers.

Gynaecologists have a long and proud tradition of using staging systems for female cancers, dating back to the League of Nations staging system for cervical cancer, first published in 1920. In 1954 FIGO assumed the patronage of the Annual Report on the Results of Treatment in Gynecological Cancer. With it also came the responsibility for overseeing the staging of gynaecological cancers, which were at the heart of the Annual Report data and information system. Since that time the FIGO Committee on Gynecologic Oncology has made several modifications to the various staging systems for gynaecological cancer, most notably those for cervix and endometrial cancer. 1954 also saw the UICC set up a committee on clinical stage
classification and applied statistics, which had as its aim the extension of the general technique of classification of cancer at all sites by anatomical extent of the disease using the TNM system.

The FIGO system of classification was originally based on clinical examination, essentially of the anatomical extent of disease. Over the years all staging systems for gynaecological cancers, with the exception of cervical cancer, have moved from a clinical basis to one of a surgical pathological nature.

Stage I has generally been referred to as early stage disease, where the lesion would appear to be confined to the organ of origin. Stage II is a general description of disease that has extended locally beyond the site of origin to involve adjacent organs or structures. Stage III then represents more extensive involvement, with stage IV representing clearly metastatic disease. The basic stages are then generally classified into substages, which are usually a reflection of specific prognostic factors within a given stage.

Tumour classifications may be based on many systems. For example, the anatomical site of the disease and the clinical and pathological extent of disease. Similarly the histologic type and grade of tumours, as well as age of patient and the duration of signs and symptoms are all known to have an influence on the outcome of disease and have all been used in various staging systems. The TNM system describes the anatomical extent of disease based on the assessment of three primary components. T refers to the extent of the primary tumour, N to the presence or absence and extent of regional lymph node metastases, and M the presence or absence of distant metastases. The TNM system is also further classified into two groups. The cTNM system, which is essentially a pre-treatment clinical classification based on evidence acquired before treatment from physical examination, imaging, biopsy, endoscopy, surgical exploration and other relevant examinations. The pTNM system is the other sub-group and is based on post-surgical histopathological classification. This uses evidence acquired before treatment once again, supplemented or modified by the additional evidence acquired from surgical and from pathological examination. After assigning T, N, M and/or pT, pN and pM categories, these items may then be grouped into stages. The classification system and stage grouping, once established, must remain unchanged in medical records. Clinical stage is essential to select and evaluate therapy, while the pathological stage provides the most precise data to estimate prognosis and calculate end results. The FIGO and TNM classifications are virtually identical. The TNM Prognostic Factor Project Committee has graciously agreed to defer to all questions regarding staging of gynaecological cancer to the FIGO Committee on Gynecologic Oncology.

In conclusion, any good staging system must have three basic characteristics. It must be valid, reliable, and, above all, it must be practical. Validity means that the staging system must allow for the creation of groups of cases that experience similar outcomes, while at the same time reflecting a full range of possible presentations for each type of cancer. Also over time, the system in order to retain its validity must be flexible so that it can adapt to important changes in medical care.

A reliable staging system should ensure that identical cases would always be assigned to the same stage category. It should be unambiguous; it should be based as far as possible on measurable entities that have been evaluated objectively. The system should also not be subject to frequent changes until sufficient data and information are obtained to warrant such changes.

Finally, a practical staging system must be suitable for day to day use in a wide range of clinical environments and must not require diagnostic procedures that are not readily available to most practitioners or extraordinary expertise or knowledge regarding a particular malignancy.

Principles Governing the Development of Good Practice Guidelines
The following basic concepts and principles were used to produce these clinical practice guidelines.

Definition:
Clinical practice guidelines (CPG) or good practice guidelines (GPG) are systematically developed statements to assist health care practitioners and providers in making decisions about appropriate care for patients in specific clinical circumstances.

Purpose:
The purpose of CPG’s may vary depending on ones perspective; eg a
clinician may see them as a method of improving patient outcomes. Administrators and other societal authorities may see them as helpful in viewing distribution and allocation of health funding and appropriated (related) resources.

**Objectives:**
The objectives of CPG should be to reduce inappropriate variation in clinical practice. This may also serve to prevent resource limitations from adversely affecting patient care and may also be used as a way of influencing health care providers to incorporate new strategies into the clinical care of their patients based on new scientific evidence.

**Statement:**
Clinical practice guidelines should always be based on evidence. However, high levels of evidence are not always available for certain situations. A distinction must be made between Evidence and Beliefs. CPG must also consider benefits versus harm versus evidence. In terms of belief, care needs to be taken because patient advocacy movements may grasp these items and use them for their own personal agendas and at times what patients want may be markedly different from what the evidence might show or what might be appropriate treatment.

**Guideline Development:**
- Guidelines capture evidence and/or beliefs at a point in time and therefore they will need to change in the future.
- Guidelines should consider social and cultural differences.
- Guidelines should be enabling and not prescriptive in nature.
- One needs to be careful that physician-created guidelines are not taken over by third parties and made mandatory. This we wish to avoid.
- Anything that is mandatory should not be called a guideline. It might be better termed recommendation.
- Guidelines should offer rather than saying “one should treat”.

**Concerns:**
a) Many groups are developing guidelines and it is difficult not to produce guidelines that may not be in conflict with some other body.

b) Guidelines should not be too rigid so as to actually hamper physician decision-making freedom.
c) Legal implications and the threat of medico-legal action need to be considered if one is practising outside of so called guidelines.
d) Individuals may be unaware that guidelines actually exist for particular scenarios.
Cancer of the Vulva

1.1 Staging

1.1.1 Anatomy.
Cases should be classified as carcinoma of the vulva when the primary site of growth is in the vulva. Tumours present in the vulva as secondary growths, from either a genital or extra-genital site, have to be excluded. Malignant melanoma should be separately reported and staged according to the system for cutaneous melanomas. A carcinoma of the vulva that extends into the vagina should be considered as a carcinoma of the vulva. There must be histologic confirmation of the cancer.

1.1.1.1 Nodal stations.
The inguinal and femoral nodes are the primary sites of regional spread.

1.1.1.2 Metastatic sites.
Involvement of pelvic lymph nodes (external, hypogastric obturator and common iliac) are considered distant metastases.

1.1.2 Surgical staging classification.
Vulvar cancer has been surgically staged since 1988. The final diagnosis is dependent upon thorough histopathologic evaluation of the operative specimen (vulva and lymph nodes). Various modifications have been made with a subdivision of Stage I in 1994.

1.1.2.1 Regional lymph nodes (N).
- NX – Regional lymph nodes cannot be assessed;
- NO – No regional lymph node metastasis;
- N1 – Unilateral regional lymph node metastasis;
- N2 – Bilateral regional lymph node metastases.

1.1.2.2 Distant metastasis (M).
- MX – Distant metastasis cannot be assessed;
- M0 – No distant metastasis;
- M1 – Distant metastasis.

<table>
<thead>
<tr>
<th>FIGO Stages</th>
<th>TNM Categories</th>
</tr>
</thead>
<tbody>
<tr>
<td>Primary tumour cannot be assessed</td>
<td>TX</td>
</tr>
<tr>
<td>No evidence of primary tumour</td>
<td>T0</td>
</tr>
<tr>
<td>0 Carcinoma in situ (preinvasive carcinoma)</td>
<td>Tis</td>
</tr>
<tr>
<td>I Tumour confined to vulva or vulva and perineum, 2 cm or less in greatest dimension</td>
<td>T1</td>
</tr>
<tr>
<td>IA Tumour confined to vulva or vulva and perineum, 2 cm or less in greatest dimension and with stromal invasion no greater than 1.0 mm*</td>
<td>T1a</td>
</tr>
<tr>
<td>IB Tumour confined to vulva or vulva and perineum, 2 cm or less in greatest dimension and with stromal invasion greater than 1.0 mm*</td>
<td>T1b</td>
</tr>
<tr>
<td>II Tumour confined to the vulva or vulva and perineum, more than 2 cm in greatest dimension</td>
<td>T2</td>
</tr>
<tr>
<td>III Tumour invades any of the following: lower urethra, vagina, anus and/or unilateral regional node metastasis</td>
<td>T3</td>
</tr>
<tr>
<td>IV Tumour invades any of the following: bladder mucosa, rectal mucosa, upper urethral mucosa; or is fixed to bone and/or bilateral regional node metastases</td>
<td>T4</td>
</tr>
<tr>
<td>IVA Any distant metastasis including pelvic lymph nodes</td>
<td></td>
</tr>
</tbody>
</table>

* The depth of invasion is defined as the measurement of the tumour from the epithelial-stromal junction of the adjacent most superficial dermal papilla, to the deepest point of invasion.
1.2 Introduction

Carcinoma of the vulva is an uncommon tumour, representing about 4% of gynaecologic malignancies. Because of the relatively small experience of individual institutions, randomised trials of therapeutic approaches are uncommon, and most studies are based on retrospective clinico-pathologic reviews\(^{(1)}\).

<table>
<thead>
<tr>
<th>FIGO Stage</th>
<th>UICC T</th>
<th>N</th>
<th>M</th>
</tr>
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<tbody>
<tr>
<td>0</td>
<td>Tis</td>
<td>N0</td>
<td>M0</td>
</tr>
<tr>
<td>IA</td>
<td>T1A</td>
<td>N0</td>
<td>M0</td>
</tr>
<tr>
<td>IB</td>
<td>T1B</td>
<td>N0</td>
<td>M0</td>
</tr>
<tr>
<td>II</td>
<td>T2</td>
<td>N0</td>
<td>M0</td>
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<tr>
<td>III</td>
<td>T1</td>
<td>N1</td>
<td>M0</td>
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<tr>
<td></td>
<td>T2</td>
<td>N1</td>
<td>M0</td>
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<td></td>
<td>T3</td>
<td>N1</td>
<td>M0</td>
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<tr>
<td>IVA</td>
<td>T1</td>
<td>N2</td>
<td>M0</td>
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<td>T2</td>
<td>N2</td>
<td>M0</td>
</tr>
<tr>
<td></td>
<td>T3</td>
<td>N2</td>
<td>M0</td>
</tr>
<tr>
<td>IVB</td>
<td>Any T</td>
<td>Any N</td>
<td>M1</td>
</tr>
</tbody>
</table>

1.1.2.3 Histopathologic types.

Squamous cell carcinoma is the most frequent form of cancer of the vulva. Malignant melanoma is the second most common tumour and should be reported separately. Other histopathologic types are:
- adenocarcinoma underlying Paget’s disease of vulva, verrucous carcinoma, Bartholin gland carcinoma, adenocarcinoma not otherwise specified (NOS), basal cell carcinoma.
- Histopathologic grades (G).
  - Gx – Grade cannot be assessed;
  - G1 – Well-differentiated;
  - G2 – Moderately differentiated;
  - G3 – Poorly or undifferentiated.
It is predominantly a disease of post-menopausal women, the age specific incidence increasing with increasing age. The external location of the vulva should prompt early presentation, but traditionally significant delays in diagnosis have been common with this cancer.

Ninety percent of cancers are squamous in origin, while melanomas, adenocarcinomas, basal cell carcinomas, verrucous carcinomas, sarcomas, and other rare malignancies also occur. Most squamous carcinomas occur on the labia majora but the labia minora, clitoris, and perineum may also be primary sites.

Vulvar intraepithelial neoplasia (VIN) tends to occur in younger women and may be associated with similar lesions of the cervix and vagina. VIN III is a precursor lesion in some patients, and should be effectively treated by superficial excision, with or without laser therapy, when diagnosed. Treatment of vulvar cancer has evolved into an individualised multidisciplinary approach, and patients should be referred centrally to a gynaecological cancer centre where all relevant expertise is available. Level of evidence B.

1.3 Screening
There is no screening procedure for vulvar cancer. However patients with a past history of cervical or vaginal cancer should have inspection of the vulva, with or without colposcopic examination, as part of their regular follow-up. Patients with lichen sclerosis or a past history VIN III should also be kept under regular surveillance.

1.4 Squamous Cell Carcinoma

1.4.1 Presenting symptoms
Vulvar cancer may be asymptomatic, but most patients present with a vulvar lump or ulcer. There is often a long standing history of pruritus, which may be due to associated vulvar dystrophy. Bleeding or discharge is an occasional presenting symptom, and patients with advanced disease may present with a lump in the groin.

1.4.2 Diagnosis
Diagnosis should be confirmed by biopsy prior to definitive treatment. A wedge biopsy under local anaesthesia in the office is usually sufficient. The biopsy should include some surrounding skin and underlying stroma.

It is preferable not to excise the entire lesion as it makes it more difficult to plan the definitive excision.

If the lesion is 2 cm or less in diameter and depth of stromal invasion is ≤ 1 mm on wedge biopsy, complete excision of the lesion must be undertaken to allow serial sectioning to properly assess the depth of invasion. The Committee on Gynecologic Oncology of FIGO measures depth of stromal invasion from the epithelial-stromal junction.

1.4.3 Investigations
1. Pap smear of the cervix if cervix is still in situ
2. Colposcopy of the cervix and vagina, because of the common association with other squamous intraepithelial lesions.
3. A CT scan of the pelvis and groins is often helpful to detect any enlarged lymph nodes in the groins or pelvis.
4. Routine full blood count, biochemical profile and chest x-ray preoperatively.

1.4.4 Clinical Practice Guidelines
The clinical findings should be recorded on a staging diagram (eg Fig 1). The material in Tables 1 and 2 is usually listed on the reverse side of the diagram.

1.4.5 Treatment

1.4.5.1 Treatment of vulvar intraepithelial neoplasia (VIN) or carcinoma in situ
Various treatment modalities are available for treating intraepithelial lesions of the vulva. Initial assessment should consist of multiple biopsies to ensure that the lesion is entirely intraepithelial. Patients with multifocal lesions should have biopsies taken from several lesions. Once the diagnosis has been established, superficial local excision of the vulvar epithelium with a 0.5-1.0-cm margin is considered adequate for lesions of the lateral aspect of the vulva. Lesions involving the labia minora may also be treated by local excision but may respond favourably to laser vaporisation or ablation. Laser treatment of the hair-bearing skin of the vulvar epithelium will usually...
produce depigmentation and destruction of hair follicles with subsequent loss of hair growth. Laser is also appropriate for clitoral lesions. Large lesions may be treated with a skinning vulvectomy and split thickness skin graft. **Level of evidence C.**

1.4.5.2 Invasive vulvar cancer
Management of vulvar cancer must be individualised. There is no standard operation and the emphasis is on performing the most conservative operation consistent with cure of the disease\(^1\).

In considering treatment options, it is necessary to consider independently the most appropriate management of
1. the primary lesion
2. the groin lymph nodes

1.4.5.2.1 Microinvasive vulvar cancer (Stage IA)
Stage IA carcinoma of the vulva is defined as a single lesion measuring 2 cm or less in diameter with a depth of invasion of 1.0 mm or less, the depth being measured from the epithelial-stromal junction of the most adjacent superficial dermal papilla to the deepest point of invasion. Lesions of this extent should be managed with wide local excision. If the local resection reveals features that are unfavourable (neural or vascular invasion), consideration should be given to more radical excision. Groin dissection is not necessary for lesions of this type\(^2\). **Level of evidence C.**

1.4.5.2.2 Early vulvar cancer
Tumours confined to the vulva without clinically suspicious lymph nodes may be considered early.

1.4.5.2.2.1 Management of the primary lesion (Figure 2)
In order to decrease psychosexual morbidity, a more conservative operation than radical vulvectomy usually is indicated. The procedure may be called a radical local excision, and for localized lesions, this operation is as effective as radical vulvectomy in preventing local recurrence\(^3\-12\).

Surgical removal should achieve lateral margins of at least 1 cm, and the deep margin should be the inferior fascia of the urogenital diaphragm, which is coplanar with the fascia lata and the fascia over the pubic symphysis\(^13\).

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**Figure 2: Management of early vulvar cancer**

If the lesion is close to the urethra, the distal 1 cm of the urethra may be resected without jeopardizing urinary continence.

If there is associated VIN or lichen sclerosis, these areas may be superficially excised to control symptoms and to exclude other areas of superficial invasion. **Level of Evidence C.**

1.4.5.2.2 Management of groin lymph nodes
Recurrence in the groin carries a very high mortality so appropriate
groin dissection is the single most important factor in reducing mortality from early vulvar cancer\(^1\).

All patients with T2 lesions and all patients with T1 tumours with >1 mm stromal invasion should have at least an ipsilateral inguino-femoral lymphadenectomy. **Level of Evidence C.**

The incidence of positive contralateral nodes in patients with lateral T1 tumours is < 1%, so unilateral groin dissection is appropriate for such lesions\(^{11}\).

Bilateral groin dissection should be performed for midline tumours, and for those involving the anterior labia minora\(^{14}\). Large lateral tumours should probably also have bilateral dissection, particularly if the ipsilateral nodes are positive\(^{11}\). Sentinel node excision is still experimental, and should be performed only within a clinical trial\(^{15}\). **Level of Evidence B.**

1.4.5.2.2.3 Groin dissection

It is recommended that both inguinal and femoral nodes be removed, as inguinal node dissection alone is associated with a higher incidence of groin recurrence\(^{16}\). **Level of Evidence A.** The femoral nodes are situated medial to the femoral vein within the fossa ovalis. There is no need to remove the fascia lata to dissect the femoral nodes\(^{17}\).

Groin dissection may be safely performed through a triple incision approach as this should improve primary healing\(^{18,19}\). **Level of Evidence B.** Alternatively, an en bloc approach may be used, particularly for clitoral or periclitoral lesions. To avoid skin necrosis, all subcutaneous tissue above the superficial fascia must be preserved.

Groin dissection (with post-operative radiation for patients with positive groin nodes) was superior to groin irradiation in one randomized trial, although the depth dose was considered to have been inappropriate in this study\(^{20}\).

**Management of patients with positive groin nodes**

The Gynecologic Oncology Group demonstrated superior results for pelvic and inguinal radiation compared to pelvic node dissection for patients with grossly positive groin nodes, or more than one microscopic positive node\(^{21}\). **Level of Evidence A.**

Subsequent studies have further emphasized the prognostic significance of the morphology of positive groin nodes, particularly the size of the metastasis and the presence or absence of extracapsular spread\(^{22-24}\).

Patients with one (and possibly two) micrometastases (< 5 mm) do not require adjuvant radiation therapy.

Patients should receive bilateral pelvic and groin irradiation for the following indications:

- one macrometastasis (> 5 mm diameter)
- extracapsular spread
- two (possibly three) or more micrometastases (≤ 5 mm)

**Level of Evidence C.**

1.4.5.2.2.4 Radiation fields and doses

In most cases, fields should include the inguino-femoral nodes and at least the lower pelvic nodes (below the SI joints).

One of a variety of radiation techniques can be selected, depending on the patient’s body habitus, and extent of disease.

Combined photon and electron techniques are often used to treat the regional nodes without overdosing the femoral heads. However, care must be taken to completely include both the superficial and deep inguinal lymph nodes.

It is important to avoid underdosage of superficial inguinal nodes by high energy photon beams. If electron beams are used, the energy must be sufficient to cover the deep inguinal nodes.

In most cases, CT-based treatment planning should be used to verify adequate coverage.

The dose of radiation is determined by the initial extent of regional disease and any known residual. After a groin dissection with microscopic inguinal metastases, 50 Gy in 1.8-2.0 Gy fractions is usually sufficient.

If there are multiple positive nodes or if there is evidence of extracapsular extension, somewhat higher doses up to 60 Gy may be given to a reduced volume. Gross residual disease may require doses of 60-70 Gy.

The role of concurrent chemotherapy in this setting is unknown.

1.4.5.2.3 Advanced vulvar cancer

Patients with T3 or T4 primary tumours or bulky positive groin nodes
are considered to have advanced vulvar cancer. For such patients, multimodality treatment planning is particularly important.

1.4.5.2.3.1 Management of the groin lymph nodes

It is desirable to determine the status of the groin nodes prior to planning the overall treatment (11).

If there are no suspicious nodes palpable in the groin, bilateral inguino-femoral lymphadenectomy should be performed. If final histologic assessment reveals positive nodes, adjuvant radiation to the groin and pelvis should follow the guidelines given for early stage disease.

If there are suspicious nodes in the groin, a pre-operative CT scan may help identify the extent of groin and pelvic lymphadenopathy (Figure 3).

Figure 3: Management of clinically suspicious groin nodes

- Clinically suspicious nodes
  - CT-scan of pelvis
  - Resection of macroscopic groin nodes and frozen section
    - Positive
      - Retroperitoneal resection of any macroscopic pelvic nodes seen on CT-scan
        - Pelvic and groin radiation therapy
    - Negative
      - Inguino-femoral lymphadenectomy
        - Two or more positive nodes or extracapsular spread
          - Observation

Resection of all enlarged groin nodes should be performed, and frozen section diagnosis obtained.

If nodes are negative, full inguino-femoral lymphadenectomy should be performed.

If nodes are positive, a complete lymphadenectomy should probably be avoided because full-groin dissection together with post-operative groin irradiation may result in severe lymphoedema. Only enlarged nodes from the groin and pelvis should be removed, and the patient given post-operative groin and pelvic radiation (12).

Level of Evidence C.

If there are ulcerated or fixed groin nodes, they should be biopsied to confirm the diagnosis then treated with primary radiation. When feasible, the nodes should be resected following the radiation (Figure 4) (12). Level of Evidence C.

Figure 4: Management of clinically obvious groin nodes:

- Fixed or ulcerated nodes
  - Surgically resectable
    - Resection of all macroscopic nodes in groin and any enlarged pelvic nodes on CT
      - Post-operative radiotherapy to groins and pelvis
  - Unresectable
    - Pre-operative radiotherapy ± chemotherapy
      - Post-operative resection of macroscopic residual disease
### 1.4.5.2.3.3 Radiation Protocol

If the groin nodes are positive and meet the requirements described earlier for adjuvant radiation, the initial radiation treatment fields should include the pelvis, inguinal nodes, and primary site, which are treated to a total dose of at least 50 Gy. Care must be taken to adequately cover the inguinal nodes.

Some clinicians prefer to treat in an open-leg position but care must be taken to apply bolus to the vulva to avoid underdosage of the skin. Areas of gross disease or particularly high risk are usually boosted with appositional fields of electrons selected to provide an adequate dose to the surface and at depth. Gross vulvar disease probably requires 60-70 Gy to achieve local control, although investigators are currently exploring a wide variety of chemo-radiation schedules, and the relationship between dose and local control remains somewhat uncertain. **Level of Evidence C.**

### 1.4.5.2.3.4 Close surgical margins

Post-operative radiation may be used for close surgical margins (< 5 mm), if the margins cannot be re-excised. Although local control is improved, overall survival is not significantly different because of the ability to salvage patients with local recurrence.

In some cases, the positive margin may be boosted with brachytherapy, although this technique requires experience to avoid an excessive risk of necrosis. Alternatively, the operative bed may be treated with an appositional electron field. **Level of Evidence C.**

### 1.4.5.2.3.2 Management of the primary tumour (Figure 5)

This should usually follow dissection of the groins.

If it is possible to resect the primary lesion with clear surgical margins and without sphincter damage leading to urinary or faecal incontinence, primary surgical excision is desirable.

If primary surgery would result in the need for a bowel or urinary stoma, it is preferable to employ primary radiation therapy, followed by a more limited resection of the tumour bed.

Chemo-radiation has been used, sometimes without need for surgical resection of the tumour bed.

The groin nodes and pelvis may need to be included in the treatment field depending on the status of the groin nodes, as determined initially.

### 1.5 Special situation

#### 1.5.1 Vulvar melanoma

Vulvar melanoma is the second most common neoplasm of the vulva. The majority of lesions involve the clitoris or labia minora. The Clark or Breslow modifications to the micro staging system should be used for the staging of vulvar melanoma rather than the more common TNM/FIGO system. These systems measure the depth of invasion in terms of the descriptive histology of the skin.

Any pigmented lesion on the vulva should be excised for diagnosis unless it has been known to be present and unchanged for some years.

In line with trends toward more conservative surgery for cutaneous melanomas, there is a trend toward more conservative resection of...
population. Most patients will present with vulvar discomfort and itching and on examination an eczematoid-weeping lesion is often seen. Diagnosis is usually confirmed by biopsy which usually will establish if one is dealing with an intraepithelial or invasive lesion (1,38).

Intraepithelial Paget’s disease requires superficial local excision. It is difficult to obtain clear margins with this disease, as often the underlying histologic change will seem to extend far beyond what is clinically visible. More recently, there has been a move to perform less radical resection for intraepithelial lesions, with re-excision at a later date should lesions become symptomatic or clinically visible. Lesions that involve or extend into the urethra or anus can be particularly difficult to manage, and may require laser therapy.

If there is an underlying adenocarcinoma, the invasive component should be treated by radical local excision with margins of at least 1 cm. At least an ipsilateral inguino-femoral lymphadenectomy should be performed for unilateral lesions with adjuvant radiation following the same indications as for squamous carcinomas. Level of Evidence C.

1.5.3 Paget’s Disease
This is predominantly an intraepithelial lesion, but on occasion it may be associated with an underlying invasive adenocarcinoma. The disease occurs predominantly in the menopausal or post-menopausal population. Most patients will present with vulvar discomfort and itching and on examination an eczematoid-weeping lesion is often seen. Diagnosis is usually confirmed by biopsy which usually will establish if one is dealing with an intraepithelial or invasive lesion (1,38).

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1.6 Pathology
The surgical specimen should be correctly orientated and photographed. The photograph should be used to indicate the origin of tissue blocks.

The size of the specimen should be measured and the dimensions of any visible tumour measured.

The macroscopic tumour-free surgical margins should be measured. Sections are taken through the tumour to measure tumour depth. Sections should be taken from urethral, anal and vaginal resection margins.

Lymph nodes should be carefully dissected out and the site from which they are removed recorded. A full cross-section of each lymph node should be embedded.

The following histological points should be noted:

a. Tumour type:
   - keratinizing
   - basaloid
   - Bowenoid
b. Depth of invasion
should be measured from the epithelial-stromal junction of the
adjacent dermal papilla to the deepest point of invasion by the
tumour.

c. Tumour grade
noted, but not proven to be prognostically significant.

d. Histological measurement of tumour-free margins and statement
as to whether tumour is completely excised.

e. Presence or absence of vascular space invasion.

f. Nature of the adjacent non-malignant squamous epithelium

VIN
Lichen sclerosus
Squamous hyperplasia
HPV associated changes

g. Sites and number of nodes examined and number of positive
nodes. Presence or absence of extracapsular spread.

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Cancer of the vagina

2.1 Staging
A description of the staging classification for primary vaginal carcinoma is detailed in Table 1 and its stage grouping in Table 2.1

Table 1: Carcinoma of the Vagina: FIGO staging

<table>
<thead>
<tr>
<th>Stage</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>Carcinoma in situ; intraepithelial neoplasia grade 3</td>
</tr>
<tr>
<td>I</td>
<td>The carcinoma is limited to the vaginal wall</td>
</tr>
<tr>
<td>II</td>
<td>The carcinoma has involved the subvaginal tissue but has not extended to the pelvic wall</td>
</tr>
<tr>
<td>III</td>
<td>The carcinoma has extended to the pelvic wall</td>
</tr>
<tr>
<td>IV A</td>
<td>Tumour invades bladder an/or rectal mucosa and/or direct extension beyond the true pelvis</td>
</tr>
<tr>
<td>IVB</td>
<td>Spread to distant organs</td>
</tr>
</tbody>
</table>

2.1.1.1 Nodal stations
The upper two-thirds of the vagina is drained by lymphatics to the pelvic nodes, with the lymphatics paralleling the course of the uterine artery and the vaginal artery to the obturator, hypogastric and external iliac nodes. The lower third of the vagina drains to the inguinal-femoral nodes. Some lesions may drain via pararectal lymphatic channels.

2.1.1.2 Metastatic sites
The most common sites of distant spread include the lungs, liver, and bony skeleton. The rules for staging are similar to those for carcinoma of the cervix.

2.1.1.3 Histopathologic types
Squamous cell carcinoma is the most common type of cancer occurring in the vagina. Infrequently an adenocarcinoma may occur.

2.1.1.4 Histopathologic grades (G)
- Gx – Grade cannot be assessed;
- G1 – Well differentiated;
- G2 – Moderately differentiated;
- G3 – Poorly or undifferentiated.

2.1.1.5 Regional lymph nodes (N)
- NX – Regional lymph nodes cannot be assessed;
- NO – No regional lymph node metastasis;
2.2 Introduction
Carcinoma of the vagina constitutes only about 2% of malignant neo-
plasms of the female genital tract. The vagina, however, can be a
common site of metastatic disease by either direct extension of cervi-
cal or vulvar tumours or through metastatic lymphatic or vascular
deposits as seen in endometrial and gestational trophoblastic disease.

Up to 30% of patients with primary vaginal carcinoma have a his-
tory of in situ or invasive cervical cancer treated at least 5 years
earlier. Some vaginal cancers are preceded by vaginal intraepithe-
elial neoplasia (VAIN), although the true malignant potential of VAIN
is not known. Prior pelvic radiation has also been considered a
possible cause of vaginal cancer.

Most vaginal cancers occur in post-menopausal or elderly
women. When occurring in younger patients, the disease seems to
be etiologically related to cervical neoplasia, and thus human papillo-
mavirus (HPV) dependent. Histologically, approximately 95% of
primary vaginal cancers are squamous cell lesions. Metastatic
tumours in the vagina can also occur from non-gynaecological sites
such as the urinary bladder, urethra or periurethral glands and rarely
breast or lung cancer.

2.3 Screening
Routine screening for vaginal cancer following hysterectomy for
benign disease is not cost effective, but women with a history of cer-
vical intraepithelial or invasive neoplasia are at increased risk, and
should be followed with regular pap smears.

2.4 Vaginal intraepithelial neoplasia (VAIN)
For patients with an abnormal pap smear and no gross abnormality,
vaginal colposcopy and use of Lugol’s iodine to stain the vagina are
necessary. Excision of colposcopically abnormal areas is usually nec-
essary under anaesthesia. This is particularly true for lesions
involving the vaginal vault, where occult carcinoma may be found in
up to 28% of patients with VAIN.

Treatment of VAIN is individualised and depends on the extent,
location and general medical condition of the patient. Numerous
treatment methods, ranging from the various methods of local tissue
destruction or ablation through to more extensive surgery and includ-
2.5.1 Treatment
Patients should all be referred to tertiary referral units, because of the limited experience with these lesions. All treatment must be individualised, and will vary depending on the stage of disease and the site of vaginal involvement.

For most patients, it is important to try to maintain a functional vagina.

2.5.1.1 Surgery
Surgery has a limited role because of the close proximity of the bladder and rectum, but may be useful in the following situations (1,16,17).

1. In patients with Stage I disease involving the upper posterior vagina. If the uterus is still in situ, radical hysterectomy, upper vaginectomy to achieve clearance of at least 1 cm, and pelvic lymphadenectomy may be performed, while if hysterectomy has been previously performed, radical upper vaginectomy and pelvic lymphadenectomy may be appropriate.

2. In young patients who require radiation therapy. Pre-treatment laparotomy may allow ovarian transposition, surgical staging and resection of any bulky positive lymph nodes.

3. In patients with Stage IV A disease, particularly if a recto-vaginal or vesico vaginal fistula is present. Primary pelvic exenteration is a suitable treatment option for such patients, either combined with pelvic lymphadenectomy or pre-operative radiation.

4. In patients with a central recurrence after radiation therapy. Surgery will usually necessitate some type of pelvic exenteration in such patients.

Level of Evidence C.

2.5.1.2 Radiation Therapy
Radiation therapy is the treatment of choice for most patients with vaginal cancer, and comprises an integration of teletherapy and intracavitary/interstitial therapy.

Selected cases of Stage I and II lesions can be treated adequately with intracavity radiation alone (18,19). For larger lesions, treatment is
usually started with approximately 5000 cGy external radiation to shrink the primary tumour and treat the pelvic nodes. Intracavitary treatment follows. There is improved local control with total tumour doses of at least 7000cGy. If the lower one-third of the vagina is involved, the groin nodes should be treated or dissected. Level of Evidence C.

There is limited reported experience with chemo-radiation for vaginal cancer. However in view of the problem with control of the central disease, the concurrent use of 5-fluorouracil and/or Cisplatin may be appropriate. Level of Evidence D.

2.5.2 Prognosis
Recent reports have indicated 5-year survival rates comparable to cervical cancer. A recent study of 193 patients from the M.D. Anderson Cancer Center in Houston reported 5-year disease-specific survival rates of 85% for 50 patients with Stage I disease, 78% for 97 patients with Stage II, and 58% for 46 patients with Stages III-IVA.

2.6 Special situations
2.6.1 Adenocarcinoma
Approximately 9% of primary vaginal carcinomas are adenocarcinomas, and they affect a younger age group, regardless of whether or not exposure to diethylstilbestrol (DES) in utero has occurred. Adenocarcinomas may arise in areas of vaginal adenosis in DES exposed patients, in Wolffian rest elements, peri-urethral glands, and foci of endometriosis.

2.6.1.1 Screening
A young woman exposed to DES in utero should be initially seen when she begins to menstruate, or at approximately 14 years of age. The entire vagina and cervix should be inspected and palpated, and a pap smear taken.

2.6.1.2 Treatment
In general, adenocarcinomas are treated in a similar manner to squamous lesions. In young patients, every effort should be made to preserve vaginal and ovarian function. This may necessitate reconstruction of the vagina, or pre-radiation ovarian transportation.

2.6.1.3 Prognosis
Prognosis for clear cell carcinomas of the vagina is generally good, with an overall survival of 78%. Survival for non-clear cell adenocarcinomas is significantly worse than for squamous cancer.

2.6.2 Vaginal Melanoma
Malignant melanomas of the vagina are rare, and almost all cases occur in white women. Most commonly occur in the distal vagina, particularly on the anterior vaginal wall. Most are deeply invasive and radical surgery has been the mainstay of treatment, often involving some type of pelvic exenteration. Recently, more conservative local excisions have been used, with comparable survival rates reported. This is often combined with post-operative radiation. Radiation therapy alone may be effective in selected cases, and high-dose fractions (>400 cGy) may yield better response rates. Overall 5-year survival is about 10%. Level of Evidence C.

2.6.3 Sarcoma botryoides
Sarcoma botryoides is a highly malignant tumour of the rhabdomyoblasts. These neoplasms are found in infants and children and usually present with discharge, bleeding or a visible mass at the introitus.

In the past, exenterative surgery was used for these lesions, but survival was poor. More recently, conservative surgery has been used in conjunction with pre-operative or post-operative chemotherapy and radiotherapy with significantly improved survival. Most reported chemotherapeutic experience has been with Vincristine, Actinomycin D and Cyclophosphamide (VAC).

If the lesion is small and can be resected with organ preservation, surgery should be the initial approach. For bulkier lesions, pre-operative chemo or radiotherapy should be given. Extended radiotherapy fields are not recommended as they may produce significant developmental problems with the bony pelvis by destroying or interfering with growth centres in these structures.

References:


Cancer of the Cervix Uteri

3.1 Staging

3.1.1 Anatomy

3.1.1.1 Primary site. The cervix is the lower third of the uterus. It is roughly cylindrical in shape, projects through the upper, anterior vaginal wall and communicates with the vagina through an orifice called the external os. Cancer of the cervix may originate on the vaginal surface or in the canal.

3.1.1.2 Nodal stations. The cervix is drained by preureteral, pos-tureteral, and uterosacral routes into the following first station nodes: parametrial, internal (obturator - hypogastric), external iliac, pre-sacral, and common iliac. Para-aortic nodes are second station and are considered metastases.

3.1.1.3 Metastatic sites. The most common sites of distant spread include the aortic and mediastinal nodes, the lungs and skeleton.

3.1.2 Rules for classification

3.1.2.1 Clinical-diagnostic staging. Staging of cervical cancer is based on clinical evaluation; therefore, careful clinical examination should be performed in all cases, preferably by an experienced examiner and under anaesthesia. The clinical staging must not be changed because of subsequent findings. When there is doubt as to which stage a particular cancer should be allocated, the earlier stage is mandatory. The following examinations are permitted: palpation, inspection, colposcopy, endocervical curettage, hysteroscopy, cystoscopy, proctoscopy, intravenous urography, and X-ray examination of the lungs and skeleton. Suspected bladder or rectal involvement should be confirmed by biopsy and histologic evidence. Conization or amputation of the cervix is regarded as a clinical examination. Invasive cancers so identified are to be included in the reports. Findings of optional examinations, e.g. laparoscopy, ultrasound, CT scan, MRI, and PET scan are of value for planning therapy but, because these are not generally available and the interpretation of results is variable, the findings of such studies should not be the basis for changing the clinical staging. Fine needle aspiration (FNA) of scan-
Table 1: Carcinoma of the Cervix Uteri – Staging

<table>
<thead>
<tr>
<th>FIGO Stages</th>
<th>TNM Categories</th>
</tr>
</thead>
<tbody>
<tr>
<td>Primary tumour cannot be assessed</td>
<td>TX</td>
</tr>
<tr>
<td>No evidence of primary tumour</td>
<td>T0</td>
</tr>
<tr>
<td>0 Carcinoma in situ (preinvasive carcinoma)</td>
<td>Tis</td>
</tr>
<tr>
<td>I Cervical carcinoma confined to uterus (extension to corpus should be disregarded)</td>
<td>T1</td>
</tr>
<tr>
<td>IA Invasive carcinoma diagnosed only by microscopy. All macroscopically visible lesions – even with superficial invasion – are Stage IB/T1b</td>
<td>T1a</td>
</tr>
<tr>
<td>IA1 Stromal invasion no greater than 3.0 mm in depth and 7.0 mm or less in horizontal spread</td>
<td>T1a1</td>
</tr>
<tr>
<td>IA2 Stromal invasion more than 3.0 mm and not more than 5.0 mm with a horizontal spread 7.0 mm or less</td>
<td>T1a2</td>
</tr>
<tr>
<td>IB Clinically visible lesion confined to the cervix or microscopic lesion greater than IA2/T1a2</td>
<td>T1b</td>
</tr>
<tr>
<td>IB1 Clinically visible lesion 4.0 cm or less in greatest dimension</td>
<td>T1b1</td>
</tr>
<tr>
<td>IB2 Clinically visible lesion more than 4 cm in greatest dimension</td>
<td>T1b2</td>
</tr>
<tr>
<td>II Tumour invades beyond the uterus but not to pelvic wall or to lower third of the vagina</td>
<td>T2</td>
</tr>
<tr>
<td>IIA Without parametrial invasion</td>
<td>T2a</td>
</tr>
<tr>
<td>IIB With parametrial invasion</td>
<td>T2b</td>
</tr>
<tr>
<td>III Tumour extends to pelvic wall and/or involves lower third of vagina and/or causes hydrenephrosis or non-functioning kidney</td>
<td>T3</td>
</tr>
<tr>
<td>IIIA Tumour involves lower third of vagina no extension to pelvic wall</td>
<td>T3a</td>
</tr>
<tr>
<td>IIIB Tumour extends to pelvic wall and/or causes hydrenephrosis or non-functioning kidney</td>
<td>T3b</td>
</tr>
<tr>
<td>IVA Tumour invades mucosa of bladder or rectum and/or extends beyond true pelvis</td>
<td>T4</td>
</tr>
<tr>
<td>IVB Distant metastasis</td>
<td>M1</td>
</tr>
</tbody>
</table>

*Note: The depth of invasion should not be more than 5 mm taken from the base of the epithelium, either surface or glandular, from which it originates. The depth of invasion is defined as the measurement of the tumour from the epithelial-stromal junction of the adjacent most superficial epithelial papilla to the deepest point of invasion. Vascular space involvement, venous or lymphatic, does not affect classification.

**Note: The presence of bullous oedema is not sufficient to classify a tumour as T4.
smooth and indurated parametrium is truly cancerous, or only inflammatory. Therefore, the case should be placed in Stage III only if the parametrium is nodular to the pelvic wall or if the growth itself extends to the pelvic wall.

The presence of hydronephrosis or non functioning kidney resulting from stenosis of the ureter by cancer permits a case to be allotted to Stage III even if, according to other findings, the case should be allotted to Stage I or Stage II.

The presence of bullous oedema, as such, should not permit a case to be allotted to Stage IV. Ridges and furrows into the bladder wall should be interpreted as signs of submucous involvement of the bladder if they remain fixed to the growth at rectovaginal examination. Finding malignant cells in cytologic washings from the urinary bladder requires further histological confirmation in order to be considered for Stage IVA.

Table 2: Carcinoma of the Cervix Uteri – Stage Grouping

<table>
<thead>
<tr>
<th>FIGO Stage</th>
<th>UICC T</th>
<th>UICC N</th>
<th>UICC M</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>Tis</td>
<td>N0</td>
<td>M0</td>
</tr>
<tr>
<td>IA1</td>
<td>T1a1</td>
<td>N0</td>
<td>M0</td>
</tr>
<tr>
<td>IA2</td>
<td>T1a2</td>
<td>N0</td>
<td>M0</td>
</tr>
<tr>
<td>IB1</td>
<td>T1b1</td>
<td>N0</td>
<td>M0</td>
</tr>
<tr>
<td>IB2</td>
<td>T1b2</td>
<td>N0</td>
<td>M0</td>
</tr>
<tr>
<td>IIA</td>
<td>T2a</td>
<td>N0</td>
<td>M0</td>
</tr>
<tr>
<td>IIB</td>
<td>T2b</td>
<td>N0</td>
<td>M0</td>
</tr>
<tr>
<td>IIIA</td>
<td>T3a</td>
<td>N0</td>
<td>M0</td>
</tr>
<tr>
<td>IIIB</td>
<td>T1</td>
<td>N1</td>
<td>M0</td>
</tr>
<tr>
<td></td>
<td>T2</td>
<td>N1</td>
<td>M0</td>
</tr>
<tr>
<td></td>
<td>T3a</td>
<td>N1</td>
<td>M0</td>
</tr>
<tr>
<td></td>
<td>T3b</td>
<td>Any N</td>
<td>M0</td>
</tr>
<tr>
<td>IVA</td>
<td>T4</td>
<td>Any N</td>
<td>M0</td>
</tr>
<tr>
<td>IVB</td>
<td>Any T</td>
<td>Any N</td>
<td>M1</td>
</tr>
</tbody>
</table>

3.1.3.2 Regional Lymph Nodes (N)
- NX – Regional lymph nodes cannot be assessed
- N0 – No regional lymph node metastasis
- N1 – Regional lymph node metastasis

3.1.3.3 Distant Metastasis (M)
- MX – Distant metastasis cannot be assessed
- M0 – No distant metastasis
- M1– Distant metastasis

3.1.4 Histopathology
Cases should be classified as carcinomas of the cervix if the primary growth is in the cervix. All histologic types must be included. Grading by any of several methods is encouraged, but is not a basis for modifying the stage groupings. When surgery is the primary treatment, the histologic findings permit the case to have pathologic staging, as described above. In this situation, the TNM nomenclature may be used. All tumours are to be microscopically verified.

3.1.4.1 Histopathologic types
- Cervical intraepithelial neoplasia, Grade III
- Squamous cell carcinoma in situ
- Squamous cell carcinoma
  - Keratinizing
  - Non-keratinizing
  - Verrucous
- Adenocarcinoma in situ
- Adenocarcinoma in situ, endocervical type
- Endometrioid adenocarcinoma
- Clear cell adenocarcinoma
- Adenosquamous carcinoma
- Adenoid cystic carcinoma
- Small cell carcinoma
- Undifferentiated carcinoma

3.1.4.2 Histopathologic grades (G)
- Gx – Grade cannot be assessed;
- G1 – Well differentiated;
- G2 – Moderately differentiated;
- G3 – Poorly or undifferentiated.

3.2 Introduction
World-wide, cervical cancer is second only to breast cancer as the most common female malignancy in both incidence and mortality.
3.3 Cervical Screening
Data from many countries have shown that screening with cervical cytology reduces the incidence and mortality from cervical cancer.

Principles:
1. The purpose of a cervical screening programme is to reduce the incidence and mortality of cervical cancer.
2. Cervical screening should be population based with wide coverage (aim for at least 80% coverage of the population).
3. Cervical cytology is the most used method of screening.

3.3.1 Screening guidelines

3.3.1.1 Age group to be screened
This depends on the particular age distribution of deaths from cervical cancer and may be “country specific”. Deaths from cervical cancer are rare before age 25. Women can be discharged from the screening programme at the age of 65 if they have had two negative smears in the previous 10 years.

3.3.1.2 Frequency of screening
There is a higher incidence of women developing an interval cancer if the time from the previous smear is extended beyond 3 years.

3.3.1.3 Management of cervical cytology results
Recommendations for management after a cervical smear:
1. Routine recall: for a reportedly normal smear
2. Repeat smear:
   i. if smear is inadequate, repeat in three months.
   ii. for mild dyskaryosis or borderline nuclear changes, repeat in six – twelve months, depending on national protocol. The recommended repeat interval allows for possible resolution of changes. If the abnormality persists on repeat cytology, colposcopy is recommended.
   An alternative approach is to perform high risk HPV typing in patients with borderline cytology. Colposcopy should then be performed if high risk HPV types are found. If high risk HPV types are not found, cytology should be repeated in 12 months.
3. Refer for colposcopy: For moderate or severe dyskaryosis, query invasive disease or query glandular neoplasia.

More than 80% of new cases are diagnosed in economically disadvantaged people.
Carcinoma of the uterine cervix grows locally and may extend in continuity to the paracervical tissues and to the pelvic organs, spread to regional lymph nodes, and only later metastasize to distant structures.
Squamous carcinoma and adenocarcinoma are the most frequent histologic types.
3.4 Management of Cervical Cancer

3.4.1 Microinvasion

The diagnosis of Stage IA1 or IA2 disease can only be made on the basis of a cone biopsy with negative margins, or on a trachelectomy or hysterectomy specimen. If the margins of the cone biopsy are positive for CIN III or invasive cancer, a second cone biopsy should be performed or the patient treated as for Stage IB1 disease (1).

Colposcopy should be performed to exclude any associated vaginal intraepithelial neoplasia (VAIN) before undertaking definitive treatment.

3.4.1.1 Stage IA1

The recommended management is total abdominal or vaginal hysterectomy (2). If there is any associated VAIN, an appropriate cuff of vagina should be removed.

If fertility is desired, observation after cone biopsy is appropriate, with Pap smear follow-up at four months, 10 months, and then annually if both previous smears are negative. Level of Evidence B.

3.4.1.2 Stage IA2

There is a definite potential for lymph node metastasis in patients with Stage IA2 disease, so pelvic lymphadenectomy should be included in the treatment protocol (3,4).

The recommended treatment is modified radical hysterectomy (Type 2) and pelvic lymphadenectomy. If there is no lymph vascular space invasion, consideration may be given to extrafascial hysterectomy and pelvic lymphadenectomy. Level of Evidence C.

If fertility is desired, options are:

i. large cone biopsy plus extra-peritoneal or laparoscopic pelvic lymphadenectomy, or

ii. radical trachelectomy and extra peritoneal or laparoscopic pelvic lymphadenectomy (5).

3.4.1.3 Follow-up:

Should be mainly with Pap smears and should be annually after two normal smears at four and 10 months.

3.4.2 Invasive carcinoma

3.4.2.1 Initial Evaluation

For the patient with a visible lesion, biopsy should be done to confirm the diagnosis. Initial evaluation includes clinical examination (if necessary under anaesthesia), and colposcopy of the vagina to exclude VAIN. Relevant clinical symptoms should be investigated, and the bladder and rectum may be evaluated by cystoscopy and sigmoidoscopy if symptomatic. Chest x-ray and renal evaluation (which may consist of renal ultrasound, IVP, CT or MRI) are mandatory. CT and/or MRI and/or PET may provide some information on nodal status or systemic spread.

3.4.2.2 Stage IB1, IIA < 4 cm

Early stage cervical cancer (IB1, IIA < 4 cm) has a good prognosis and can be controlled with surgery or radiotherapy (6,7). Level of Evidence A.

The treatment of choice will depend on the availability of resources, the oncologist involved, and the age and the general health of the patient. It is desirable to have multidisciplinary consultation where possible and patients should be informed about the therapeutic alternatives, including their toxicity and the expected outcomes.

Morbidity is usually higher when both surgery and radiation are combined. In order to minimize morbidity, primary therapy should avoid the planned use of both radical surgery and radiation therapy. Level of Evidence A.

3.4.2.2.1 Surgery

The standard surgical treatment of stage IB1/IIA (< 4 cm diameter) is modified radical or radical abdominal hysterectomy (Class II and Class III according to Piver Rutledge classification) and pelvic lymphadenectomy.

In younger patients, the ovaries may be preserved and suspended outside the pelvis if post-operative radiation is likely to be given.

A vaginal radical hysterectomy and laparoscopic pelvic lymphadenectomy may be done in particular cases (8,9). Level of Evidence C.
3.4.2.2.2 Radiotherapy
The standard radiation treatment of stage IB1/IIA (≤ 4 cm diameter) is external pelvic irradiation plus brachytherapy. Suggested doses, including both external beam radiation and LDR brachytherapy, are 80-85 Gy to point A and 50-55 Gy to point B. The dose of external pelvic radiation should be 45 to 50 Gy in 180 to 200 cGy fractions. Using HDR brachytherapy, doses should be defined according to biological equivalence.

3.4.2.2.3 Adjuvant therapy post-surgery
The risk of recurrence after radical surgery is increased with the presence of positive nodes, positive parametria, or positive surgical margins. Adjuvant concurrent chemoradiation (using 5FU + Cisplatin or Cisplatin alone) improves survival compared with pelvic irradiation alone in such patients. Level of Evidence A.

Risk is also increased in those with uninvolved nodes but large tumour volume, capillary – like space (CLS) involvement, and outer one-third invasion of the cervical stroma. Adjuvant whole pelvic irradiation reduces the local failure rate and improves progression-free survival compared with patients treated with surgery alone. Radiation therapy appears to be particularly beneficial for patients with adenocarcinoma or adenosquamous histologies (20). Level of Evidence A.

Two groups have reported comparable tumour control with decreased morbidity using a small field of pelvic radiation, designed to cover the vaginal vault and parametrical tissues (12,13). The upper border on this field extends to about S1-2 rather than L5-S1. Level of Evidence C.

3.4.2.3 Stage IB2 – IIA (> 4 cm)
Options for primary therapy include:
1) Primary chemoradiation (14).
2) Primary radical hysterectomy and bilateral pelvic lymphadenectomy, which usually needs to be followed with adjuvant radiation.
3) Neoadjuvant chemotherapy (three rapidly delivered courses of platinum based chemotherapy) followed by radical hysterectomy and pelvic lymphadenectomy ± adjuvant post-operative radiation or chemoradiation (15).

3.4.2.3.1 Concurrent chemoradiation
The most commonly used treatment is external beam radiation plus intracavitary brachytherapy with concurrent weekly platinum chemotherapy. Suggested doses of radiation should be 85 to 90 Gy to point A and 55 to 60 Gy to point B. Cisplatin is given in a dose of 40 mg per m² weekly during external beam therapy. In patients with positive common iliac or paraaortic nodes, extended field radiation should be considered (16,17). Little data are yet available on the toxicity associated with concurrent chemotherapy and extended field irradiation. Level of Evidence A.

3.4.2.3.2 Primary surgery and probable adjuvant radiation
Primary radical hysterectomy offers the advantage of allowing proper surgical staging, while simultaneously removing the primary tumour, thereby obviating the need for brachytherapy (18). It also allows resection of any bulky positive lymph nodes, which are less likely to be sterilized with primary radiation (19).

Because these tumours are bulky by definition, adjuvant radiation is more likely to be necessary. Patients at particular risk of local recurrence are those with extensive CLS involvement and those with invasion to the outer third of the cervical stroma (20). The high-risk node-negative patients may be treated by whole pelvic (11) or small field pelvic radiation (12,13). Patients with positive common iliac or paraaortic nodes may be treated by extended field radiation (16,17) with or without chemotherapy. Level of Evidence C.

3.4.2.3.3 Neoadjuvant chemotherapy followed by radical hysterectomy and pelvic lymphadenectomy
Data from randomised trials would suggest that neoadjuvant platinum-based chemotherapy prior to definitive surgery is associated with better results than primary radiation (15,21). There are no data available to compare the results of concurrent chemoradiation with those of neoadjuvant chemotherapy followed by surgery. Level of Evidence B.

The chemotherapy used in the Buenos Aires study was as follows (15):
Cisplatinum 50 mg/m² IV in 15 minutes on day 1
Vincristine 1 mg/m² IV push on day 1, and
3.4.2.4 Advanced cervical cancer

3.4.2.4.1 Definition
Includes stage IIB, III and IV A.

3.4.2.4.2 Primary treatment
Standard primary treatment is irradiation given as a combination of external radiation and intracavitary brachytherapy with concurrent chemotherapy (14,22). Level of Evidence A.

Primary pelvic exenteration may be considered for Stage IV A disease not extending to the pelvic sidewall, particularly if a vesicovaginal or rectovaginal fistula is present. Level of Evidence C.

3.4.2.4.3 Irradiation dose and technique
Doses and field technique are given in Table 3. Irradiation should be given by an appropriate energy causing a uniform dose distribution (± 5%) within the primary and secondary target volume. The target volume should be determined by clinical examination and CT-scanning when possible. The field technique should consist of at least four fields. Brachytherapy may be given as high or low dose rate. The standard treatment is external beam radiation plus intracavitary brachytherapy concurrent with platinum-based chemotherapy. Cisplatin is given weekly during the external beam therapy at a dose of 40 mg/m². Suggested doses of radiation should be 85 to 90 Gy to point A and 55 to 60 Gy to point B. In patients with positive common iliac or paraaortic nodes extended field radiation should be considered (16,17,23). Level of Evidence C.

3.4.2.5 Stage IVB or recurrent disease

3.4.2.5.1 Background
Recurrence may be pelvic, distant or both. As the bulk of the primary pelvic tumour increases, the proportion of patients with disease recurrent or persistent in the pelvis as the only site of failure increases compared with the proportion developing distant metastases.

The majority of recurrences occur within two years of diagnosis and the prognosis is poor with most patients dying as a result of uncontrolled disease (24). The median duration of survival is seven months.

Symptoms of recurrent/metastatic cervical cancer may include pain.

**Table 3: Management of Advanced Cervical Cancer**

<table>
<thead>
<tr>
<th>Stages:</th>
<th>Stage IIB – IVA</th>
</tr>
</thead>
</table>
| Staging: | Examination under general anaesthesia  
Chest x-ray  
Renal imaging  
Optional CT/MRI scan of abdomen and pelvis, PET scan |
| Radiation technique: |  
A. Primary target  
Tumour + uterus  
B Secondary target  
Pelvic lymph nodes and common iliac lymph nodes  
Field technique: 4 fields  
Field borders for external irradiation  
A. Tumour determined by palpation and CT scan (if available) + 2 cm margin  
B. A-P fields:  
Lateral: 2 cm lateral to the bony margin of the pelvis  
Superior: Between L5 and S1  
Inferior: 2 cm below the obturator foramen (or 2 cm below lower extent of clinical tumour)  
C. Lateral fields  
Anterior = individually determined by tumour  
Posterior = individually determined by tumour:  
Primary target dose: External irradiation 50 Gy/5-6 weeks + LDR intracavitary boost 30-35 Gy point A.  
(for IIB - IVA, 35-40 Gy)  
Secondary target: External irradiation 50 Gy/5 Weeks  
Total treatment time: 6-7 weeks  
Concurrent chemotherapy: Cisplatin 40 mg/m² every week during external irradiation |

Bleomycin 25 mg/m² by continuous infusion over six hours on days 1-3. The regime is repeated at 10-day intervals for three cycles.
leg swelling, anorexia, vaginal bleeding, cachexia and psychological problems among others.

The coordinated efforts of a team of professionals are optimal; the team may include gynaecologic oncologists, radiation and medical oncologists, palliative care physicians, specialised nursing staff, psychologists, and possibly stomal therapists. Relief of pain and other symptoms, along with comprehensive support for the patient and her family, are paramount.

### 3.4.2.5.2 Management of patients who relapse after primary treatment

Treatment decisions should be based on the performance status of the patient, the site of recurrence and/or metastases, the extent of metastatic disease and the prior treatment.

#### 3.4.2.5.2.1 Locally recurrent cervical cancer following radical surgery

<table>
<thead>
<tr>
<th>Guidelines – Locally Recurrent Cervical Cancer Following Surgery</th>
<th>Level of Evidence</th>
</tr>
</thead>
<tbody>
<tr>
<td>Radiation therapy is indicated in patients with locally recurrent cervical cancer following radical surgery</td>
<td>C</td>
</tr>
<tr>
<td>Concurrent chemotherapy with either 5-Fluorouracil and/or Cisplatin with radiation should be considered and may improve outcome</td>
<td>B</td>
</tr>
<tr>
<td>Pelvic exenteration may be an alternative (particularly if a fistula is present) to radical radiotherapy and concurrent chemotherapy in selected patients without pelvic sidewall involvement.</td>
<td>C</td>
</tr>
</tbody>
</table>

#### 3.4.2.5.2.2 Therapeutic options for local relapse after primary surgery

Relapse in the pelvis following primary surgery may be treated by either radical radiation or pelvic exenteration. Radical irradiation (± concurrent chemotherapy) may cure a substantial proportion of those with isolated pelvic failure after primary surgery.

Radiation dose and volume should be tailored to the extent of disease. Fifty Gray in 180 cGy fractions should be delivered to microscopic disease and using field reductions 64 to 66 Gy should be delivered to the gross tumour volume.

Where disease is metastatic or recurrent in the pelvis after failure of primary therapy and not curable, a trial of chemotherapy with palliative intent or symptomatic care is indicated. Cisplatin is the single most active agent for the treatment of cervical cancer.

The expected median time to progression or death is three to seven months.

#### 3.4.2.5.2.3 Local recurrence following definitive (radical) radiation

<table>
<thead>
<tr>
<th>Guideline – Local Recurrence Following Prior Radiotherapy</th>
<th>Level of Evidence</th>
</tr>
</thead>
<tbody>
<tr>
<td>Selected patients with small disease (≤ 2 cm) confined to the cervix may be suitable for radical hysterectomy</td>
<td>C</td>
</tr>
<tr>
<td>Patients with a central recurrence and no evidence of metastatic disease should be considered for pelvic exenteration</td>
<td>C</td>
</tr>
</tbody>
</table>

Radical hysterectomy may be used for patients with small disease (≤ 2 cm in diameter) confined to the cervix. The morbidity is high, but some patients can be cured without need for a stoma.

Patients with a central recurrence that involves the bladder and/or rectum, without evidence of intraperitoneal or extra pelvic spread, and who have a tumour-free space along the pelvic sidewall, are potentially suitable for pelvic exenteration. The triad of unilateral leg oedema, sciatic pain and ureteral obstruction almost always indicates unresectable disease on the pelvic sidewall, and palliative measures are indicated.

The prognosis is better for patients with a disease-free interval greater than six months, a recurrence 3 cm or less in diameter, and no sidewall fixation. The five-year survival for patients selected for treatment with pelvic exenteration is in the order of 30 - 60% and the operative mortality should be < 10%.
for stages IA2 and beyond, further treatment should be as follows:

(i) If margins are positive, or if there is deep stromal infiltration and vascular space invasion, pelvic radiation with or without concurrent chemotherapy should be given.

(ii) For patients without these high risk features, radical para-

metrectomy, upper vaginectomy and pelvic lymph node dissection is an alternative option.

Level of evidence C

3.5.2 Cervical Cancer during Pregnancy

In general, the management of cervical cancer in pregnant women follows the same principles as in non-pregnant women. There are a few special considerations. Cone biopsy should be used only if cytology and histology suggest possible invasion, because of the problems of haemorrhage, abortion or premature labour. A multidisciplinary approach with involvement of obstetrician and neonatologist is essential. All management plans should be decided only after full discussion with the woman and preferably her partner, and their wishes must be respected.

For patients with suspected microinvasive carcinoma of the cervix, a therapeutic delay does not appear to be detrimental to maternal outcome and can result in significant foetal salvage. For Stage IA1 disease confirmed by cervical conization with clear surgical margins may be followed until term, and undergo vaginal delivery. There is no evidence that the route of delivery influences the outcome for women with microinvasive carcinoma.

For stage IA1 disease, no further treatment is required. For patients with advanced disease, management should be individualised and is influenced by the stage of disease and the duration of the pregnancy. MRI scan should be obtained to provide information on the extent of disease.

(2) If the diagnosis is made before 20 weeks, the disease should generally be treated without delay. For appropriate patients, radical hysterectomy and pelvic lymphadenectomy may be performed with the foetus insitu. For patients having treatment with chemoradiation, spontaneous abortion will usually occur.

For those diagnosed after 28 weeks, the recommendation should be
to await foetal viability. If the diagnosis is made between 20 and 28 weeks, treatment delay appears to be an option at least for Stages IA2 and IB1, with no significant impairment of prognosis (39, 40). For more advanced disease, it is not known whether treatment delay will affect survival. In addition, there is no standard definition on what constitutes significant treatment delay.

In practice, the duration of the treatment delay should be influenced by clinical stage and histopathologic findings of the tumour, gestational age at diagnosis, and the parents’ desire regarding their unborn child. If treatment delay is planned in women with locally advanced disease, neoadjuvant chemotherapy should be considered in an attempt to prevent disease progression (40), and close clinical surveillance is mandatory. Delivery should be performed not later than 34 weeks of gestation.

Unless the lesion has been removed by conization, the recommended mode of delivery is classical caesarean section, although several retrospective studies have demonstrated no evidence that vaginal delivery adversely affects prognosis (42).

### Appendix 1 FIGO Staging

<table>
<thead>
<tr>
<th>Stage</th>
<th>Characteristics</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>Carcinoma in situ, intraepithelial carcinoma; cases of stage 0 should not be included in any therapeutic statistics for invasive carcinoma</td>
</tr>
<tr>
<td>I</td>
<td>The carcinoma is strictly confined to the cervix (extension to the corpus should be disregarded)</td>
</tr>
<tr>
<td>IA</td>
<td>Invasive cancer identified only microscopically. All gross lesions, even with superficial invasion, are stage IB cancers. Invasion is limited to measured stromal invasion with a maximum depth of 5 mm taken from the base of the epithelium, either surface or glandular, from which it originates. Vascular space involvement, either venous or lymphatic, should not alter the staging</td>
</tr>
<tr>
<td>IA1</td>
<td>Measured invasion of stroma no greater than 3 mm in depth and no wider than 7 mm</td>
</tr>
<tr>
<td>IA2</td>
<td>Measured invasion of stroma &gt; 3 mm and no greater than 5 mm in depth and no wider than 7 mm</td>
</tr>
<tr>
<td>IB</td>
<td>Clinical lesions confined to the cervix or preclinical lesions &gt; IA</td>
</tr>
<tr>
<td>IB1</td>
<td>Clinical lesions no greater than 4 cm in size</td>
</tr>
<tr>
<td>IB2</td>
<td>Clinical lesions &gt; 4 cm in size</td>
</tr>
<tr>
<td>II</td>
<td>The carcinoma extends beyond the cervix, but has not extended onto the pelvic wall; the carcinoma involves the vagina, but not as far as the lower third</td>
</tr>
<tr>
<td>II1</td>
<td>No extension onto the pelvic wall, but involvement of the lower third of the vagina</td>
</tr>
<tr>
<td>II2</td>
<td>Extension onto the pelvic wall or hydronephrosis or nonfunctioning kidney</td>
</tr>
<tr>
<td>III</td>
<td>The carcinoma has extended beyond the true pelvis or has clinically involved the mucosa of the bladder or rectum</td>
</tr>
<tr>
<td>IIIA</td>
<td>No extension onto the pelvic wall, but involvement of the lower third of the vagina</td>
</tr>
<tr>
<td>IIIB</td>
<td>Extension onto the pelvic wall or hydronephrosis or nonfunctioning kidney</td>
</tr>
<tr>
<td>IVA</td>
<td>Spread of the growth to adjacent organs</td>
</tr>
<tr>
<td>IVB</td>
<td>Spread to distant organs</td>
</tr>
</tbody>
</table>

(1) FIGO staging is based on clinical data (clinical examination and colposcopy), chest x-rays, IVP, biopsy and D&C.
(2) Cystoscopy and sigmoidoscopy may be used for clinical stage (bladder and/or rectal mucosal biopsy).
(3) Lymphangiogram, CT, MRI, PET, laparotomy, laparoscopy cannot be used for clinical staging.
(4) Pathological IVP defines a cancer as stage IIIB.
(5) Paracervical, parametrical, hipogastric, obturator, internal, external and common iliac, presacral and sacral are the regional nodes.
Figure 1: Stage IBI

Work-up
- Biopsy
- Clinical staging
- CT of pelvis and abdomen
- MR or PET optional

Therapy
- Radical hysterectomy (class II-III) with pelvic LND

Pathology
- No risk factors
- Negative nodes, LVSI
- Outer 1/3 invasion
- Pelvic RT with Cisplatin
- Positive nodes
- Positive parametrium
- Positive margins
- Pelvic RT with Cisplatin

Follow-up
- Negative nodes, LVSI
- Outer 1/3 invasion

Primary Radiation
- External beam pelvic RT 45 Gy/4-5 weeks
- Intracavitary brachytherapy
  - LDR 35-40 Gy
  - HDR 7 Gy/w x 4

External beam pelvic RT 45 Gy/4-5 weeks + intracavitary LDR boost 35-40 Gy point A or HDR equivalent biological dose

Concurrent Chemoradiation
- Concurrent chemotherapy:
  - Cisplatin 40 mg/m² q week during external irradiation
- Total treatment time: 6-7 weeks

Follow-up
- CR
- PR
- Adjuvant RT ± concurrent chemotherapy
- Progression
- Palliative pelvic RT ± concurrent chemotherapy

*The role of neoadjuvant CT followed by class II-III radical hysterectomy and pelvic lymphadenectomy (+ adjuvant concurrent CT/RT) remains to be further defined.*
References:


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Cancer of the Corpus Uteri

4.1 Staging

4.1.1 Anatomy

4.1.1.1 Primary site. The upper two-thirds of the uterus above the level of the internal cervical os is called the corpus. The Fallopian tubes enter at the upper lateral corners of a pear-shaped body. The portion of the muscular organ that is above a line joining the tubouterine orifices is often referred to as the fundus.

4.1.1.2 Nodal stations. The major lymphatic trunks are the utero-ovarian (infundibulo-pelvic), parametrial, and presacral, which drain into the hypogastric, external iliac, common iliac, presacral, and para-aortic nodes.

4.1.1.3 Metastatic sites. The vagina and lungs are the common metastatic sites.

4.1.2 Rules for classification

The FIGO Committee on Gynecologic Oncology, following its meeting in 1988, recommended that endometrial cancer be surgically staged. There should be histologic verification of grading and extent of the tumour.

4.1.3 Notes about the staging

4.1.3.1 Histopathology – degree of differentiation.

Cases of carcinoma of the corpus should be grouped with regard to the degree of differentiation of the adenocarcinoma as follows:

- **G1**: \( \leq 5\% \) of a nonsquamous or nonmorular solid growth pattern.
- **G2**: 6-50\% of a nonsquamous or nonmorular solid growth pattern.
- **G3**: > 50\% of a nonsquamous or nonmorular solid growth pattern.
4.1.3.4 Notes on pathologic grading

- Notable nuclear atypia, inappropriate for the architectural grade, raises the grade of a Grade 1 or Grade 2 tumour by 1.
- In serous and clear cell adenocarcinomas, nuclear grading takes precedence.
- Adenocarcinomas with squamous differentiation are graded according to the nuclear grade of the glandular component.

4.1.3.5 Rules related to staging

- Corpus cancer is now surgically staged, therefore procedures previously used for determination of stages are no longer applicable (e.g. the findings of fractional curettage to differentiate between Stage I and Stage II).
- It is appreciated that there may be a small number of patients with corpus cancer who will be treated primarily with radiation therapy. In these cases, the clinical staging adopted by FIGO in 1971 would still apply, but designation of that staging system would be noted.
- Ideally, width of the myometrium should be measured along with the depth of tumour invasion.
- As a minimum, any enlarged or suspicious lymph nodes should be
removed in all patients. For high risk patients (Grade III, deep myometrial invasion, cervical extension, serous or clear cell histology), complete pelvic lymphadenectomy and resection of any enlarged para-aortic nodes is recommended.

4.1.4 Histopathology (according to WHO/ISGP classification)
All tumours are to be microscopically verified.
The histopathologic types are:
• Endometrioid carcinoma
  - Adenocarcinoma
  - Adenoacanthoma (adenocarcinoma with squamous metaplasia)
  - Adenosquamous carcinoma (mixed adenocarcinoma and squamous cell carcinoma)
• Mucinous adenocarcinoma
• Papillary serous adenocarcinoma
• Clear cell adenocarcinoma
• Undifferentiated carcinoma
• Mixed carcinoma

4.1.4.1 Histopathologic grades (G)
• Gx – Grade cannot be assessed;
• G1 – Well differentiated;
• G2 – Moderately differentiated;
• G3 – Poorly or undifferentiated.

4.2 Introduction
In developed countries, where deaths from cervical cancer have been reduced by up to 50% because of screening, endometrial cancer ranks alongside ovarian cancer as the leading types of gynaecological cancer. The incidence of endometrial cancer rises from 2 per 100,000 women per year under the age of 40 years to 40-50 per 100,000 women per year in the 6th, 7th and 8th decades. Deaths from endometrial cancer in the United States doubled between 1988 and 1998, probably due to a combination of increased life expectancy and an epidemic of obesity, which predisposes to this disease.

The aetiology of endometrial cancer is unclear, although endometrioid carcinoma is thought to progress through a premalignant phase of intraendometrial neoplasia in a large proportion of cases. Other forms such as papillary serous and clear cell carcinoma probably arise as a result of a sequence of poorly understood genetic mutations; we know, for example, that mutant p53 commonly stains positive in papillary serous carcinoma.

Until recently there was very little research on which to base clinical practice guidelines, but the last 10 years have seen greater interest in endometrial cancer clinical trials. Its early presentation following post-menopausal bleeding results in a generally good prognosis, but
it should be treated by tightly governed protocols, and where appropriate, by expert multidisciplinary teams.

There is no means of effective screening available for endometrial cancer, although certain high risk groups such as those with Lynch type 2 syndrome can undergo endometrial surveillance by hysteroscopy and biopsy, or transvaginal ultrasonography if post-menopausal. Following presentation, ultrasound is an effective first test with a high negative predictive value when the endometrial thickness is less than 5 mm. In one of the largest studies undertaken, there was a negative predictive value of 96% amongst 1168 women in whom the results of transvaginal ultrasound were correlated with an endometrial biopsy obtained by curettage. When a biopsy is required this can be obtained usually as an office procedure using a number of disposable instruments developed for the purpose. In certain cases hysteroscopy may be helpful, and with flexible instruments can also be done without recourse to general anaesthesia. If cervical stenosis or patient tolerance does not permit an office procedure, then curettage under anaesthesia is necessary. Individuals whose pelvic examination is unsatisfactory may also be evaluated with transvaginal or abdominal ultrasound to rule out concomitant adnexal pathology.

Following a histopathologic diagnosis of endometrial carcinoma, the local extent of the tumour, and the risk for metastatic disease should be determined, and the perioperative risk assessed.

As a minimum, the pathology report should indicate both the tumour type and the degree of differentiation. A chest X-ray, full biochemistry and blood count are routine. A serum CA-125 may be of value in advanced disease for follow-up. Evaluation for metastasis is indicated particularly in patients with abnormal liver function tests, and clinical findings such as parametrial or vaginal tumor extension. In certain situations, cystoscopy and/or proctoscopy may be helpful, if direct extension to bladder or rectum is suspected.

4.3 Prognostic tumour characteristics for high risk
The recommended histopathological criteria for determining poorer prognosis are as follows:
- Tumour grade 3 (poorly differentiated)
- Deep myometrial invasion (FIGO stage 1C)
- Lymphvascular channel involvement
- Positive peritoneal cytology
- Serous papillary tumour
- Clear cell tumors
- Cervical involvement (stage II)

The most accurate means of assessing both depth of myometrial invasion and cervical involvement is MR scanning. CT and MR are equivalent in terms of evaluating nodal metastases, but neither is good enough to replace surgical lymph node assessment.

Non-surgical staging for endometrial cancer, where extraterine disease exists, is inherently inaccurate, particularly in respect of small nodal involvement, intraperitoneal implants, and adnexal metastasis. Furthermore, it is widely accepted that a curettage specimen may under-classify the tumour when the hysterectomy specimen has been fully examined. Up to 20% of tumours may have a worse histologic grade and occasionally different tumour type based on the hysterectomy specimen.

4.4 Surgical staging procedure for endometrial cancer
In 1988, the FIGO system changed from clinical to surgical staging for endometrial cancer. Since that recommendation, considerable debate has ensued as to what constitutes an internationally acceptable approach. A generally recommended protocol would be that the abdomen should be opened with a vertical midline abdominal incision and peritoneal washings taken immediately from the pelvis and abdomen, followed by careful exploration of the intra-abdominal contents. The omentum, liver, peritoneal cul-de-sac and adnexal surfaces should be examined and palpated for any possible metastases, followed by careful palpation for suspicious or enlarged nodes in the aortic and pelvic areas. The standard surgical procedure should be an extraperitoneal total hysterectomy with bilateral salpingo-oophorectomy. Adnexal removal is also recommended even if the adnexae appear normal, as they may contain micrometastases. Vaginal cuff removal is not necessary nor is there any benefit from excising parametrial tissue in the usual case. If cervical stromal involvement is demonstrated pre-operatively or if unsuspected cervical involvement is noted and can be encompassed by a modified radical hysterectomy, then this may be the most appropriate action in experienced hands.
surgical staging is to be undertaken.

4.7 Adjuvant Radiotherapy
Historically, two basic approaches have been adopted regarding the use of radiotherapy in the initial management of endometrial carcinoma. The earlier approach was to administer preoperative irradiation followed by surgery. More recently, findings at laparotomy are used to determine the need for radiotherapy in a purely adjuvant setting following definitive surgery.

In Europe it has been common practice to base the need for adjuvant radiotherapy on risk determined by grade of tumour and myometrial invasion. In North America and Australia the decision is generally based more on whether surgical staging has excluded extrauterine disease and hence the risk of recurrence. The argument is that rationalising the use of radiotherapy in this way reduces morbidity and maintains survival rates. Several recent studies have reported excellent results for patients with stage I endometrial cancer, in whom external radiation was avoided in node negative women (14-16).

Low risk disease does not require adjuvant radiotherapy as demonstrated in a Danish cohort study of low risk women, with a 96% 5-year survival (17). A seminal Norwegian trial (18) indicated 20 years ago that overall survival was not improved by adjuvant external beam therapy, although it did reduce the risk of pelvic recurrence. The trial included 621 women with all categories of FIGO stage I disease and all women had vaginal brachytherapy. The failure to improve overall survival was due to a higher risk of distal metastases in women who had brachytherapy.

The PORTEC trial from the Netherlands reported 715 women with either grade 1 (outer 1/2), all grade 2 or grade 3 (inner 1/2) disease, who were randomised after surgery (without lymphadenectomy) to pelvic lymphadenectomy or no lymphadenectomy, showed no therapeutic benefit (19).

Laparoscopically assisted vaginal hysterectomy is permitted, for surgeons experienced in this technique in low grade disease, but would need to be converted to an open procedure if unexpected metastases are identified. It can be accompanied by a laparoscopic lymphadenectomy, if surgical staging is to be undertaken.

4.5 Who should perform the surgery?
Low risk tumours will have positive nodes in less than 5% cases (well differentiated and <1/2 myometrial invasion) and do not require full surgical staging. These women can generally be safely operated on by a general gynaecologist, but those at greater risk of extrauterine disease who require lymphadenectomy should be referred to a specialist gynaecological oncologist. This triaging of women can be done most effectively by a thorough pre-operative assessment, paying particular attention to the pathology and to radiological features.

4.6 Is lymphadenectomy therapeutic?
Although required for accurate staging, a therapeutic benefit for lymphadenectomy is controversial. One case control study suggested that it may be therapeutic (20) and another showed a good prognosis even in node positive women (21). In the UK, the MRC ASTEC trial, which randomised women undergoing surgery for presumed stage I endometrial cancer to pelvic lymphadenectomy or no lymphadenectomy, showed no therapeutic benefit (22).

Laparoscopically assisted vaginal hysterectomy is permitted, for
Algorithm for Histopathologically confirmed Endometrial Cancer

- CXR
- MR Scan

High risk
- G3 and/or
- >1/2 myometrial invasion and/or
- suspicion of nodal mets on MR/CT scan
- papillary serous/clear cell type

Refer to gynaecological oncologist

Low risk
- G1
- G2 < 1/2 myometrial invasion

Can be managed by generalist

Staging laparotomy - (pelvic +/- para aortic lymphadenectomy*)
Peritoneal washing
Total abdominal hysterectomy/BSO

If nodes positive
- prescribe radiotherapy

If nodes negative and serosa intact
- adjuvant therapy is probably not required*, although many would give vault brachytherapy

Low vertical incision
Total abdominal hysterectomy/BSO
Peritoneal washing
Palpation of lymph nodes plus sampling if suspicious

If higher risk features are found on histopathology
- consider adjuvant radiotherapy in conjunction with clinical oncologist

*Randomised trials are needed to provide conclusive evidence regarding the clinical effectiveness of lymphadenectomy and adjuvant radiotherapy in the presence of high risk factors.

the control group. Ten year locoregional relapse rates were 5% after radiation and 14% in controls (P < 0.0001), and 10-year overall survival was 66% and 73%, respectively (P = 0.09). Endometrial cancer death rates were 11% (RT) and 9% (controls) (P = 0.47) (20).

The published data suggest that adjuvant radiotherapy is not indicated in the presence of low or intermediate risk stage I disease. This would certainly include a) all G1 tumours without serosal involvement, and G2 < 50% myometrial invasion. In higher risk women in whom full surgical staging has excluded extraterine disease, radiotherapy is of uncertain benefit and many would reserve external beam radiotherapy in these women for pelvic recurrence. Others would advocate adjuvant radiotherapy for the very high risk cases, such as G3 tumors with > 50% myometrial invasion. Many would give vaginal brachytherapy alone for high risk patients with negative pelvic nodes.

4.8 Progestogen therapy
This has been widely prescribed in the past, but a meta-analysis of 6 randomised trials involving a total of 3,339 women has shown no survival benefit for adjuvant progestogen therapy in endometrial cancer (21). A subsequently published randomised trial of 1012 women also failed to demonstrate any survival benefit (22).

4.9 Stage II
Patients with clinically occult Stage II disease are generally managed in a similar fashion as patients with Stage I disease.

Surgery can be used as a primary treatment for clinically overt cervical involvement; a radical hysterectomy with bilateral pelvic lymphadenectomy and selective aortic node dissection should be performed. If this approach is to be used, pre-operative MR scanning is advisable to ensure local resectability without bladder involvement. Recent studies indicate excellent results for this approach, with no benefit from the addition of radiation for patients with negative nodes (23-25).

If surgery is not considered feasible initially, full pelvic radiotherapy and intracavitary brachytherapy, followed by adjunctive hysterectomy, with selective lymphadenectomy of the aortic pelvic nodes, may be employed.
4.10 Stage III
Patients with Stage III endometrial carcinoma, by virtue of vaginal or parametrial extension, are best treated by pelvic irradiation after thorough metastatic work-up. Once therapy is complete, exploratory laparotomy is advised on those patients whose disease seems to be resectable. Extended field radiation therapy or systemic therapy with cytotoxic medications or hormones is warranted in the presence of extrapelvic metastases, depending on the patient’s condition. If an individual is in a clinical Stage III category, because of adnexal mass or involvement, as noted on ultrasound, these individuals should undergo surgery without preoperative radiotherapy, to determine the nature of the mass and to perform surgical pathological staging. In many instances cytoreductive surgery can be carried out and if the uterus is resectable then hysterectomy and adnexectomy should be performed. In some instances, rather than metastases to the ovary, one may find that the patient has a primary lesion of both the endometrium and also of the ovary after histologic examination of the removed tissue.

4.11 Stage IV
Patients with evidence of extrapelvic metastases are usually managed with systemic chemotherapy or hormonal therapy. However, the GOG recently reported a randomized trial to compare whole abdominal radiation with doxorubicin-cisplatin (AP) chemotherapy in women with Stage III of IV endometrial cancer having a maximum of 2 cm of postoperative residual disease. Chemotherapy significantly improved progression-free and overall survival compared with WAI. At 60 months, 55% of AP patients were predicted to be alive compared with 42% of WAI patients.

Local irradiation may be beneficial, particularly in brain or bone metastases, and occasionally pelvic radiotherapy may help in providing local tumor control and preventing bleeding or complications from local disease.

4.12 Special considerations

4.12.1 Diagnosis post-hysterectomy
Diagnosis of endometrial carcinoma post-hysterectomy can present some difficult management problems, particularly if the adnexae have not been removed. This situation most often arises following vaginal hysterectomy for pelvic prolapse. Recommendations for further postoperative therapy are based on known risk factors for extraterine disease related to the histologic grade and depth of myometrial involvement. Individuals with grade 3 lesions, deep myometrial invasion, or LVS invasion are candidates for additional surgery to remove the adnexa and to complete surgical staging. Alternatively, the empiric use of external beam radiation to the pelvis may be used. Patients with a grade 1 or 2 lesion with minimal myometrial invasion and no LVS involvement generally require no further therapy.

4.12.2 Medically inoperable patient
Morbid obesity and severe cardiopulmonary disease are the general reasons a patient with endometrial carcinoma is usually thought to be medically inoperable. Uterine brachytherapy can achieve cure rates in excess of 70% and may be combined with external beam radiotherapy in the presence of prognostic factors suggesting a high risk of involved nodes.

For patients with a well-differentiated lesion and contraindications to general anaesthesia and unsuited for radiotherapy, high-dose progestins may be used.

4.12.3 Diagnosis in the young woman
The diagnosis of endometrial carcinoma during the reproductive years should be made with caution, since this malignancy is uncommon in women < 35 years and Grade I endometrial carcinoma may be confused with severe atypical hyperplasia in these situations. In these women consideration should be given to an oestrogen related underlying condition such as granulosa cell tumour, polycystic ovaries or obesity. Atypical hyperplasia can be treated successfully with progestins, and the decision to use progestins in these situations may be appropriate if preservation of fertility is desirable. Equivocal lesions should be examined by an experienced pathologist. If carcinoma is confirmed, hysterectomy with adnexal removal remains the treatment of choice. When uncertainty remains regarding the presence of true carcinoma, the ultimate decision rests with the patient and the patient...
should undergo thorough counselling and documentation if a conservative approach is followed. A recent paper reported that 4 of 12 patients aged 40 years and younger with grade I tumors had complete resolution of disease using medroxyprogesterone acetate 600 mg per day. Two of the 4 went on to have children (27).

4.12.4 Positive peritoneal cytology
The presence of positive peritoneal cytology, which is often a difficult diagnosis, given the malignant appearance of reactive mesothelial cells, should only be made after a thorough cytopathology review. Management in this situation is controversial if no other features of extraperitoneal disease have been documented at the time of surgical staging, because insufficient data exist regarding recurrence risk and treatment results.

4.13 Follow-up
The conventional reasons for follow-up of treated cancer patients are accepted and involve providing reassurance, diagnosing early recurrence and collecting data. A range of protocols for following up women with endometrial cancer exists but the evidence base in endometrial cancer does not provide much support for follow-up in terms of improving survival. One prospective (28) and several retrospective studies (29-32) internationally have addressed follow-up. Overall, very few recurrences were identified as a direct result of clinic review and neither recurrence-free nor overall survival were improved in these cases compared with those detected at clinical presentation. A Canadian study (33) concluded that the use of routine follow-up Pap smears and chest x-rays was probably not cost effective. In non-irradiated patients, a strong case can be made for regular follow-up to detect vaginal recurrence at the earliest opportunity, given the high salvage rate following radiotherapy (33).

4.14 Recurrence
Localised recurrences are managed preferentially by surgery, irradiation, or a combination of the two, depending on the primary therapy. Large lesions should be excised, if feasible, with isolated pelvic recurrence of any grade being potentially curable, particularly if it occurs more than 1 or 2 years after initial therapy. In this setting, extended or radical surgery may be justified if the patient has already received prior irradiation. The results of pelvic exenteration in properly selected cases of this sort are similar to those obtained in cervical cancer.

Patients with non-localised recurrent tumours may be candidates for progestin therapy (medroxyprogesterone acetate 50-100 mg tid or megestrol acetate 80 mg two to three times per day). The progestin therapy is continued as long as the disease is static or in remission. Maximum clinical response may not be apparent for 3 or more months after initiating therapy. Chemotherapy with cisplatinum, taxol and adriamycin has been recommended for patients with advanced or recurrent disease, non-amenable to cure by surgery and/or radiotherapy (26,34).

4.15 Recommendations for practice
1. Preoperative assessment by endometrial pathology is required to differentiate between tumors at low and high risk of lymph node metastasis, and imaging can be useful in determining depth of invasion, cervical involvement and suspicion of involved nodes. Level of Evidence C.
2. Outside clinical trials, lymphadenectomy should be performed for staging only in high risk cases. There is little evidence to support a therapeutic benefit, but it may be used to select women with positive nodes for radiotherapy. Level of Evidence C.
3. There is evidence that adjuvant radiotherapy is not effective in women with low or moderate risk endometrial cancer in terms of overall survival, though it does reduce the rate of pelvic recurrence. Level of Evidence A. It is certainly indicated in cases with positive nodes and in advanced stage disease. Outside clinical trials, most would advocate its use in the presence of high risk prognostic factors to ensure pelvic control. For surgically staged patients with negative nodes, vaginal brachytherapy may be recommended in higher risk cases. Level of Evidence B.
4. There is no evidence to support the use of adjuvant hormone therapy (progestogen). Level of Evidence A.
5. High risk and advanced stage endometrial cancer should be managed where possible by gynaecological oncologists as part of a multidisciplinary specialist team. Professional consensus.
The configuration of the carcinoma (polypoid or flat), its length, thickness and the number of slices involved (which when multiplied by their estimated thickness, can be translated into an estimate of its width) should be recorded. The depth of myometrial invasion and the width of uninvolved myometrium should be measured. These measurements should be repeated for each plane of the myometrium invaded by tumour (ie: anterior, posterior, left and right lateral walls and fundus). The involvement of lower uterine segment (uterine isthmus) and left and right cornua should be recorded.

The macroscopic estimate of the depth of myometrial invasion will accord with the histological assessment in almost 90% of cases provided the assessment is confined to an estimate of involvement of the inner or outer half (Doeving et al 1989, MK Heatley, personal observation). One or more blocks should be taken through the full thickness of the tumour and uninvolved myometrium. If the myometrium is too thick to fit into one cassette, two should be used. A frozen section study has shown that one (or occasionally two) blocks provide a correct indication of the extent of myometrial invasion in over 90% of cases (Atad et al 1994). It is recognised however, that more extensive histological assessment is desirable if histotechnology resources permit. A histological assessment of the depth of invasion is desirable because pathologists may experience difficulty in determining the true extent of myometrial invasion if, eg, there are associated conditions such as adenomyosis (Jacques et al 1998). It is also desirable to assess the background endometrium to establish if hyperplasia is present (Beckner et al 1985).

Histological examination
The extent to which the specimen is examined histologically will be best determined locally depending upon the availability of histotechnology resources. At a minimum, blocks should be selected to allow adequate FIGO staging for each patient (FIGO 1989). In the case of the uterine specimen, the usual blocks of cervix (midline from the anterior and posterior lips) should suffice unless there is associated cervical pathology. A transverse block from the junction of cervix and uterine isthmus should be processed to exclude or document tumour involvement at this position. It is often possible to identify this junction, since an admixture of cervical and endometrial glands can be
visualised on macroscopic examination of parallel slices from the cut end of the uterus or previously amputated cervical stump.

Blocks of Fallopian tube (to exclude intraluminal tumour extension), ovary (to exclude metastatic or synchronous endometrioid neoplasia) and suspicious serosal lesions should be sampled. Many pathologists routinely examine the cornua histologically since myometrial invasion at these locations represents the point at which the tumour extends closest to the serosa and may upstage the tumour from a stage IB to a IC lesion.

Reports should include:
- tumour type with details of minor components
- grade
- depth of myometrial invasion
- width of tumour-free myometrium
- presence or absence of lymphatic invasion
- involvement of cervical epithelium or stroma.

Other samples which may be recovered include ascitic fluid or peritoneal washings for cytological assessment and tissue from the vagina, bladder, bowel, peritoneum or lymph nodes. If a tumour deposit is macroscopically identifiable, it may be sufficient to sample the tumour only. If no tumour is macroscopically identifiable, it is usually necessary to sample all the submitted tissue histologically to confirm or exclude its presence.

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Cancer of the Fallopian Tube

5.1 Staging

5.1.1. Anatomy

5.1.1.1. Primary Site
The Fallopian tube extends from the posterior superior aspect of the uterine fundus laterally and anteriorly to the ovary. Its length is approximately 10 cm. The lateral end opens to the peritoneal cavity.

5.1.1.2. Metastatic sites
Carcinoma of the Fallopian tube metastasizes in three manners:
- Intraperitoneal seeding with peritoneal implants even with an intact tube
- To the regional lymph node including the para-aortic nodes and pelvic lymph nodes
- Direct extension to the surrounding organs.

5.1.2. Descriptive aspects
The Fallopian tube is a hollow viscus. Disease is often incidentally found at laparotomy or presents as an adnexal mass. Fallopian tube cancer should be staged surgically with a histological confirmation of disease. This includes carcinoma in situ. Tumour extension into the submucosa or muscularis and to and beyond the serosa can therefore be defined. These facts, together with the laterality, in addition to the presence or absence of ascites, must also be taken into consideration.

5.1.3. Surgical staging classification
Staging for the Fallopian tube is by the surgical-pathological system. Operative findings prior to tumour debulking may be modified by histopathologic as well as clinical or radiologic evaluation. The most common staging system used is the FIGO staging system. However, it is useful to know the UICC TMN staging system (See Tables 1 and 2).

5.1.4. Histopathologic types
More than 90% of Fallopian tube carcinoma is papillary serous adenocarcinoma. Other tumours include clear cell carcinoma and endometrioid carcinoma. All these are treated essentially the same way. Rare types include sarcoma, germ cell tumours and lymphoma.

5.1.4.1 Histopathologic grades
- Gx – Grade cannot be assessed
- G1 – Well differentiated (papillary)
- G2 – Moderately differentiated (papillary-alveolar)
- G3 – Poorly differentiated (alveolar-medullary)

Table 1: Carcinoma of the Fallopian tube – Staging

<table>
<thead>
<tr>
<th>FIGO</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>T0</td>
<td>Primary tumour cannot be assessed</td>
</tr>
<tr>
<td>T1</td>
<td>Tumour confined to Fallopian tubes</td>
</tr>
<tr>
<td>T1a</td>
<td>Tumour limited to one tube, without penetrating the serosal surface; no ascites</td>
</tr>
<tr>
<td>T1b</td>
<td>Tumour limited to both tubes, without penetrating the serosal surface; no ascites</td>
</tr>
<tr>
<td>T1c</td>
<td>Tumour limited to one or both tubes, with extension onto/through the tubal serosa; or with positive malignant cells in the ascites or positive peritoneal washings</td>
</tr>
<tr>
<td>T2</td>
<td>Tumour involves one or both Fallopian tubes with pelvic extension</td>
</tr>
<tr>
<td>T2a</td>
<td>Extension and/ or metastasis to uterus and/or ovaries</td>
</tr>
<tr>
<td>T2b</td>
<td>Extension to other pelvic organ</td>
</tr>
<tr>
<td>T2c</td>
<td>IIIB/C with positive malignant cells in the ascites or positive peritoneal washings</td>
</tr>
<tr>
<td>T3and/orN1</td>
<td>Tumour involves one or both Fallopian tubes with peritoneal implants outside the pelvis and/or positive regional lymph nodes</td>
</tr>
<tr>
<td>T3a</td>
<td>Microscopic peritoneal metastasis outside the pelvis</td>
</tr>
<tr>
<td>T3b</td>
<td>Macroscopic peritoneal metastasis outside the pelvis 2 cm or less in greatest dimension</td>
</tr>
<tr>
<td>T3c and/orN1</td>
<td>Peritoneal metastasis more than 2 cm in greatest dimension and/or positive regional lymph nodes</td>
</tr>
<tr>
<td>M1</td>
<td>Distant metastasis beyond the peritoneal cavity</td>
</tr>
</tbody>
</table>
been studied, but there have been no consistent factors identified. However, the similarities in the age group, association with low parity, and frequent infertility status, suggest that the aetiology may be similar to ovarian carcinoma. Indeed, studies have demonstrated similar genetic abnormalities as in ovarian cancer, such as c-erb, p53, k-ras mutations, as well as a recent possible association with BRCA1 and BRCA2.

5.3 Screening
Fallopian tube carcinoma is a rare entity. There is no recommendation for screening.\textsuperscript{(1,4,5,11)}

5.4 Diagnosis

5.4.1 Pre-operative
Abnormal vaginal bleeding is the most common presenting complaint and it is present in more than 50% of patients. This may be associated with watery vaginal discharge, vague lower abdominal pain, distention, and pressure. Ten percent of patients present with ‘hydrops tubae profuens’, a palpable pelvic mass that resolves during examination associated with watery vaginal discharge. More than 50% of patients present with Stage I or Stage II disease, most likely due to its pattern of presentation. Although it has been diagnosed as an incidental finding during pap smear and during CA 125 screening (as part of a randomised control trial), Pap smear and CA 125 cannot be recommended as screening modalities. However, CA125, being raised in a significant percentage of patients, acts as an adjunctive to transvaginal ultrasonography, CT or MRI scan\textsuperscript{(1,2,4)}.

The Fallopian tubes’ close proximity to the ovary and the uterus makes it difficult sometimes to identify the true primary. This is particularly so in the advanced stages. The most widely accepted and therefore most commonly used diagnostic criteria for the diagnosis of primary Fallopian tube carcinoma are shown in Table 3. It was first developed by Hu and later modified by Sedlis\textsuperscript{(11,12)}.

5.4.2 Staging laparotomy
Retrospective analyses have suggested that advanced stages at presentation and the presence of residual tumour at the end of treatment with chemotherapy are associated with poorer prognosis. Therefore

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### Table 2: Carcinoma of the Fallopian tube – Stage grouping

<table>
<thead>
<tr>
<th>FIGO Stage</th>
<th>T</th>
<th>N</th>
<th>M</th>
</tr>
</thead>
<tbody>
<tr>
<td>IA</td>
<td>T1a</td>
<td>N0</td>
<td>M0</td>
</tr>
<tr>
<td>IB</td>
<td>T1b</td>
<td>N0</td>
<td>M0</td>
</tr>
<tr>
<td>IC</td>
<td>T1c</td>
<td>N0</td>
<td>M0</td>
</tr>
<tr>
<td>II A</td>
<td>T2a</td>
<td>N0</td>
<td>M0</td>
</tr>
<tr>
<td>II B</td>
<td>T2b</td>
<td>N0</td>
<td>M0</td>
</tr>
<tr>
<td>III C</td>
<td>T2c</td>
<td>N0</td>
<td>M0</td>
</tr>
<tr>
<td>II A</td>
<td>T3a</td>
<td>N0</td>
<td>M0</td>
</tr>
<tr>
<td>III B</td>
<td>T3b</td>
<td>N0</td>
<td>M0</td>
</tr>
<tr>
<td>III C</td>
<td>T3c</td>
<td>N0</td>
<td>M0</td>
</tr>
<tr>
<td>IV</td>
<td>Any T</td>
<td>N1</td>
<td>M0</td>
</tr>
</tbody>
</table>

### Table 3: Diagnostic criteria

Pathologic criteria for primary Fallopian tube malignancy

1. The tumour arises from the endosalpinx
2. The histologic pattern reproduces the epithelium of tubal mucosa
3. There is transition from benign to malignant epithelium
4. The ovary and endometrium are either normal or with a tumour smaller than the tumour in the tube

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5.2 Overview
Carcinoma of the Fallopian tube is a rare malignancy. It forms between 0.1% to 1.8% of all gynaecological cancers diagnosed and the annual incidence of 3.6 per million women in the USA has not changed much in the past years. More than 60% of Fallopian tube cancer occurs in post-menopausal women. Predisposing factors have

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careful surgical staging at presentation is paramount in the treatment of early Fallopian cancer. The para-aortic nodes above the inferior mesenteric artery are the most frequently involved retroperitoneal nodes (13). For advanced disease, there should also be optimal removal of the primary tumour and involved adjacent organs (1-5,7,14). The following must be performed through a midline incision:

- Careful evaluation of the entire abdomino-pelvic cavity to delineate extent of disease
- Total abdominal hysterectomy and bilateral salpingo-oophorectomy
- Sampling of the pelvic and para-aortic lymph nodes
- Infracolic omentectomy
- Washings of the peritoneal cavity
- Biopsies of any suspicious areas including the abdominal and pelvic peritoneum

In the unlikely event that a patient is young and wishes to retain fertility, limited surgery can be considered for carcinoma in-situ only, after detailed evaluation and careful discussion. However, this limited approach is not encouraged in view of the high incidence of bilateral involvement. In established cancer, there is no role for conservative surgery (1-2,14).

5.5. Clinical Practice Guidelines

5.5.1. Management of Fallopian tube adenocarcinoma

Since the first reported case in 1847, there have only been 1,500 cases documented in literature. Further, retrospective reviews have been hampered with non-uniform clinical and surgical staging, non-central pathological reviews, and lack of diagnostic criteria. As more retrospective reviews were reported, especially after the wide acceptance of the FIGO staging, it became apparent that the histology, prognostic indicators and survival mirror ovarian cancer. The only exceptions seem to be that firstly, stage for stage, patients with early Fallopian carcinoma have a poorer prognosis. Secondly, Fallopian tube carcinoma has a significant rate of lymph node metastases. In view of the similarities with ovarian cancer and the impracticality of performing prospective randomized trials in a relatively rare cancer, the management of Fallopian tube carcinoma has been essentially identical to that of ovarian cancer. The guidelines for post-operative management are also essentially identical to those of ovarian cancer.

5.5.1.1. Chemotherapy regimes

The success of the paclitaxel and platinum combination in ovarian cancer has led to greater usage of this combination in Fallopian cancer. Many retrospective reviews seem to suggest that this regime is superior to historical controls using a combination of alkylating agent with platinum. It is therefore the view of many clinicians that chemotherapy for ovarian cancer should be acceptable for Fallopian cancer in the absence of prospective studies (1-4,5,7,14-19). For chemotherapy regimes please refer to ovarian cancer regimes.

5.5.1.2. Management of early disease

5.5.1.2.1. Management of carcinoma in situ

Patients should undergo laparotomy with resection of the tumour as outlined above. There is no recommendation for adjuvant therapy for carcinoma in-situ of the Fallopian tube after primary surgical therapy (1,2,14).

5.5.1.2.2. FIGO Stage I and Stage II

Patients with early stage disease should undergo surgical staging. Patients with a final histologic diagnosis of adenocarcinoma in situ or Stage I grade 1 tumour do not require postoperative adjuvant chemotherapy. All other patients should be considered for adjuvant platinum-based chemotherapy. Patients whose diagnosis was incidental (i.e., patients who underwent surgery for a benign condition, and the histologic specimen contained malignancy) should undergo repeat surgical staging, and optimal debulking, should there be metastatic disease. These patients should then receive adjuvant platinum-based chemotherapy (1,3,15-19). Level of Evidence C.

5.5.1.3. Management of advanced disease

5.5.1.3.1 FIGO Stage III

All carcinoma is taken to be adenocarcinoma unless otherwise stated and platinum-based chemotherapy regimes are as per ovarian cancer. Patients who have undergone surgical debulking should be considered for adjuvant platinum-based chemotherapy (1-5,15-18). Patients who
have not undergone optimal debulking at initial diagnosis because of medical contraindications should receive platinum-based chemotherapy followed by re-evaluation. After 3 cycles of chemotherapy, these patients should be considered for an interval debulking procedure, should there be any residual tumour. However, this practice has not been validated with any prospective trial in Fallopian tube cancer. **Level of Evidence C.**

5.5.1.3.2. FIGO Stage IV
- Patients with distant metastases should have the primary site of disease confirmed histologically.
- As much tumour as possible should be resected at laparotomy, and a symptomatic pleural effusion should be drained preoperatively.
- Patients who are of adequate performance status should receive platinum-based chemotherapy, as per ovarian cancer. For patients too ill to receive chemotherapy, symptomatic treatment should be offered. **Level of Evidence C.**

5.5.2. Management of choriocarcinoma of the Fallopian tube
This entity is extremely rare, but has been reported in tubal pregnancy and also associated with in-vitro fertilisation. Patients should be treated as for uterine choriocarcinoma, which is curable, with primary surgery followed by chemotherapy according to the prognostic factors. In this case, conservative surgery can be considered for limited disease if fertility is to be preserved. **Level of Evidence D.**

5.5.3. Management of germ cell tumours of the Fallopian tube
These are extremely rare in the Fallopian tube. However, they do occur in young women with fertility potential, and although highly curable, they tend to progress rapidly. Therefore, it is pertinent to diagnose and treat early. Treatment is with primary surgery followed by chemotherapy according to the prognostic factors. Limited conservative surgery should be considered for disease of all stages if fertility is to be preserved. Chemotherapy protocols are as for ovarian germ cell tumours. **Level of Evidence D.**

5.5.4. Management of sarcoma of the Fallopian tube
Tubal sarcomas are extremely rare lesions. Most sarcomas histologically are classified as mixed Mullerian tumours. Treatment again is with primary surgery and chemotherapy as per sarcoma of uterine origin. **Level of Evidence D.**

5.5.5. Management of miscellaneous rare histologies
The Fallopian tube is not exempted from other rare tumours such as mucosa related lymphoma MALT. Treatment is with primary surgery and systemic chemotherapy. The chemotherapy protocols are directed by the individual histologies. **Level of Evidence D.**

5.5.6. Follow-up
There is no evidence to show that intensive clinical monitoring in asymptomatic women has any positive impact on overall survival or on the quality of life. Nonetheless, early diagnosis of recurrence after a prolonged progression free interval is thought to offer the best results. The objectives of follow-up are as follows:
- Determination of the patient’s immediate response to the treatment employed;
- Early recognition and prompt management of any treatment-related complications, including any psychological sequelae;
- Early detection of persistent or recurrent disease;
- Collection of data regarding the efficacy of treatment;
- For patients with early disease, it serves as an opportunity for breast cancer screening, and for patients treated with conservative surgery, for cervical cancer screening.

In general, during the first year following treatment, patients should be seen every three months with a gradual increase in intervals to every four to six months and annually after the fifth year. At each follow-up, the patient should have her history retaken and complete physical examination (including breast, pelvic and rectal examination) performed to exclude any clinical signs of recurrence. The serum CA125 titer may also be checked at regular intervals, especially if it was raised at primary diagnosis, although the literature in this area is unclear as to the impact of such a practice on survival. Radiological tests such as ultrasonography of the pelvis, CT scans or MRI scans should only be performed when the clinical findings or the tumour markers suggest possible recurrence. **Level of Evidence D.**

All patients with an intact cervix should receive a regular Pap
smear. All patients above the age of 40 should undergo a yearly routine mammogram, as should younger patients with a strong family history of breast cancer.

References
Cancer of the Ovary

6.1. Staging

6.1.1. Sites of ovarian cancer

6.1.1.1 Primary site

The ovaries are a pair of solid oval-shaped organs, 2-4 cm in diameter, that are connected by a peritoneal fold to the broad ligament and by the infundibulopelvic ligament to the lateral wall of the pelvis.

6.1.1.2. Nodal drainage

The lymphatic drainage occurs by the utero-ovarian, infundibulopelvic, and round ligament trunks and an external iliac, common iliac, hypogastric, lateral sacral, para-aortic nodes and, occasionally, to the inguinal nodes (1,5).

6.1.1.3. Metastatic sites

The peritoneum, including the omentum and pelvic and abdominal viscera, is the most common site for seeding. This includes involvement of the diaphragmatic and liver surfaces. Pleural involvement is also frequent. Other extra peritoneal or extra pleural sites are relatively uncommon but can still occur (1,3).

6.1.2. Rules for classification

Although CT scans can delineate the intra-abdominal spread of disease to a certain extent, ovarian cancer should be staged surgically. There should be histological confirmation of the disease. Operative findings determine the stage and therefore the prognosis of the patient (1,3).

Chest X-rays serve as a screen for pleural metastases. As extra-pulmonary and extra-peritoneal metastases are infrequent, there is no requirement for other radiological evaluation unless symptomatic. Serum CA 125 titer is useful in determining response to chemotherapy, although it does not contribute to staging.

6.1.2.1. Evaluation of surgical staging

If the pre-operative suspicion is malignancy, the laparotomy should 
be performed via a midline incision. The following should be performed for adequate staging:\(^{(1,8)}\):
- Careful evaluation of all peritoneal surfaces
- 4 washings of the peritoneal cavity: diaphragm, right and left abdomen, pelvis
- Infracolic omentectomy
- Selected lymphadenectomy of the pelvic and para-aortic lymph nodes
- Biopsy and/or resection of any suspicious lesions, masses and any adhesions
- Random blind biopsies of normal peritoneal surfaces, including that from the undersurface of the right hemidiaphragm, bladder reflection, cul-de-sac, right and left paracolic recesses, and both pelvic sidewalls
- Total abdominal hysterectomy and bilateral salpingo-oophorectomy
- Appendectomy for mucinous tumours

6.1.2.2. Post-surgical treatment – pathologic staging
The biopsies taken as outlined above would form the basis of staging. However, any other suspicious area, such as pleural effusion, and rare but obvious involvement of extra pulmonary or pleural and extra peritoneal sites must be biopsied\(^{(1,8)}\).

6.1.2.3. International Federation of Gynecology and Obstetrics (FIGO) staging
The most commonly utilised staging system is the FIGO system modified in 1988. It is based on findings made mainly through surgical exploration as outlined above. See Table 1 for the complete FIGO staging\(^{(1,3,4)}\). However, it is also useful to be aware of the equivalents within the UICC TNM classification (See Table 1, and Table 2).

<table>
<thead>
<tr>
<th>FIGO</th>
<th>TNM</th>
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</thead>
<tbody>
<tr>
<td>Primary tumour cannot be assessed</td>
<td>TX</td>
</tr>
<tr>
<td>No evidence of primary tumour</td>
<td>T0</td>
</tr>
<tr>
<td>Tumour confined to ovaries</td>
<td>T1</td>
</tr>
<tr>
<td>Tumour limited to one ovary, capsule intact</td>
<td>T1a</td>
</tr>
<tr>
<td>No tumour on ovarian surface</td>
<td></td>
</tr>
<tr>
<td>No malignant cells in the ascites or peritoneal washings</td>
<td></td>
</tr>
<tr>
<td>Tumour limited to both ovaries, capsules intact</td>
<td>T1b</td>
</tr>
<tr>
<td>No tumour on ovarian surface</td>
<td></td>
</tr>
<tr>
<td>No malignant cells in the ascites or peritoneal washings</td>
<td></td>
</tr>
<tr>
<td>Tumour limited to one or both ovaries, with any of the following:</td>
<td>T1c</td>
</tr>
<tr>
<td>Capsule ruptured, tumour on ovarian surface, positive malignant cells in the ascites or positive peritoneal washings</td>
<td></td>
</tr>
<tr>
<td>Tumour involves one or both ovaries with pelvic extension</td>
<td>T2</td>
</tr>
<tr>
<td>Extension and/or implants in uterus and/or tubes</td>
<td>T2a</td>
</tr>
<tr>
<td>No malignant cells in the ascites or peritoneal washings</td>
<td></td>
</tr>
<tr>
<td>Extension to other pelvic organ</td>
<td>T2b</td>
</tr>
<tr>
<td>No malignant cells in the ascites or peritoneal washings</td>
<td></td>
</tr>
<tr>
<td>II A/B with positive malignant cells in the ascites or positive peritoneal washings</td>
<td>T2c</td>
</tr>
<tr>
<td>Tumour involves one or both ovaries with microscopically confirmed peritoneal metastasis outside the pelvis and/or positive peritoneal washings</td>
<td></td>
</tr>
<tr>
<td>Microscopic peritoneal metastasis beyond the pelvis</td>
<td>T3a</td>
</tr>
<tr>
<td>Macroscopic peritoneal metastasis beyond the pelvis 2 cm or less in greatest dimension</td>
<td>T3b</td>
</tr>
<tr>
<td>Peritoneal metastasis beyond pelvis more than 2 cm in greatest dimension and/or regional lymph nodes metastasis</td>
<td>T3c</td>
</tr>
<tr>
<td>Staging IV. Pleural effusion must have positive cytology.</td>
<td></td>
</tr>
</tbody>
</table>

Note: Liver capsule metastasis is T3/Stage III, liver parenchymal metastasis M1/Stage IV.
they are too poorly differentiated to be placed in any other group.
- Mixed epithelial tumours
  - These tumours are composed of two or more of the five major cell types of common epithelial tumours. The types are usually specified.
- Cases with intraperitoneal carcinoma in which the ovaries appear to be incidentally involved and not the primary origin should be labelled as extra-ovarian peritoneal carcinoma.

Epithelial tumours of the ovary are also further sub-classified by grading. This is important because histological grading is proportional to prognosis. This grading system does not apply to non-epithelial tumours.
- Gx – Grade cannot be assessed
- G1 – Well differentiated
- G2 – Moderately differentiated
- G3 – Poorly differentiated

Non-epithelial malignancy, although more uncommon, is also extremely important. These include granulosa cell tumours, germ cell tumours, sarcomas and lymphomas. They shall be discussed as separate entities.

6.1.3. Histopathologic classification

Majority of ovarian cancer is of the epithelial origin. The task forces of FIGO endorse the histological typing of epithelial ovarian tumours as presented in the WHO publication n.9 in 1971. It is recommended that all ovarian epithelial tumours be subdivided according to a simplified version.2

Epithelial ovarian neoplasms are classified as follows:
- Serous tumours
- Mucinous tumours
- Endometrioid tumours
- Clear cell tumours
- Brenner tumours
- Undifferentiated carcinomas
  - This group of malignant tumours is of epithelial structure but

### Table 2: Carcinoma of the ovary – Stage grouping

<table>
<thead>
<tr>
<th>FIGO Stage</th>
<th>T</th>
<th>N</th>
<th>M</th>
</tr>
</thead>
<tbody>
<tr>
<td>IA</td>
<td>T1a</td>
<td>N0</td>
<td>M0</td>
</tr>
<tr>
<td>IB</td>
<td>T1b</td>
<td>N0</td>
<td>M0</td>
</tr>
<tr>
<td>IC</td>
<td>T1c</td>
<td>N0</td>
<td>M0</td>
</tr>
<tr>
<td>II A</td>
<td>T2a</td>
<td>N0</td>
<td>M0</td>
</tr>
<tr>
<td>IIIB</td>
<td>T2b</td>
<td>N0</td>
<td>M0</td>
</tr>
<tr>
<td>IIIC</td>
<td>T2c</td>
<td>N0</td>
<td>M0</td>
</tr>
<tr>
<td>IIIA</td>
<td>T3a</td>
<td>N0</td>
<td>M0</td>
</tr>
<tr>
<td>IIIB</td>
<td>T3b</td>
<td>N0</td>
<td>M0</td>
</tr>
<tr>
<td>IIIC</td>
<td>T3c</td>
<td>N0</td>
<td>M0</td>
</tr>
<tr>
<td>IV</td>
<td>Any T</td>
<td>N1</td>
<td>M0</td>
</tr>
</tbody>
</table>

**Regional Nodes (N)**
- Nx Regional lymph nodes cannot be assessed
- N0 No regional lymph node metastasis
- N1 Regional lymph node metastasis

**Distant Metastasis (M)**
- Mx Distant metastasis cannot be assessed
- M0 No distant metastasis
- M1 Distant metastasis (excluding peritoneal metastasis)

6.2. Introduction

Malignant tumours of the ovaries occur at all ages. Major histologic types occur in different age groups. For women less than 20 years of age, germ cell tumours constitute the majority of cases, while epithelial ovarian cancers (EOC) are primarily seen in women older than 50 years.

EOC is a relatively common disease in the USA. The lifetime risk of a woman in the United States developing ovarian cancer is approximately 1 in 70. Approximately 23% of gynaecologic cancers are ovarian in origin, but 47% of all deaths from cancer of the female genital tract occur in women with ovarian cancer. Overall, epithelial ovarian cancer accounts for 4% of all new cancer diagnoses in women and 5% of all cancer related deaths.

The overall incidence of EOC varies from 9 to 17/100,000 and is highest in industrialised countries, with the exception of Japan. How-
Borderline tumours occur most commonly in the perimenopausal age group. For younger patients, tumour markers such as human gonadotropin (hCG) and alpha-fetoprotein (AFP) are mandatory to exclude germ cell tumours.

EOC in its early stages does not usually produce symptoms or signs that would alert the clinician to this diagnosis. Approximately two-thirds of all EOCs are Stage III or Stage IV at diagnosis. Symptoms that include vague abdominal pain or discomfort, menstrual irregularities, dyspepsia and other mild digestive disturbances, which may only have been present for a few weeks, are usually the presenting complaint. Therefore, a high index of suspicion is required for all women between the ages of 40 to 69. As the disease progresses, the ascites, abdominal distention and discomfort generally worsen and may be associated with respiratory symptoms from increased intra-abdominal pressure or from the transudation of fluid into the pleural cavities. Abnormal vaginal bleeding is uncommon as a symptom or sign of disease.

A detailed medical history must be taken to ascertain possible risk factors, history of other cancers and history of cancer in the family. Then a complete physical examination, including breast, pelvic and rectal examination, must be performed. A Pap smear is also usually performed at the same time.

Prior to surgery a chest radiograph should be taken to screen for pulmonary and pleural metastases while a CT scan of the abdomen and pelvis should be performed to delineate the extent of intra-abdominal disease or the presence of another primary. However, in the absence of extra-abdomino-pelvic disease, radiological scanning does not replace surgical staging with laparotomy. Barium enema or colonoscopy is indicated should symptoms suggest possible bowel cancer. Tumour markers including CA 125, and carcinoembryonic antigen (CEA) should be considered. With a high CA 125, the most common diagnosis would be EOC. However, a stomach or colon primary with metastases to the ovaries may mimic ovarian cancer. A current mammogram should also be considered as patients are frequently in the age group where breast cancer is prevalent.

6.3. Screening
To date no cost-effective screening program for ovarian cancer is available. Studies using CA 125, ultrasonography of the pelvis and pelvic examination have not produced an acceptable level of sensitivity and specificity in women of normal risks. Patients with a strong family history of epithelial ovarian cancer, especially of the syndromes described above, should consult a genetic counselor to stratify their risks, and when appropriate, be placed on prospective screening trials. Presently there are no screening studies for non-EOC.

6.4. Diagnosis
The success of treatment depends on early diagnosis. However, the ability to achieve early diagnosis remains an unsolved problem. The clinician must be mindful of the different neoplasms that occur in different age groups in order to have a high index of suspicion.
6.4.1. Staging laparotomy and surgical management
Generally, the prognosis of epithelial ovarian tumours is independently affected by the following:
• Stage of the cancer at diagnosis;
• The histological sub-type and grading;
• The volume of residual disease
Of the three, the most important are the stage and the volume of residual disease. This holds true regardless of the histological diagnosis.
A thorough staging laparotomy is therefore the most important part of early management. Upon entering the abdominopelvic cavity through a midline incision, the peritoneal fluid should be sent for cytology. In the absence of ascites, irrigation should be performed and washings should be sent for cytology.
The laparotomy should then proceed with a detailed examination of the contents including all the peritoneal surfaces. In addition to all the suspicious sites, random biopsies from the peritoneal reflection of the bladder, the posterior cul-de-sac, both paracolic gutters, sub-diaphragmatic surfaces and both pelvic sidewalls should be taken. The primary tumour, if limited to the ovary, should be examined to look for capsular rupture. All obvious sites of tumour must be removed wherever possible in addition to total hysterectomy and bilateral salpingo-oophorectomy. Further, the omentum, pelvic and para-aortic lymph nodes should also be removed for histological examination. This procedure allows for accurate staging of early disease and has the objective of optimal debulking of advanced disease.
In younger women fertility is an issue. In these patients, the prognosis according to the extent of the tumour should be discussed, and conservative surgery can be considered with informed consent.
Conservative surgery should require the following:
• Laparotomy assessment as outlined above in section 6.1.2.1, except total hysterectomy and bilateral salpingo-oophorectomy;
• Intraoperative finding of unilateral ovarian involvement with capsule intact (stage IA);
• Normal examination of the opposite ovary (without wedge biopsy)

6.5. Clinical Practice Guidelines
6.5.1. Management of patients in the reproductive age group with a suspicion of cancer diagnosis
Clinical judgment is a major factor in surgical decision-making and this is the most pertinent in the approach to a pelvic mass in the young, reproductive aged woman. In the past, the options were essentially limited to either laparotomy with assessment and removal, or observation with regular pelvic ultrasonography. Advances in laparoscopic surgery have provided another option for evaluation and potential treatment.
If the suspicion is strong for malignancy, open laparotomy is indicated. In the presence of good surgical skills, laparoscopy is more appropriate if the suspicion is more for benign disease in a young woman, where tumour markers (including human chorionic gonadotropin and alpha-feto protein) are normal.
The following factors point to the presence of a malignancy, and are useful in the clinical assessment of masses:
• Age of the patient (young for germ cell, older for EOC)
• Bilaterality
• Tumour fixation clinically
• Ascites
• Ultrasonographically complex
• CT finding of metastatic nodules
• Elevated tumour markers

6.5.2. Management of epithelial ovarian cancer (EOC)
6.5.2.1. Early stages
About a quarter of patients will present with apparent Stage I or Stage II disease. Although the radiological findings may seem to correlate with gross findings at laparotomy, it is imperative that these patients undergo a thorough surgical staging. The details of the staging laparotomy have been outlined above. For patients with obviously limited disease (Stage IA), who desire to retain their reproductive potential, a wedge biopsy of the contralateral unaffected ovary is not recommended as it can affect the subsequent fertility.
The prognosis of adequately staged patients with Stage IA and Stage IB Grade I cystadenocarcinoma is extremely good such that
adjuvant chemotherapy would not provide further benefits. For higher grade tumours and for patients with Stage IC disease, adjuvant platinum-based chemotherapy should be considered, although this practice remains controversial. All patients with Stage II disease should receive adjuvant chemotherapy. The number of cycles of chemotherapy has also not been clarified but usually between 3-6. Level of Evidence A.

6.5.2.2 Advanced stages
Three quarters of all patients with ovarian cancer present with Stage III or IV disease. These patients are usually quite symptomatic from the intra-abdominal disease. This may affect the performance status and fitness for surgery. However, one of the most critical prognostic indicators in patients with advanced stage ovarian cancer is the volume of residual disease. Therefore, all patients whose medical condition permits should undergo primary surgical laparotomy with maximal attempt at optimal cytoreduction. Systemic pelvic and para-aortic lymphadenectomy does not improve overall survival, when compared to removal of bulky nodes only, although there is a modest improvement in progression-free survival. Level of Evidence A.

In certain patients whose primary cytoreduction is considered suboptimal, interval debulking may be considered after three cycles of systemic chemotherapy. This also applies to patients who could not, in view of physical fitness, undergo primary cytoreduction prior to chemotherapy.

Patients who have had cytoreduction should receive adjuvant chemotherapy. For systemic chemotherapy, a combination of a paclitaxel or docetaxel with carboplatin is the first choice. Docetaxel is considered because of its favourable neurotoxicity profile. At the end of six cycles of chemotherapy, maintenance chemotherapy with paclitaxel has been shown to improve disease-free interval but not overall survival. However, this treatment must only be offered if a patient achieves complete response to treatment, and understands the aim of treatment and its potential toxicities.

The role of intraperitoneal chemotherapy remains controversial, but the recent Gynecologic Oncology Group trial compared intravenous paclitaxel plus cisplatin with intravenous paclitaxel plus intraperitoneal cisplatin and paclitaxel in patients with Stage III ovarian or primary peritoneal carcinoma, with no residual disease greater than 1 cm in diameter. Only 42% of patients in the intraperitoneal group completed 6 cycles of the assigned therapy, but the intraperitoneal group had an improvement in progression-free survival of 5.5 months (23.8 vs. 18.3 months; P = 0.05) and an improvement in overall survival of 15.9 months (65.6 vs. 49.7 months; P = 0.03). Further studies of intraperitoneal therapy are ongoing. The controversy relates to the increased toxicity and catheter-related complications. Level of Evidence A.

6.5.2.3 Chemotherapy for EOC
The following chemotherapy regimes are recommended for the treatment of ovarian cancer:

- Paclitaxel 175mg/m^2 over 3 hours/ Carboplatin AUC 6 over 1 hour
- Docetaxel at 75mg/m^2 over 1 hour/ Carboplatin AUC 5 over 1 hour

The following chemotherapy schedules can also be considered:

- Paclitaxel 135mg/m^2 over 24 hours/ Cisplatin 75mg/m^2 over 6 hours
- Paclitaxel 175mg/m^2 over 3 hours/ Cisplatin 75mg/m^2 over 6 hours

In the GOG study of intraperitoneal chemotherapy, the IP arm of the study was as follows:

IV paclitaxel 135mg/m^2 over 24 hours on day 1
IP cisplatin 100 mg/m^2 on day 2
IP paclitaxel 60 mg/m^2 on day 8.

6.5.2.4 Second-look laparotomy
A second-look laparotomy (or laparoscopy) is performed on a patient who has no clinical evidence of disease after a prescribed course of chemotherapy, in order to determine response to treatment. It has not been shown to influence survival, although the information obtained correlates with subsequent outcome. Level of Evidence C.
6.5.2.5 Secondary cytoreduction
Secondary cytoreduction may be defined as an attempt at cytoreductive surgery at some stage following completion of first-line chemotherapy. Retrospective studies suggest that patients benefit if all macroscopic disease can be removed, which usually means patients with a solitary recurrence. Patients with a disease-free interval \( \geq 24 \) months appear to derive most benefit\(^{25,26} \), although if all macroscopic disease can be removed, benefits can even occur at the time of a second-look laparotomy\(^{27} \). Level of Evidence C.

6.5.2.6 Follow-up for malignant EOC
There is no evidence to show that intensive clinical monitoring in asymptomatic women has any positive impact on overall survival or on the quality of life. Nonetheless, early diagnosis of recurrence after a prolonged progression-free interval is thought to offer the best results. The objectives of follow-up are as follows:
- Determination of a patient’s immediate response to the treatment employed;
- Early recognition and prompt management of any treatment-related complications, including any psychological sequelae;
- Early detection of persistent or recurrent disease;
- Collection of data regarding the efficacy of any treatment and the complications associated with those treatments;
- Facilitation of screening for breast cancer in patients with early disease, and to facilitate screening for cervical cancer for patients with conservative surgery.

In general, during the first year following treatment, patients should be seen every three months with a gradual increase in intervals to every four to six months and annually after the fifth year. At each follow-up, the patient should have her history retaken; complete physical examination (including breast, pelvic and rectal examination) performed to exclude any clinical signs of recurrence. The CA125 should also be checked at regular intervals. It is unclear as to the utility of such a practice on survival\(^{10} \). Radiological tests such as ultrasonography of the pelvis, CT, MRI, and/or PET scans should only be performed when the clinical findings or the tumour markers suggest possible recurrences Level of Evidence C.

All patients with intact cervix should receive a regular Pap smear.
effusion, and unusual metastases to organs such as the brain, liver or bone.

The treatment of an asymptomatic patient with recurrent disease based on tumour marker alone is difficult. Close observation or hormonal therapy with agents such as Tamoxifen can be considered. It is important that the patient understands that responses to chemotherapy do not necessarily translate into meaningful survival prolongation. Often improvement in quality of life and optimisation of functionality become the goals of treatment. Any therapy that may compromise the latter goals may not be justified. At these difficult times, it is of utmost importance to involve the patient’s friends, family and loved ones in the decision-making. Level of Evidence C.

6.5.3 Management of epithelial cancer of low malignant potential (borderline tumour)

Compared to obviously malignant neoplasms, these borderline tumours tend to affect a younger population. They constitute 15% of all EOC. Nearly 75% of these are Stage I at the time of diagnosis. The following can be said for these tumours:
• The diagnosis must be based on the original tumour.
• Extensive sectioning of the neoplasm is necessary to rule out truly invasive characteristics.
• The prognosis of these tumours is extremely good, with a 10-year survival of about 95%.
• Lesions that behave in a malignant fashion usually have an indolent course.
• There are occasional spontaneous regressions of peritoneal implants.
• Early stage, serous histology, and younger age at diagnosis are associated with a more favourable prognosis.
• Although gross residual disease after primary laparotomy is associated with poorer prognosis, mortality from the disease remains unusual.

The cause of death has been determined to be benign complications of disease (e.g., small bowel obstruction), complications of therapy, and only rarely malignant transformation. The mainstay of treatment is primary surgical staging and cytoreduction. For patients with Stage I disease that still desire to have children, conservative surgery with unilateral oophorectomy can be considered after intra-operative inspection of the contralateral ovary to exclude involvement. For patients with only one ovary, or bilateral cystic ovaries, a partial oophorectomy can be considered to retain fertility. For all other patients, total hysterectomy and bilateral salpingo-oophorectomy is recommended, with maximal cytoreduction if the disease is metastatic.

Optimally cytoreduced patients in all stages of disease should receive only expectant treatment without adjuvant chemotherapy, provided the metastases are also borderline tumours histologically. There have been no prospective studies that suggest benefit of adjuvant chemotherapy. A small percentage of patients may benefit from chemotherapy. These are patients in whom the histological specimens reveal invasive implants on the peritoneal surfaces or omentum. Patients who develop a rapid recurrence of intraperitoneal disease also require chemotherapy. Such patients probably had undetected invasive cancers.

For patients with slow recurrence of the disease, especially after a long disease-free interval, cytoreduction should be repeated, and chemotherapy only given if frankly invasive disease is now found histologically.

Follow-up of patients with no evidence of disease is as those of malignant EOC, but less frequent intervals are appropriate. If the contralateral ovary has been retained, it should be followed by transvaginal ultrasonography, at least on an annual basis. Level of Evidence C.

6.5.4 Management of granulosa cell tumors

Granulosa cell tumors account for about 70% of sex-cord stromal tumours, and 3-5% of all ovarian neoplasms. There are two types of granulosa cell tumour: the juvenile and the adult type. Due to the high oestrogen production, the juvenile type presents with sexual precocity, while the adult type may present with post-menopausal bleeding. Because the presenting symptoms are usually prominent, and the tumours are usually slow growing, the majority of patients are diagnosed with Stage I disease. The peak incidence is in the first post-menopausal decade.

The nature of these tumours, like those of borderline tumours, is
generally slow growing with a tendency to late recurrence. Stage at
diagnosis is the most important prognostic indicator. Others of signif-
ificance include age of patient, tumour size, and histological features.
If metastatic, adequate cytoreduction is the mainstay of treatment. If
the patient is young and the disease is confined to one ovary, conser-
ватive surgery should be performed.

The infrequency of the disease, and its protracted course, has result-
ed in a lack of prospective studies. There is no evidence that adjuvant
chemotherapy or radiotherapy improves the results obtained with
good surgery alone for Stage I disease. However, some clinicians
advocate adjuvant chemotherapy for Stage II and Stage III disease
This strategy is based on small studies.

Follow-up is clinical. Serum inhibin is a useful tumour marker.

**Level of Evidence C.**

6.5.5 Management of germ cell tumors

6.5.5.1 Introduction

This group of ovarian tumors consists of a variety of histologically
different entities that are all derived from the primitive germ cells of
the embryonic gonad. Malignant germ cell tumors represent a rela-
tively small proportion of all ovarian tumors. Prior to advances in
chemotherapy, the prognosis for these aggressive tumors was poor.
Over the past decade, new chemotherapeutic regimes have made
germ cell tumors among the most highly curable cancers.

6.5.5.2 Presentation

The highest incidence of germ cell tumors occurs in the second and
third decades of life. It is frequently diagnosed by finding a palpable
abdominal mass in a young woman who complains of abdominal
pain. The following are the symptoms of germ cell tumors in order
of frequency:

- **Acute abdominal pain**
- **Chronic abdominal pain**
- **Asymptomatic abdominal mass**
- **Abnormal vaginal bleeding**
- **Abdominal distention**

6.5.5.3 Histological classification

The classification of germ cell tumours of the ovary is important for
prognostication and for treatment with chemotherapy. Germ cell
tumours are classified as follows:

- **Germ cell tumours**
  - Dysgerminoma
  - Non-dysgerminoma (embryonal cancer)
    - Embryonal differentiation
    - Mixed
    - Mature
    - Immature
    - Extraembryonal differentiation
      - Choriocarcinoma
      - Endodermal sinus tumour (yolk sac tumour)
      - Extraembryonal carcinoma

6.5.5.4 Diagnosis, staging and surgical management

Germ cell tumours are staged the same as epithelial ovarian cancer.
The treatment is dependent not only on the stage. Dysgerminoma is
the equivalent of seminoma in testicular cancer. It is exquisitely
chemotherapy and radiotherapy sensitive. The cure rate is high irre-
spective of the stage of the neoplasm. The other histologies are really
equivalent to non-seminomas in testicular cancer. The aggressiveness
of the disease is dependent on the type, the most aggressive being
endodermal sinus and choriocarcinoma. With chemotherapy, they are
also highly curable.

As chemotherapy can cure the majority of patients even with
advanced disease, conservative surgery is standard in all stages of all
germ cell tumors. Conservative surgery means laparotomy with care-
ful examination and biopsy of all suspicious areas, with limited
cytoreduction, thereby avoiding major morbidity. The uterus and the
contralateral ovary should be left intact if they are normal. Wedge
biopsy of a normal ovary is not recommended as it defeats the pur-
pose of conservative therapy by causing possible infertility. Patients
who receive conservative surgery with the preservation of one ovary
retain acceptable fertility rates despite adjuvant treatment with
chemotherapy. There has been no report of higher adverse obstetric
outcome or long-term unfavourable sequelae in the offspring.
Secondary surgery is of no proven benefit except in a minority of patients whose tumour was not completely resected at the initial surgical procedure and who had teratomatous elements in their primary tumour. Surgical resection of residual masses detected by clinical examination or by radiographic procedures may be beneficial as such masses may contain mature teratoma or residual tumour\(^{[52,54]}\).

6.5.5.5 Post-operative management and follow-up of dysgerminoma
Patients with Stage IA disease may be observed after surgery. A small proportion of patients may recur, but they can be treated successfully at the time of recurrence with a high rate of cure\(^{[1]}\). Patients with disease beyond the ovary should receive adjuvant chemotherapy. Radiotherapy is probably as effective as chemotherapy but ovarian failure makes it undesirable for patients with an intact ovary. However, for patients with any contraindication for chemotherapy, radiotherapy remains an effective option.

Dysgerminoma is extremely sensitive to chemotherapy, and treatment with chemotherapy cures the majority of patients even with advanced disease. The recommended chemotherapy regime is etoposide 100mg/m\(^2\) per day for 5 days with cisplatin 20mg/m\(^2\) per day for 5 days, with or without bleomycin at 10U per day for days 1/8/15 (EP or BEP; various schedules of bleomycin are utilised).

When there is bulky residual disease, it is common to give 3 to 4 courses of combination BEP chemotherapy. As BEP chemotherapy is associated with a lower relapse rate and shorter treatment time\(^{[42]}\), it is preferred compared to an older regime VAC\(^{[43]}\), a combination of vincristine, dactinomycin, and cyclophosphamide. Other tested chemotherapy regimes include combinations of ifosfamide and doxorubicin; vinblastine, ifosfamide and cisplatin; cyclophosphamide, doxorubicin and cisplatin, and POMB-ACE, a combination of cisplatin, vincristine, bleomycin, methotrexate, actinomycin D, cyclophosphamide and etoposide\(^{[48]}\). All patients who do not respond to standard therapy are candidates for clinical trials. Level of Evidence B.

All patients should be followed every 1-2 months for year 1, every 2 months for year 2, every 3 months for year 3, every 4 months for year 4, every 6 months for year 5, and once a year subsequently. At each visit, there should be a medical history, physical examination and tumour markers estimation. Although tumour markers are important, radiological imaging is also pertinent, especially for patients whose tumour markers were not raised at diagnosis. CT scans should be performed as clinically indicated.

Patients who did not receive chemotherapy should be followed-up more closely. Ninety percent of relapses in patients treated with chemotherapy occur within the first two years. At relapse, these patients can be successfully treated\(^{[55]}\). Level of Evidence D.

6.5.5.6 Post-operative management and follow-up of non-dysgerminoma
With chemotherapy, these tumours are also highly curable, even with advanced disease. Patients with Stage IA Grade II immature teratoma have a very good prognosis and should only be observed after primary conservative surgery. Patients with Stage IA Grade I immature teratoma have a very good prognosis. It is controversial whether adjuvant chemotherapy adds any further overall survival benefit in this subgroup of patients. All other patients with higher stage and higher-grade tumours should receive postoperative adjuvant chemotherapy\(^{[50]}\).

The recommended chemotherapy regime is etoposide 100mg/m\(^2\) per day for 5 days with cisplatin 20mg/m\(^2\) per day for 5 days, with or without bleomycin at 10U per day for days 1/8/15 (EP or BEP; various bleomycin schedules are utilised). When there is bulky residual disease it is common to give 3 to 4 courses of combination chemotherapy and 2 additional courses after achieving serological remission. As BEP chemotherapy is associated with a lower relapse rate and shorter treatment time\(^{[42]}\), it is preferred compared to the older regime VAC\(^{[44]}\), a combination of vincristine, dactinomycin, and cyclophosphamide.

Patients who do not respond to BEP may still attain a durable remission with cisplatin/vinblastine/ifosfamide (VIP) as salvage therapy\(^{[45]}\). Newer potential treatments include an ifosfamide combination or high-dose chemotherapy and autologous marrow rescue. After chemotherapy, patients with metastatic immature teratomas can sometimes have residual masses, which are composed entirely of mature elements. These masses can grow, and often require surgical
removal for symptom control. All patients who do not respond to
standard therapy are candidates for clinical trials. **Level of Evidence B.**

All patients should have lactate dehydrogenase (LDH), alpha-feto
protein (AFP), and human gonadotropin (beta hCG) performed to
monitor the response to treatment. All patients treated with
chemotherapy should be followed-up with medical history, physical
examination and appropriate tumour markers in the same way as dys-
germinomas. CT scans should be performed as clinically indicated.

Relapses in patients usually occur within the first two years after
diagnosis* (4,5). **Level of Evidence D.**

6.5.6 Management of sarcoma of the ovary
Primary sarcoma of the ovary is a rare entity that occurs primarily in
post-menopausal patients* (49). Nevertheless, an accurate diagnosis and
differentiation from other types of primary ovarian cancer is impor-
tant, as the prognosis is generally poor. There are two types of
sarcoma. Mixed Mullerian tumours (MMTs), the most common of the
two, are defined by the presence of both carcinomatous and sarcoma-
tous components* (46). Pure sarcomas are rarer. They can be categorised
as stromal cell sarcomas, fibrosarcomas, leiomyosarcomas, neurofi-
brosarcomas, rhabdomyosarcomas, chondrosarcomas, angiosarcomas,
and liposarcomas.

Patients with early stage disease have survival advantage compared
to those with advanced stage disease. The role of histological sub-
type and residual disease as prognostic indicators are controversial,
although many studies suggest that larger volume of residual disease
is associated with shorter survival. Different chemotherapeutic
regimes have been reported with wide ranging response rates.
Platinum-based regimes seem to have the better outcome compared to
non-platinum regimes. The overall prognosis remains poor. The rarity
of this entity prohibits any prospective randomised trials to test the
efficacy of any treatment strategy.

Despite the absence of more concrete data, it is still recommended
that patients with ovarian sarcomas undergo complete surgical stag-
ing or optimal cytoreduction if metastatic disease is present* (50). In
the absence of a trial of adjuvant chemotherapy, patients should receive
platinum-based chemotherapy post-operatively* (35-57).

The follow-up schedule is recommended as for EOC. However, the
benefit of surveillance with CA 125 is unknown. **Level of Evidence C.**

6.5.7 Management of primary lymphoma of the ovary (POL)
Primary lymphoma of the ovary is also rare. The majority of lym-
phomas that involve the ovary usually do so as part of a systemic
disease. Therefore, when the diagnosis of ovarian lymphoma is made
it is pertinent to exclude systemic disease. Fox and Langlely had pro-
posed a three-point diagnostic criteria making this entity even rarer.
The criteria are as follows:

- At the time of the diagnosis, the lymphoma is clinically confined
to the ovary and full investigations fail to reveal evidence of lym-
phoma elsewhere. A lymphoma can still be considered as ovarian
primary if spread has occurred to immediately adjacent lymph
nodes or if there has been direct spread to infiltrate immediately
adjacent structures.
- The peripheral blood and bone marrow should not contain any
abnormal cells.
- If further lymphomatous lesions occur at sites remote from the
ovary then at least several months should have lapsed between the
appearance of ovarian and extraovarian lesions.

Recent studies have shown that benign lymphoid tissue can be found
in up to 50% of normal ovaries. It is therefore possible that POL may
have arisen from malignant transformation of these benign aggre-
gates. POL has a propensity to metastasize to the contralateral ovary
and intraperitoneally. Diagnosis requires excisional or incisional
biopsy following by appropriate fixation to allow accurate sub-classi-
fication. Fine needle biopsies of masses will not be adequate for this
purpose. Diffuse large cell non-Hodgkin’s lymphoma has been
reported to be the commonest sub-type although the rarity of POL as
a whole prohibits a detailed understanding of the significance of the
sub-classifications.

One entity that demands greater attention is ovarian involvement in
Burkitts lymphoma in countries where Burkitts disease is endemic.
Enlargement of one or both ovaries is the second most common form
of presentation after involvement of the jaw.

Treatment of POL after surgical removal is no different from treat-
References


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Trophoblastic Disease

7.1 Introduction

Before 1969 metastatic choriocarcinoma was nearly invariably fatal, whereas most patients are now cured and usually retain reproductive function. The basis for this dramatic change is earlier diagnosis, the ability to precisely measure human chorionic gonadotrophin (hCG) and the availability of effective chemotherapy. Trophoblastic disease needs to be treated by, or at least in consultation with, physicians experienced in the management of this disease spectrum. The morbidity and mortality is 9 times higher when the inexperienced physician treats such patients.

Precise follow-up of patients and precise monitoring using a reliable assay of hCG is essential to good results. The protocol presented here will help outline these principles.

7.1.1 Definitions:
The term Gestational Trophoblastic Neoplasia (GTN) replaces the terms chorioadenoma destruens, metastasizing mole and choriocarcinoma. These were pathologic diagnoses. While histologic verification is desirable, it is not essential for the clinical classification we now use. Hydatidiform mole is gestational trophoblastic disease GTD). Nine to twenty per cent of patients with complete hydatidiform mole go on to have gestational trophoblastic neoplasia. This may be chemical only or associated with evidence of invasive mole. If the process is confined to the uterus, it is termed non-metastatic trophoblastic neoplasia. If metastases are demonstrated in the lung or vagina and/or in brain, liver, kidney or elsewhere, the diagnosis is that of metastatic gestational trophoblastic neoplasia. Besides post-molar trophoblastic neoplasia, trophoblastic tumour may also follow abortion (30%) or normal pregnancy (20%). Placental site trophoblastic tumour (PSTT) is a variant of GTD but should be classified separately as it has a distinct clinical presentation and its course and management differs from GTD. Non-gestational trophoblastic disease is choriocarcinoma of the ovary or testis.

7.1.2 Aetiology of hydatidiform mole

Our perspectives of the pathophysiology of hydatidiform mole and trophoblastic neoplasia have also become significantly focused. The distinction between complete mole and partial mole has been validated by genetic analysis and DNA fingerprinting. Complete mole occurs in one in 1500 pregnancies in the USA but still in 1 in 400 pregnancies in Korea, Indonesia and among Native Americans in the United States! With complete mole, the chromosomal material from the ovum is lost and the genetic material in the conceptus is paternally derived. Fertilization of this “empty” ovum by one sperm results in a 46XX androgenic conceptus. Fertilization may be by 2 sperms giving an XX or XY androgenic conceptus. A YY fertilization will not develop beyond a few cell embryo. With complete mole no foetus develops from this androgenic fertilization. The placenta develops hydatidiform changes and trophoblastic hyperplasia, resulting in a mole with a 9% - 20% chance of subsequent neoplasia.

Partial mole is being increasingly recognized as a cause of foetal loss. Many first trimester abortions are associated with triploidy and are in fact partial moles when examined histologically and confirmed by flow cytometry. Clinically there is high hCG, a foetus is present that is abnormal and there are hydatidiform changes in the placenta. Early pregnancy pre-eclampsia may occur and there is usually less trophoblastic hyperplasia than with a complete androgenic mole. The incidence of trophoblastic sequelae is 4%.

It should be noted that hydatidiform degeneration is a sign of a poorly functioning early placenta and may occur with spontaneous abortion without trophoblastic hyperplasia.

The diagnosis of metastatic gestational trophoblastic neoplasia in a patient who has not had a hydatidiform mole may be difficult. All physicians need to be aware that any female patient of reproductive years who has an obscure disease may have choriocarcinoma. This applies particularly to women presenting with cerebrovascular accidents or persistent pneumonia. All such patients need to have an hCG.

7.2 Diagnosis, evacuation and follow-up after evacuation of hydatidiform mole

Ultrasound examination of the first trimester uterus and particularly vaginal colour Doppler flow ultrasound has made possible the detection of abnormalities of early pregnancy. The diagnosis of hydatidiform mole is nearly always made by ultrasound. After mole evacuation, patients are followed closely with weekly hCG. The diag-
nosis of gestational trophoblastic neoplasia is made on the basis of an elevated hCG plateau or rising hCG titers over a period of several weeks. Histologic choriocarcinoma and/or the appearance of metastases with a persistently raised serum hCG level is an absolute indication for chemotherapy.

Physical examination and investigations such as chest x-ray, ultrasound, CT scanning, or MRI of the brain, chest, abdomen and pelvis particularize the type of disease present. Gestational trophoblastic neoplasia responds excellently to chemotherapy and even with extensive high risk metastatic disease a mortality of 90% has been converted to a cure rate of 92% or better.

Hydatidiform mole should be treated by evacuating the uterus. The patient must then be followed by serial weekly hCG titers to undetectable levels. In the 20% of patients in whom the levels of hCG remain elevated, several courses of chemotherapy may be required. In general, patients are no longer treated for moles with prophylactic chemotherapy as this exposes 80% to unnecessary chemotherapy. Prophylactic or rather adjuvant chemotherapy (i.e. not one course, but several courses to non-detectable hCG) should only be offered to patients who cannot be followed.

7.2.1 hCG Assays, nicked hCG, phantom hCG
A reliable assay for total hCG is central to the management of patients with trophoblastic disease. The assay must measure all portions of the hCG molecule particularly free beta subunit, hyperglycosylated hCG (hCG-H), nicked hCG and hCG missing the terminal carboxyl segment. These products are more common in neoplasia than is total hCG. Several commercial assay kits do not measure free beta subunit or nicked hCG or differentially recognised hyperglycosylated hCG. Physicians treating patients with trophoblastic disease must ensure that the laboratory used provides accurate assay results, otherwise false low values may result in inappropriate management.

In the last few years we have also encountered patients who have positive serum pregnancy tests but have no trophoblastic disease and in fact no pregnancy associated events. These patients’ serum contains heterophilic antibody that reacts with the antibody of certain assay kits to give (false) positive hCG results. This is called phantom hCG. It is important for physicians to be aware of this problem because such patients do not require any therapy. The appended bibliography contains relevant references to false positive hCG.

The presence of real hCG may be easily confirmed and phantom hCG ruled out by performing hCG assays in dilution that should provide proportionate results or by demonstrating the similarity presence of hCG in serum and in urine. However hCG of less than 50mIU/ml in urine is inaccurate. When there is doubt the US hCG Reference Laboratory should be consulted (larry@hcglab.com).

7.2.2 Quiescent Gestational Trophoblastic Disease
In the last five years clinicians have encountered patients with elevated hCG usually about 50 to 100 mIU/ml that follows an episode of molar pregnancy or occasionally is discovered incidentally. This is real hCG and not false positive hCG. The patients have no abnormal clinical findings and imaging is negative. The hCG persists in spite of treatment with chemotherapy or surgery. In some 20% of these patients the hCG becomes re-elevated after a period of several weeks to several years and overt tumour becomes detectable. During the quiescent period the patient has no detectable hyperglycosylated hCG (hCG-H) but as soon as the hCG rises a significant proportion is hCG-H frequently prior to the appearance of clinically detectable neoplasia by examination or imaging. At this time therapy is effective. Two articles describing details of this syndrome are cited in the bibliography section.

7.2.3 Pathology
The histologic diagnosis of both complete and partial hydatidiform mole is well recognized. If there is doubt in distinguishing between partial mole and complete mole, flow cytometry may be helpful and is now a well established technique. The major current difficulty is the recognition of tissue obtained from the uterus at 4 to 8 weeks gestational age as hydatidiform mole. The criteria for making the diagnosis of these early moles, made possible by the earlier ultrasound diagnosis of the hydatidiform mole, have been extensively described by Paradinas. It is important to realize that the classical features may not yet be present and that foetal membrane and even foetal erythrocytes may still be present.

The histologic diagnosis of placental site tumour may also be diffi-
cult on curettage material and may require the expertise of a gynaecologic pathologist with extensive experience in this refined area of gynaecologic pathology. This becomes particularly important when hysterectomy is indicated in young nulliparous women in whom this diagnosis is believed to be present.

7.3 Detailed discussion of trophoblastic disease management

7.3.1 Hydatidiform mole

7.3.1.1 Diagnosis of hydatidiform mole
1. History
2. Clinical examination
3. Ultrasound examination preferably with vaginal colour doppler flow ultrasound
4. Radiologic examination by magnetic resonance imaging or CT scan is indicated only when ultrasound examination is inconclusive.
5. Serum hCG levels are helpful.
Note: Bleeding or excess vomiting in the first trimester merits ultrasound examination to allow a positive diagnosis of mole, multiple pregnancy or foetal abnormality. “No foetal heart or high hCG above 80,000 mIU/ml equals mole.”

7.3.1.2 Required studies for patients with hydatidiform mole
1. Clinical examination including neurological examination, eye fundus examination and blood pressure.
2. Chest x-ray
3. Blood count with platelet count, BUN, creatinine and liver function tests on admission. Blood group and hold clot. Thyroid function tests may be indicated. Clinical thyrotoxicosis is very rare. PT, PTT, prothrombin, fibrinogen, if clinically indicated,
4. Serum hCG immunoassay. A specimen of serum for hCG should be obtained (1) prior to and (2) one day after the evacuation of the mole before the patient leaves the hospital.
5. Digital oximetry, blood gases, and lung scan are mandatory if there is suspicion of pulmonary embolization or pulmonary metastases which are not demonstrable by chest x-ray.

7.3.1.3 Management of hydatidiform mole
The hydatidiform mole is surgically evacuated as soon as possible after diagnosis. If necessary the patient is stabilised after the diagnosis is established. If haematologic, thyroid or pulmonary problems are present these are treated: the essential principle is mole evacuation. Evacuation should be done by suction curettage with accompanying syntocinon infusion plus ergonovine if necessary. The cervix may be dilated gently and slowly. With complete mole, a 9 mm or 10 mm suction curettage usually suffices and greater dilatation of the cervix is usually not necessary. A careful, “light” sharp curettage should be performed following the suction procedure to ensure the uterus has been completely evacuated. Hydatidiform moles of gestational size greater than 16 weeks should be evacuated at the Trophoblast Centre because of the risk of pulmonary embolization of molar tissue.

Hysterectomy may be done in patients who have finished childbearing.

If the patient wishes to retain her uterus she should be allowed to do so as hysterectomy does not improve the prognosis. RHOGAM SHOULD BE GIVEN IF INDICATED.

7.3.1.4 Management post-evacuation
1. The patient is followed by weekly hCG measurements until hCG becomes undetectable. Anaemia or infection is treated if present. When the hCG becomes undetectable, two further specimens are obtained at weekly intervals. Subsequently, the patient is tested monthly for six months and then at two monthly intervals for a further six months to insure that the hCG levels remain undetectable.
2. AN ASSAY FOR hCG SENSITIVE TO 2 mIU/ml OR LESS, IS ESSENTIAL FOR FOLLOW-UP. The assay must be able to detect all portions of the hCG molecule.
3. The patient is given reliable contraception, preferably in the form of the pill. If the fall in hCG was logarithmic the patient may be allowed to become pregnant after 6 months of follow-up. If there is very slow fall in hCG post-mole, a longer wait is indicated. It is useful to obtain an ultrasound scan early during the subsequent pregnancy and to follow hCG early in that next pregnancy to doc-
7.3.2 Gestational trophoblastic neoplasia

Principal

Gestational trophoblastic neoplasia (GTN) follows hydatidiform mole (60%), previous miscarriage/abortion (30%), normal pregnancy or ectopic gestation (10%). GTN most commonly follows hydatidiform mole as a persistently elevated hCG titer. There may also be continuing and recurring bleeding after mole. Metastatic gestational trophoblastic neoplasia will frequently manifest itself by symptoms from the metastases, such as intracranial neoplasia or "pneumonia."

7.3.2.1 Diagnosis of post-molar GTN

The diagnosis of gestational trophoblastic neoplasia is made on the basis of elevated hCG levels supported, if possible, but not necessarily, by histologic or radiologic evidence. If, after mole, hCG remains elevated, falls and becomes elevated again, therapy is instituted. Management should only be undertaken by an experienced team: mortality outside a trophoblastic centre is much greater than morbidity in a centre.

7.3.2.2 Management of gestational trophoblastic neoplasia

Required studies

1. Clinical examination: (watch for the vaginal metastasis)
2. Serial weekly hCG measurements on serum
3. Complete blood count and platelets. PT, PTT, fibrinogen, BUN, creatinine, liver function tests.
4. Chest x-ray
5. Brain, MR (or CT) scan when there is any suspicion of cerebral metastases.
6. Liver CT scans when indicated. A whole body CT scan is normally done in patients who have lung metastases.
7. Curettage should be done if there is uterine bleeding. Biopsies may be obtained from accessible sites. THERE IS GRAVE DANGER OF HAEMORRHAGE AT THE BIOPSY SITE.
8. Magnetic resonance imaging when indicated.
9. T4, thyroid studies when indicated.
10. Selective scanning using anti hCG antibody linked to radioactive iodine or indium may be done if there is persistent chemotherapy resistant disease.

7.3.2.3 Staging

7.3.2.3.1 The Staging of gestational trophoblastic neoplasia by the International Federation of Gynecology and Obstetrics (FIGO).

In 2000 FIGO recommended a clinical staging of gestational trophoblastic tumours and requested that such cases should be reported to the Annual Report on the Results of Treatment in Gynecological Cancers. For this purpose the definitions of the clinical stages of gestational trophoblastic tumours are:

**FIGO Staging**

Stage I Gestational trophoblastic tumours strictly confined to the uterine corpus.

Stage II Gestational trophoblastic tumours extending to the adnexa or to the vagina, but limited to the genital structures.

Stage III Gestational trophoblastic tumours extending to the lungs, with or without genital tract involvement.

Stage IV All other metastatic sites.

According to FIGO, hydatidiform mole should be registered but not be staged as Stage 0 because if hCG persists and the patient requires chemotherapy, restaging would be required. Such staging transgresses the present rules of the present FIGO staging system. Patients with hydatidiform mole are placed on record but staging only applies to trophoblastic neoplasia.

Cases which do not fulfill the criteria for any given stage should be listed separately as unstaged. It should be realized that most cases of low risk metastatic disease are comprised by Stage 3, while the high risk group of metastatic tumours first described by Hammond is the group that comes under Stage 4.

7.3.2.3.2 A Modified WHO scoring system has been combined with the FIGO staging.

In 2000 FIGO accepted the World Health Organization scoring sys-
The agreed criteria to diagnose Gestational Trophoblastic Neoplasia (GTN) include:
1) At least 4 values of persistently elevated hCG plateau (days 1,7,14 and 21) or longer or sequential rise of hCG for two weeks (days 1,7,14) or longer. The actual values of hCG are left to the discretion of individual physicians.
2) Lung metastases are diagnosed by chest x-ray.

Treatment of gestational trophoblastic neoplasia (trophoblastic tumour)

Low risk GTN
Gestational non-metastatic trophoblastic neoplasia, low risk metastatic neoplasia (lung metastases only) in patients who have gestational trophoblastic neoplasia for less than 4 months and whose hCG value is less than 40,000 mIU/ml hCG. WHO score 6 or less.

FIGO Stage I, II, and III.
1. Drug schedules: single agent chemotherapy:
   (a) Methotrexate 0.4 mg/kg IM for 5 days, repeated every 2 weeks. This is one of the original protocols used in GTD and is still used at Yale University. It is the standard protocol at the Brewer Trophoblast Centre in Chicago. It is associated with a 10% primary failure rate.
   (b) Methotrexate with Leucovorin rescue (Table 1). Methotrexate 1.0 mg/kg im every other day for 4 doses with leucovorin 0.1 mg/kg 24 hours after each dose of Methotrexate. This is a widely used protocol in the United Kingdom and the United States but has a 20% - 25% primary failure rate.
   (c) Methotrexate 50 mg/m² IM given weekly. This regimen is associated with a 30% primary failure rate. If this occurs Methotrexate 0.4 mg/kg IM for 5 days may be administered or the medication may be changed to Actinomycin-D 12 mcg/kg for 5 days.
   (d) Actinomycin-D, 1.25 mg/m² given every 2 weeks. This protocol carries a 20% primary failure rate. It is an alternative to the pulsed weekly methotrexate protocol (c).
   (e) Actinomycin-D, 12 micrograms/kg IV daily for 5 days, repeated every 2 weeks. This protocol is an alternative to the 5-day
MTX protocol. It may be used with patients who have hepatic dysfunction. It carries an 8% primary failure rate.

(f) Methotrexate 250 mg infusion over 12 hours. This is the MTX portion of the EMA-CO protocol. It is associated with a 30% primary failure rate.

NOTE: ACTINOMYCIN-D CAUSES SEVERE SLOUGH IF INFILTRATED AND MUST BE INJECTED VIA A NEW FREE RUNNING INTRAVENOUS INFUSION. IF ANY EXTRAVASATION DOES OCCUR, THE AREA SHOULD BE INFILTRATED WITH 100 mg HYDROCORTISONE AND 2 cc OF 1% XYLOCAINE.

The primary failure rate of the “pulsed” regimens is significantly greater than that of single agent 5-day courses because of insufficient drug exposure to cells in the S phase of replication. For example, the primary failure rate of the 5-day Actinomycin course is 8% compared with 20% with the pulsed Actinomycin-D 1.25 mg/m².

2. Repeat complete blood count, platelets, creatinine, BUN and SGOT are obtained on the first day of each course.

3. At least one course, and usually two to three courses of chemotherapy should be given beyond first negative hCG level particularly if the fall of hCG is slow or there has been extensive disease.

7.4 Chemotherapy “for the road”

Three further courses of chemotherapy, at least the first of which will be combination therapy, are given beyond the first non-detectable hCG value. A negative hCG value implies that the number of malignant cells present in the body is less than 10⁷. It does not mean the disease at that time is completely eradicated.

Certain metastatic sites may require special therapy. For example, brain lesions are treated with an increased dose of Methotrexate to 1 gm/m² in the EMA-CO protocol. With this relatively high dose of methotrexate the urine must be maintained alkaline. Depending on the size and number of brain metastases patients may be treated with 25 to 30 GY whole brain irradiation or excisional alkaline. Patients with liver metastases may be treated with 20 GY liver radiation or hepatic artery infusion. Both with brain and liver metastases, the radiation serves to prevent catastrophic haemorrhage more than controlling the trophoblastic disease.

Patients resistant to EMA-CO or recurring after previous multiagent chemotherapy may be treated by the EMA-EP (EP-EMA) protocol: this is EMA alternating with etoposide and platinum. On occasion EMA-PA (P=cis-platinum, A=adriamycin) may be used.

For EMA-EP resistant cases Taxol with Cisplatin alternating with Taxol-Etoposide or Taxol-5-FU or ifosphamide, cisplatinium etoposide (ICE) or vinblastine etoposide cisplatin (BEP) have been used.

7.4.1 Surgery for chemotherapy resistant and persistent metastases

Metastases to lung, liver, brain or other sites that do not regress with chemotherapy may be amenable to surgical extirpation.

7.4.2 Pregnancy after metastatic trophoblastic disease

Patients need to wait for twelve months after ceasing chemotherapy before undertaking pregnancy.

7.4.3 Placental site trophoblastic tumour

1. Placental site trophoblastic tumour should be separated from gestational trophoblastic tumour such as hydatidiform mole and
choriocarcinoma. It should be treated by a trophoblastic centre. Tumour load with Placental Site Trophoblastic Tumour is not reflected by hCG and while hPL may be seen immunohistochemically it is rarely detectable in serum. Patient management with both chemotherapy and surgery requires individualisation. It has recently been discovered that free beta hCG is a reliable marker for placental site trophoblastic tumour, especially in situations when there is uncertainty whether the patient has choriocarcinoma (high proportion hCG-H) or placental site trophoblastic tumour (high proportion free beta hCG). See bibliography.

Figure 1: Guidelines for the management of Gestational Trophoblastic Disease

<table>
<thead>
<tr>
<th>Hydatidiform Mole</th>
<th>Initial Assessment</th>
</tr>
</thead>
<tbody>
<tr>
<td>OBGYN History</td>
<td>CBC, Coagulation studies</td>
</tr>
<tr>
<td>Physical Examination</td>
<td>when indicated</td>
</tr>
<tr>
<td>Chest X-Ray</td>
<td>BUN, Creatinine, LFT</td>
</tr>
<tr>
<td>hCG</td>
<td></td>
</tr>
</tbody>
</table>

**Mole Evacuation**

- Complete Mole
  - Suction Evacuation Uterus
  - Laminaria: optional
- Partial Mole
  - Laminaria: optional
  - Dilatation & Evacuation D&C
- No Metastases

- WEEKLY hCG [plot graph on semi-log paper]

- Spontaneous Resolution
  - Follow for 6 to 12 months before permitting pregnancy

Note: Gestational Trophoblastic Disease (GTD) includes all aspects of this entity. The term Gestational Trophoblastic Neoplasia (GTN) is reserved for entities that require chemotherapy and replaces invasive mole, malignant GTD and other such terms. Choriocarcinoma is a pathologic term; PSTT is classified separately.

Figure 2: Guidelines for the Management of Trophoblastic Neoplasia

- Diagnosis of GTN
  - hCG, CBC, plateletes, BUN creatinine, liver function tests, coagulation studies (if indicated), chest X-ray, pelvic ultrasound. If chest X-ray is positive -> CT/USG scan abdomen, particularly liver CT or MRI brain as indicated.

- Stage I
  - Risk Factor ≤ 6
  - Single agent Chemotherapy
  - Investigation, staging and risk factor scoring
  - Diagnosis of Gestational Trophoblastic Neoplasia (GTN)
  - NO RESPONSE

- Stage II
  - Risk Factor ≤ 6
  - Change single agent chemotherapy scheduling or agent (if pulsed Act-D or MTX change to 5-day course; if one agent 5-day course fails change to alternate MTX or Act-D 5-day course)
  - NO RESPONSE

- Stage III
  - Risk Factor ≤ 6
  - Combination chemotherapy (consider TAH with uterine lesion)
  - NO RESPONSE

- Stage IV and/or
  - Risk Factor > 7
  - see Figure 3
  - NO RESPONSE

Go to Figure 3
Figure 3: Guidelines for the management of Trophoblastic Neoplasia

Post-hydadiform mole
Stage IV or risk factor ≥ 7
OR
Non-molar GTN diagnosed from metastases

Investigation, staging and risk factor scoring

hCG, CBC, platelets, BUN creatinine, liver function tests, coagulation studies (if indicated), chest X-ray, pelvic ultrasound. If chest X-ray is positive -> CT/USG scan abdomen, particularly liver CT or MRI brain as indicated.

Stage I, II, III with risk factor ≥ 7 or Stage IV

Multiple agent chemotherapy (eg EMA-CO)
[for cerebral metastases MTX dose is increased to 1g/m²]

Resolution

Follow with hCG and clinical surveillance for one year

Persistent neoplasia

Consider surgery for isolated resectable lesions (usually lung, brain or liver)

Second line multiple agent chemotherapy (EP-EMA)

CONSULT Trophoblast Centre

Figure 4: Guidelines for the management of Trophoblastic Disease

Placental Site Trophoblastic Tumour (PSTT)
(diagnosis based on expert histology)

Uterus only strongly consider TAH

Metastases EMA-CO or EP-EMA chemotherapy consider TAH if there is tumour in uterus

Appendix

Chemotherapy Protocols

Single-agent chemotherapy protocols for low risk trophoblastic neoplasia. WHO Score 6 or less.
1. Methotrexate 0.4 mg/kg daily for 5 days given IM.

<table>
<thead>
<tr>
<th>DAY</th>
<th>THERAPY</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>CBC, PLATELET COUNT, SGOT MTX, 1 mg/kg, IM</td>
</tr>
<tr>
<td>2</td>
<td>CF, 0.1 mg/kg, IM</td>
</tr>
<tr>
<td>3</td>
<td>MTX, 1 mg/kg, IM</td>
</tr>
<tr>
<td>4</td>
<td>CF, 0.1 mg/kg, IM</td>
</tr>
<tr>
<td>5</td>
<td>MTX, 1 mg/kg, IM</td>
</tr>
<tr>
<td>6</td>
<td>CF, 0.1 mg/kg, IM</td>
</tr>
<tr>
<td>7</td>
<td>MTX, 1 mg/kg, IM</td>
</tr>
<tr>
<td>8</td>
<td>CF, 0.1 mg/kg, IM</td>
</tr>
</tbody>
</table>

CBC = COMPLETE BLOOD COUNT
MTX = METHOTREXATE
CF = CITROVORUM FACTOR RESCUE

3. Methotrexate 50 mg/kg given weekly by IM injection.
4. Actinomycin 12 mcg/kg daily for 15 days.
5. Act-D 1.25 mg/m² given every two weeks. (Pulsed Act D)
6. Methotrexate 250 mg infusion over 12 hours.

Suggestions for the management of primary failure of single agent chemotherapy in Low Risk GTN (Risk Factor Score 6)

If pulsed single agent chemotherapy, either Methotrexate 50mg/m² or

7.5 Trophoblastic patient record

Physicians treating trophoblastic disease are urged to maintain an active record of weekly hCG measurements on semi-log graph paper. Treatment events such as chemotherapy and radiologic investigations are recorded on the same record.
Actinomycin 1.25 mg/m² or Methotrexate with leucovorin rescue do not effect response, it may be worthwhile to use the same agent given as a 5-day course; ie Methotrexate 0.4 mg/kg daily for 5 days or Actinomycin 12 microg/kg daily for 5 days, before switching to the alternate agent. The failure of pulsed single agent chemotherapy is thought to be associated with insufficient time of exposure of cells in cycle during the relatively brief time effective levels are present in the circulation. This practice may avoid having to give multi agent chemotherapy in such a situation in more then 50% of these patients.

Current multi-agent chemotherapy for high-risk GTN, EMA-CO WHO Score 7 or greater
Etoposide (VP-16), Methotrexate, Actinomycin D, alternating weekly with Cyclophosphamide and Oncovin (vincristine)
EMA-CO is administered on a weekly basis with anticipated cycling between each course of 14 days.
Day 1 (A) Actinomycin D 500 micrograms IV push new IV. Etoposide 100 mg/m² over 30-50 minutes Methotrexate 100 mg/m² IV infusion over 1 hour and then Methotrexate 200 mg/m² IV infusion over 12 hours by pump.
Day 2 (A) Actinomycin D 500 micrograms IV push new IV Etoposide 100 mg/m² over 30-50 minutes Folinic Acid 15mg IV push Q 6 hours for 8 doses beginning 24 hours after Methotrexate bolus. Some physicians administer the folinic acid Q 12 hours for 4 doses orally 15 mg commencing 24 hours after commencing methotrexate.
Day 8 (B) Vincristine (Oncovin) 1 mg/m² IV Cyclophosphamide 600 mg/m² IV.

NOTE:
1. Neupogen may be administered. Note that this must be started 24 hours after day 2 chemotherapy and then be stopped 24 hours before CO.
2. If the creatinine is greater than 2.0, creatinine clearance should be done prior to therapy and should be 50 or more.
3. Cycles are repeated on day 15 of cycle.

4. Chemotherapy is administered when WBC is greater than 3000 per cc. Granulocytes are greater than 1500 per cc. Platelets are greater than 100,000 and a Grade 3 gastrointestinal infection and mucositis morbidity has cleared. If toxicity necessitates a delay in course B for longer than 6 days, course A is recycled.

Combination chemotherapy for trophoblastic neoplasia with brain metastases – EMA-CO with high dose methotrexate
For GTN with brain metastases the EMA-CO protocol is modified. The dose of MTX is increased to 1000 mg/m² (one gm/m²). The MTX infusion is given over 24 hours. The urine must be kept alkaline with a measured pH of greater than 7.5 at all times by administration of bicarbonate IV. Urinary volume and pH must be followed assiduously.
If Neupogen is given, it needs to be given commencing 24 hours after the last chemotherapy and ceasing 24 hours prior to the next planned chemotherapy.

Combination chemotherapy for high risk trophoblastic neoplasia resistant to EMA-CO or recurring after combination chemotherapy
The majority of trophoblast centres use EP-EMA (EMA-CO) under such circumstances. EMA is administered in the standard way and etoposide and platinum are substituted for CO. This is a more demanding and more toxic regimen. The following schedule is taken from the Charing Cross protocol as they have had the most experience and the greatest success.

Combination chemotherapy for EMA-CO failure in trophoblastic neoplasia: EP-EMA
Cisplatin is given on Day 1 by infusion of 80 mg/kg in 1 litre by infusion pump over 12 hours. Etoposide 100 mg/m² is given over 1 hour. Day 8 – EMA is given in standard doses but the second day Actinomycin-D and etoposide are omitted. The cycle is repeated every 15 days.
Neupogen is commenced 24 hours after first methotrexate infusion and stopped 24 hours before day 8 platinum. It is recommenced 24 hours after stopping platinum and stopped 24 hours before next EMA. The timing of Neupogen has to be carefully planned.
Alternate combination chemotherapy for high risk GTN: Methotrexate, Actinomycin, Cyclophosphamide (MAC) protocol for high risk GTN. This regimen has been largely superseded by EMA-CO

<table>
<thead>
<tr>
<th>DAY</th>
<th>THERAPY</th>
</tr>
</thead>
</table>
| Day 1 | CBC, platelet count, SGOT  
Compazine, 25 mg, IM, PO or PR  
MTX, 1.0 mg/kg, IM  
ACT-D, 12 mcg/kg, stat IV  
Cyclophosphamide, 3 mg/kg, stat IV |
| Day 2 | Compazine, 25 mg, IM, PO or PR  
Leucovorin, 0.1 mgm/kg, IM  
ACT-D, 12 mcg/kg, stat IV  
Cyclophosphamide, 3 mg/kg, stat IV |
| Day 3, 4, 5 | repeat day 1 & 2 |
| Day 6 | CF, 0.1 mg/kg, IM |
| Day 7 | CBC, platelet count, SGOT  
MTX, 1.0 mg/kg, IM |
| Day 8 | CF, 0.1 mg/kg, IM |

Courses are repeated every 2 weeks or as soon as white cells and platelet recover.

Bagshawe 9-day multi-agent chemotherapy
This protocol has been reported to be more toxic than MAC and will probably be used only in exceptional circumstances. However, several trophoblast centres use it for high-risk disease as initial therapy because of the leucemogenic association of EMA-CO.

Bleomycin, Etoposide, Cisplatin (BEP) for chemotherapy resistant gestational trophoblastic neoplasia and for primary ovarian germ cell tumour

Etoposide 100 mg IV in 500 ml N saline over 1 hour Days 1, 2, 3, 4.

Cisplatin 100 mg/m² day one IV continuous infusion x 24 hours with NSx6 at 250 ccs/hr; add 20 mEgn KCl and 2 mg MgSO4 to each of last 2 litres.

Bleomycin 10 units/m² per day for 3 days; days 2, 3 and 4; IV continuous infusion for 96 hours.

Give day 1, VP-16 prior to cisplatin. Give day 2, VP-16 as part of cisplatin post-hydration; then complete post-hydration (concurrent with 1st bleomycin infusion) with D5/NS at 150 ml/hr for 11 hours. Give day 3, and 4 VP-16 prior to starting 2nd and 3rd Bleomycin infusion. Kytril, Compazine and Benadryl are given with these medications.

Validity of studies on which the protocols are based
In an age of evidence-based medicine very few of the chemotherapy protocols used to treat trophoblastic disease have undergone the rigors of a prospective randomised study. No study fulfills a Cochrane category I or II for clinical studies. The evidence on which management of Trophoblastic Neoplasia (GTN) is presently based throughout the world fulfills only a Cochrane category III. This applies to all the management protocol in this document.

Consultation
Physicians wishing for consultation concerning case management should contact their nearest Trophoblast Centre early rather than late following mole evacuation. A list of telephone numbers and e-mail addresses will be found on the web at: www.isstd.org. The bibliography provides an overview of the present situation. Those wishing to have a more detailed outline may get in touch with one of the Trophoblast Centres.

Selected Bibliography

*Comprehensive review of GTD*

*Definitions*

*hCG Measurements*
Cole LA. Use of hCG tests for evaluating Trophoblastic Disease in...


Imaging

Classification and Staging


Quiescent gestational trophoblastic disease


Pathology


Placental site trophoblastic tumours

Interesting papers


Examples of FIGO 2002 staging: scoring

1. Low stage: low score
   Forty-five year old patient requiring chemotherapy 6 weeks after hydatidiform mole. The hCG was 900 milli IU/ml. There were no metastases. The FIGO Stage Score is I: 1.

2. High stage: high score
   40 year old patient, 7 months after full term pregnancy who presents with lung (4 Mets), brain (1 met; 5 cm size), liver (2 mets) and kidney (one met each) metastases with an hCG of 42,000 milli IU/ml. This patient is FIGO Stage IV: 18.

3. High stage: low score
   20 year old patient, who presents 8 weeks post-hydatidiform mole with one metastasis to the lung and one to one kidney (4 cm in size) with an hCG of 800 mIU/ml. The FIGO Stage Score is IV: 4.

4. Low stage: high score
   Patient, aged 44, is 8 months post miscarriage, has uterine bleeding and by ultrasound is found to have a 9 cm mass in the uterus. D & C shows histologic choriocarcinoma. The hCG is 18000 milli IU/ml and there is also a 5 cm nodule in the vagina. Single agent chemotherapy with methotrexate has failed. There are no other metastases. The FIGO Stage Score for this patient is II: 10.

Acknowledgements:

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USA

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Hong Kong SAR
China

S Sasaki
Nippon Medical School
Tokyo
Japan

J Soper
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Duke University Medical Center
Durham, North Carolina
USA
Cancer of the Breast

8.1 Staging

8.1.1 Anatomy

8.1.1.1 Primary site
The breast is situated on the anterior chest wall and is composed of glandular tissue with a dense fibroareolar stroma. The glandular tissue consists of approximately twenty lobes, each of which terminates in a separate excretory duct in the nipple.

8.1.1.2 Regional lymph nodes
Lymphatic drainage of the breast is via three major pathways: axillary, transpectoral and internal mammary. The intramammary lymph nodes are considered with the axillary lymph nodes for staging purposes. Metastases to other lymph nodes, including supraclavicular cervical and contralateral internal mammary nodes are considered distant metastases (M1).

1) Axillary
Interpectoral nodes and lymph nodes along the axillary vein and its tributaries may be divided into the following levels:
   i) Level I (low-axilla): Lymph nodes that are lateral to the lateral border of the pectoralis minor muscle.
   ii) Level II (mid-axilla): Lymph nodes which lie between the medial and the lateral borders of the pectoralis minor muscle and the interpectoral (Rotter’s) lymph nodes.
   iii) Level III (apical axilla): Lymph nodes medial to the medial margin of the pectoralis minor muscle, including those designated as subclavicular, infraclavicular, or apical.

2) Internal mammary
Internal mammary lymph nodes in the intercostal spaces along the edge of the sternum in the endothoracic fascia. Any other lymph node metastasis is coded as a distant metastasis (M1) and includes supraclavicular, cervical or contralateral internal mammary lymph nodes.

8.1.1.3 Metastatic sites
All distant visceral sites are potential sites for metastases. The major sites of involvement are bone, lung, brain and liver. Metastatic disease has been found in almost every remote site.

8.2 Rules for classification

8.2.1 Clinical staging
Clinical staging includes physical examination with careful inspection and palpation of the skin, mammary gland itself and lymph nodes (axillary, supraclavicular and cervical), as well as pathological examination of the breast or other tissues, and imaging to establish the actual diagnosis of breast cancer. The extent of tissues examined pathologically for clinical staging is less than that required for pathological staging. Appropriate operative findings are also elements of clinical staging including the size of the primary tumour and chest wall invasion and the presence or absence of regional or distant metastases.

8.2.2 Pathological staging
Pathological staging will include all of the data used for clinical staging and surgical resection as well as pathological examination of the primary carcinoma, including not less than excision of the primary cancer with no tumour in any margin of resection by gross pathological examination. The case may be included in the pathologic stage if there is only microscopic but not gross involvement at a resection margin. Lesions with a resection margin, on gross examination, are coded as TX because the extent of the primary tumour cannot be assessed. Resection of at least the lower axillary lymph nodes (level I) should be carried out. Such a resection ordinary will include six or more lymph nodes.

8.3 TMN Classification
The clinical measurement used to classify the primary tumour (T) should be the one that provides the most accurate information (e.g. physical examination or mammogram). Pathologically the tumour size for classification is a measurement of the invasive component. Lesions that show a large in situ component (e.g. 3-4 cm) and a small invasive component (e.g. 0.5 cm) are classified as T1a.
8.4 Definition of TMN

8.4.1 Primary tumour (T)
Definitions for classifying the primary tumour (T) are the same for clinical and for pathologic classification. The telescoping method of classification can be applied. If the measurement is made by physical examination, the examiner will use the major headings (T1, T2, or T3). If other measurements, such as mammographic or pathologic, are used, the examiner can use the telescoped subsets of T1. (See Table 1).

Table 1: Breast cancer pre-treatment clinical classification

<table>
<thead>
<tr>
<th>T</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>TX</td>
<td>Primary tumour cannot be assessed</td>
</tr>
<tr>
<td>T0</td>
<td>No evidence of primary tumour</td>
</tr>
<tr>
<td>Tis</td>
<td>Carcinoma in situ; intraductal carcinoma, lobular carcinoma in situ, or Paget's disease of the nipple with no tumour</td>
</tr>
<tr>
<td>T1</td>
<td>Tumour 2 cm or less in greatest dimension</td>
</tr>
<tr>
<td>T1a</td>
<td>0.5 cm or less in greatest dimension</td>
</tr>
<tr>
<td>T1b</td>
<td>More than 0.5 cm but not more than 1 cm in greatest dimension</td>
</tr>
<tr>
<td>T1c</td>
<td>More than 1 cm but not more than 2 cm in greatest dimension</td>
</tr>
<tr>
<td>T2</td>
<td>Tumour more than 2 cm but not more than 5 cm in greatest dimension</td>
</tr>
<tr>
<td>T3</td>
<td>Tumour more than 5 cm in greatest dimension</td>
</tr>
<tr>
<td>T4</td>
<td>Tumour of any size with direct extension to chest wall or skin</td>
</tr>
<tr>
<td>T4a</td>
<td>Extension to chest wall</td>
</tr>
<tr>
<td>T4b</td>
<td>Oedema (including peau d’orange) or ulceration of the skin of the breast or satellite skin nodules confined to the same breast</td>
</tr>
<tr>
<td>T4c</td>
<td>Both (T4a and T4b)</td>
</tr>
<tr>
<td>T4d</td>
<td>Inflammatory carcinoma (See the definition of inflammatory carcinoma in the introduction)</td>
</tr>
</tbody>
</table>

Note: Paget's disease associated with a tumour is classified according to the size of the tumour.

8.4.2 Regional lymph nodes (N)
NX Regional lymph nodes cannot be assessed (e.g., previously removed)
N0 No regional lymph node metastasis
N1 Metastasis to movable ipsilateral axillary lymph node(s)
N2 Metastasis to ipsilateral axillary lymph node(s) fixed to one another or to other structures
N3 Metastasis to ipsilateral internal mammary lymph node(s). (See Table 2).

Table 2: Breast cancer post-surgical classifications

<table>
<thead>
<tr>
<th>Pathologic classification (pN)</th>
</tr>
</thead>
<tbody>
<tr>
<td>pNX Regional lymph nodes cannot be assessed (e.g., previously removed, or not removed for pathologic study)</td>
</tr>
<tr>
<td>pN0 No regional lymph node metastasis</td>
</tr>
<tr>
<td>pN1 Metastasis to movable ipsilateral axillary lymph node(s)</td>
</tr>
<tr>
<td>pN1a Only micrometastasis (none larger than 0.2 cm)</td>
</tr>
<tr>
<td>pN1b Metastasis to lymph node(s), any larger than 0.2 cm</td>
</tr>
<tr>
<td>pN1bi Metastasis in one to three lymph nodes, any more than 0.2 cm and all less than 2 cm in greatest dimension</td>
</tr>
<tr>
<td>pN1bii Metastasis to four or more lymph nodes, any more than 0.2 cm and all less than 2 cm in greatest dimension</td>
</tr>
<tr>
<td>pN1biii Extension of tumour beyond the capsule of a lymph node metastasis less than 2 cm in greatest dimension</td>
</tr>
<tr>
<td>pN1biv Metastasis to a lymph node 2 cm or more in greatest dimension</td>
</tr>
<tr>
<td>pN2 Metastasis to ipsilateral axillary lymph nodes that are fixed to one another or to other structures</td>
</tr>
<tr>
<td>pN3 Metastasis to ipsilateral internal mammary lymph node(s)</td>
</tr>
</tbody>
</table>

8.4.3 Distant metastasis (M)
MX Presence of distant metastasis cannot be assessed
M0 No distant metastasis
M1 Distant metastasis [includes metastasis to ipsilateral supraclavicular lymph node(s)]

8.5 Introduction
Carcinoma of the female breast represents one of the most frequent malignant tumours in women, in both developing and industrialised nations. In recent years there has been an increased emphasis on the diagnosis of breast cancer through instructions on breast self-examination, regular physical examinations by the patient’s physician, and the use of mammography. This activity, coupled with extensive public education, is leading to earlier diagnosis of disease in a sizable proportion of women with breast cancer and should lead to better overall survival and prognosis.
One result of earlier diagnosis has been the increasing number of women seen with in situ disease. This has led to a reevaluation of both the risk to these individual women and the subsequent recommendations for management of in situ disease while still conserving the breast. Wide local excision, followed by radiation therapy, produces survival and local control, which is equivalent to that obtained with mastectomy so that breast conservation is now considered standard therapy for most women with operable breast cancer. Adjuvant systemic therapy has also been demonstrated to reduce the risk of cancer recurrence and improve survival in women with breast cancer. Adjuvant chemotherapy for younger women and tamoxifen for women over the age of 50 should be offered to all but the lowest risk women. The role of combined chemotherapy and tamoxifen is still evolving. For patients with locally advanced disease, early use of systemic and local regional therapies may be helpful. In the management of metastatic disease patients the major objective should be to reduce and possibly prevent morbidity with an emphasis on the quality of life for that individual.

8.6 Screening

Approximately one in nine women, living to the age of 90, will eventually be diagnosed as having breast cancer. Mammography for screening is directed towards women at high risk. Risk factors for breast cancer, apart from hereditary factors (i.e. two or more first degree relatives), include obesity, advancing age, early menarche, late menopause, nulliparity, delayed age of first birth and alcohol consumption.

Women are encouraged to perform regular breast self-examination (BSE). For pre-menopausal women this is best done in the week following the menstrual period and for post-menopausal women a specific day of the month should be chosen. The examination should include inspection of the breast and palpation of the breast and axilla. In order to perform adequate BSE the patient should be provided with the proper instruction as to the correct technique and the manner in which this should be performed.

8.7 Screening mammography

Individuals under the age of 40, with a strong family history of breast
cancer (i.e. two or more family members), should participate in screening programs if available. Genetic counselling and testing, if available, may also be of benefit in individuals at high risk for hereditary breast cancer. In these situations, annual screening mammography is recommended. Women between the age of 40 and 50 should also attend on an annual basis. In women aged 50 to 74, research has shown that 25% fewer breast cancer deaths can be expected in women if they have regular screening mammograms between the age of 50 and 69. To achieve this rate, at least 70% of eligible women in this age group must have regular screening mammography. Currently, we would recommend that women aged 50 to 74 have a screening mammogram at least every 24 months. Women aged 75 to 79 should have screening mammograms every two years as the incidence of breast cancer increases with age.

The use of ultrasound or thermography as screening methods are not recommended as these techniques do not have the sensitivity or specificity of mammography at this time.

8.8 Hereditary breast cancer
A woman with a sister or mother with bilateral breast cancer would be at a four-fold risk of breast cancer; if the case were post-menopausal a nine-fold risk; if the case was pre-menopausal she would be at an even higher risk if in addition to the family history she met any of the following criteria: family history of ovarian cancer, male breast cancer or Ashkenazi Jewish heritage.

8.9 Clinical practice guidelines

8.9.1 Management of a breast abnormality

8.9.1.1 Mass detected by mammogram only
In the case of a discrete mass lesion noted on examination may benefit from ultrasound examination to help distinguish between a cystic and a solid lesion. If the lesion is thought to be cystic it may be aspirated under ultrasound control and material sent for cytologic interpretation.

Solid masses found on ultrasound can be managed either by fine needle aspiration, stereotactic core needle biopsy or open surgical biopsy guided by fine wire localization.

8.9.1.2 Fine calcifications noted on mammogram
When this abnormality is identified and if the opinion of the diagnostic radiologist is such that the appearance is suspicious, stereotactic core needle biopsy or open biopsy is required. If the finding is thought to be less suspicious then a follow-up mammogram in 4-6 months may be recommended.

8.9.1.3 Palpable mass
All palpable masses require biopsy assessment. A mammogram may also be helpful in the assessment of a palpable mass to evaluate the rest of the breast tissue and also to assess the contralateral breast. A normal mammogram should not be a cause for delay in biopsy in these situations.

8.9.1.4 Cystic lesion
It is thought that if a lesion is cystic, it should be aspirated and if the cyst disappears following the aspiration and the fluid is free of blood then biopsy is not necessary. If, however, the cyst occurs repeat aspiration may be done. Cytologic examination of the fluid is usually not helpful.

8.9.1.5 Solid lesion
Any solid lesion needs to be assessed with either a fine needle aspiration, stereotactic core needle biopsy or open surgical biopsy.

8.9.2 Pre-operative investigations
Investigations that are recommended to determine the presence or absence of blood-borne metastatic disease prior to definitive surgical management include bilateral mammography, CBC, liver enzymes including alkaline phosphatase, chest x-ray. A bone scan is not ordinarily recommended for clinical stage T1, T2 and N0 cancers since asymptomatic patients are unlikely to have a positive bone scan due to metastatic disease. Tumours that are greater than 5 cm or where there are palpable axillary lymph nodes or an elevated alkaline phosphatase, a bone scan should be carried out in these situations. If the liver enzymes are elevated then ultrasound of the liver should also be performed.

8.9.3 Handling the surgical specimens
To assist treatment planning, sufficient detail must be obtained from
carcinoma in situ (DCIS). For patients treated with breast conservation, close cooperation and communication is required between the surgeon, radiologist/mammographer and pathologist to ensure that local therapy has been adequate.

The larger the focus of DCIS the higher the chance of a focus of microinvasive disease. Therefore an axillary lymph node dissection may be appropriate in patients with DCIS greater than 5 cm in diameter.

Total mastectomy remains an option for patients with DCIS, however recent evidence has demonstrated that radiotherapy reduces incidence of subsequent in situ and invasive breast recurrences. Currently, adjuvant radiotherapy is recommended with women with DCIS tumours greater than 1 cm in diameter or comedocarcinoma who are interested in breast conservation and in any patient who has a resection but in whom close margins (less than 5 mm) is present.

Women with well differentiated DCIS less than 1 cm in diameter, with complete radiographic and pathologic excision of the lesion, may be managed by wide local excision alone.

Those women with very diffuse areas of DCIS (greater than 5 cm or greater than or equal to 1/4 of the breast on mammography) have a substantial risk of recurrence even after excision and radiotherapy, and for these patients mastectomy is generally recommended. Tamoxifen or other adjuvant systemic therapy is not currently recommended, however when the results of controlled trials are available with this agent, tamoxifen may be subsequently recommended in an adjuvant setting.

8.9.4 Hormone receptor levels
The oestrogen receptor status of breast cancer can be determined by immunocytochemical staining of tissue specimens or aspirates. When possible it is preferable to submit the specimens, unfixed, immediately to pathology for selection of the most appropriate technique for handling each individual specimen. Immunocytochemical staining is the standard technique to test for oestrogen receptors. It can be performed on fresh or frozen tissue, scrapings obtained from the surface of small lesions, and from aspirates with reliable results. Oestrogen receptor staining may also be a useful technique in evaluating metastasis in patients with unknown primaries, in whom the possibility of metastatic breast carcinoma is in the differential diagnosis.

8.9.5 Tumour management based on TNM classification
8.9.5.1 Paget’s disease of the breast
Surgical excision remains the standard management for Paget’s disease of the nipple or breast. In selected patients partial mastectomy may be offered if this will lead to a satisfactory cosmetic result when the lesion is completely excised. A sample of underlying breast tissue will be removed with the nipple to assess if any associated invasive or in situ component is present.

8.9.5.2 Ductal carcinoma in situ
Detailed mammographic examination of the breast to obtain a preoperative assessment of the extent of the lesion is required for ductal carcinoma in situ (DCIS). For patients treated with breast conservation, close cooperation and communication is required between the surgeon, radiologist/mammographer and pathologist to ensure that local therapy has been adequate.

The larger the focus of DCIS the higher the chance of a focus of microinvasive disease. Therefore an axillary lymph node dissection may be appropriate in patients with DCIS greater than 5 cm in diameter.

Total mastectomy remains an option for patients with DCIS, however recent evidence has demonstrated that radiotherapy reduces incidence of subsequent in situ and invasive breast recurrences. Currently, adjuvant radiotherapy is recommended with women with DCIS tumours greater than 1 cm in diameter or comedocarcinoma who are interested in breast conservation and in any patient who has a resection but in whom close margins (less than 5 mm) is present.

Women with well differentiated DCIS less than 1 cm in diameter, with complete radiographic and pathologic excision of the lesion, may be managed by wide local excision alone.

Those women with very diffuse areas of DCIS (greater than 5 cm or greater than or equal to 1/4 of the breast on mammography) have a substantial risk of recurrence even after excision and radiotherapy, and for these patients mastectomy is generally recommended. Tamoxifen or other adjuvant systemic therapy is not currently recommended, however when the results of controlled trials are available with this agent, tamoxifen may be subsequently recommended in an adjuvant setting.
8.9.5.4 Stage I or II invasive cancer
Partial or total mastectomy combined with level I and level II axillary node dissection is recommended. Axillary nodal status is still the most powerful predictor for the need for adjuvant therapy.

8.9.5.5 Partial mastectomy, axillary dissection and radiation therapy
This combination is the standard treatment for patients with tumour less than 5 cm, providing that the tumour is unifocal and is sufficiently small in relationship to the size of the breast, in that wide local excision with a margin of normal tissue is possible and results in a reasonable cosmetic effect. In addition there should be no contraindication to radiotherapy. Removal of skin is a major factor affecting the cosmetic outcome in such cases. In general it is not necessary to remove skin unless it is involved by tumour. The cavity of the local excision should also be marked with small metallic hemoclips to help in localizing the area for subsequent radiotherapy should this prove to be necessary.

Even with pathologically negative margins, 25-40% of women treated with partial mastectomy alone will recur in the breast within 5-10 years. Therefore all patients treated by partial mastectomy and axillary dissection should be reviewed regarding the need for radiation therapy to the breast.

8.9.5.6 Modified radical mastectomy
This traditional method of surgical management is equivalent to breast conservation. When a modified radical mastectomy is performed the scar should be located with appropriate consideration of the site of the primary tumour but at the same time recognizing that some patients will desire breast reconstruction. The extent of the scar and any drains should be medial to the midaxillary line and above the sixth rib. Modified radical mastectomy is the treatment of choice if there are multiple tumours in a single breast and the size of the tumour is such that removal of the primary tumour with an adequate margin of normal tissue will lead to considerable distortion of the breast. Modified radical mastectomy is also the treatment of choice if there are absolute or relative contraindications to radiation therapy. Also elderly patients may find surgery easier than a partial mastectomy and node dissection and the subsequent three to five weeks of daily radiation therapy. In addition, modified radical mastectomy is also the treatment of choice in patients who are not available for follow-up and if the patient is not interested in breast conservation.

8.9.5.7 Radical mastectomy
The classic radical mastectomy procedure is rarely indicated today, although for tumours which are tethered to the underlying pectoral fascia over a small area, extension of the modified radical mastectomy to include removal of some portion of the underlying muscle is not infrequently performed. Classical radical operation may be indicated in an occasional patient where an adequate margin cannot be otherwise accomplished.

8.9.5.8 Locally advanced tumours
These tumours are generally regarded as inoperable. Occasional exceptions may occur in patients with large breasts and mobile T3 lesions without palpable nodes (T3, N0, M0). In these patients modified radical mastectomy may well be the treatment of choice. Surgery after chemotherapy and radiotherapy may be helpful in the prevention of local recurrence. These patients should be assessed for operability after response to therapy and a metastatic work-up repeated prior to considering modified radical mastectomy. This excludes patients with initial supraclavicular node involvement. Failure to respond to the initial chemotherapy is not an indication for surgery. Local control is not enhanced by operating on inoperable patients as cutting through tumour may actually worsen the prospects for local control by producing massive involvement through the tissue planes of the axilla.

These patients also have a high risk of widespread micrometastatic disease and chemotherapy may help eradicate micrometastasis and improve the local regional results of radiation therapy. A variety of intensive chemotherapy regimens are available but a standard commonly employed treatment would consist of six courses of cyclophosphamide, doxorubicin (adriamycin) and 5-fluorouracil (this is the CAF regimen).

Radiation therapy is usually given after the course of chemotherapy, however if the patients fail to show any improvement in their local
condition following the third course of chemotherapy then chemotherapy should be discontinued and local regional radiotherapy instituted. Radiotherapy is given to the whole of the breast and the lymph node areas. A boost dose to the site of the primary lesion is usually given. A dose to the axilla is also increased by approximately 10% if bulky axillary involvement was present.

Tamoxifen, 20 mg PO daily for five years, starting approximately four weeks after the completion of chemotherapy, should be offered to all patients with locally advanced tumours except patients under the age of 50 with oestrogen receptor negative tumours.

8.9.6 Special situations

8.9.6.1 Inflammatory breast cancer
Inflammatory breast cancer usually presents with rapid development of swelling redness in the classic peau d’orange (skin oedema) which may be mistaken for an infection and treated with antibiotics before the correct diagnosis is suspected. A mass is often palpated and the breast may be diffusely involved. Mammogram may show a discrete mass but often there is only diffuse increase in the density and skin thickening. Although the diagnosis is primary clinical, a distinctive pathological finding is the involvement of dermal lymphatic vessels by tumour cells which in turn produce the skin erythema and oedema.

Inflammatory breast cancer is the most aggressive form of malignant disease at this site with a median survival of 18-24 months irrespective of intensive combined modality treatment. Surgery is not generally advised as initial treatment apart from diagnostic biopsy. Patients who do not show clinical evidence of metastatic disease and who do not develop such evidence during treatment, mastectomy is considered after the completion of chemotherapy and radiation therapy. Chemotherapy and radiation are combined using cyclophosphamide, doxorubicin and 5-fluorouracil (CAF) every three weeks for six treatments. This is then followed by radiation to the breast, chest wall and regional lymph nodes. If the response to chemotherapy is poor after the initial six weeks, chemotherapy should be discontinued and radiation therapy given at that time. Tamoxifen, 20 mg daily for five years, is recommended for all patients except for those under the age of 50 who have oestrogen receptor negative tumours.

8.9.6.2 Local regional recurrence
Local regional recurrence may follow prior treatment with modified radical mastectomy or partial mastectomy, node dissection or radiation therapy. Local regional recurrence generally carries a poor prognosis but approximately 15% of patients will be long-term survivors after further local regional therapy. Metastatic disease and extensive local recurrence are incurable and the major aim of therapy is palliation. There is no evidence to suggest that early or aggressive treatment of recurrent or metastatic disease will add to survival. Occasionally preventing or eliminating impending or otherwise inevitable problems may be an indication for treatment in the asymptomatic patient. On occasion, with asymptomatic patients, local radiation may be given to the involved area if one is dealing with a solitary metastasis.

8.9.6.3 Carcinoma of the breast in pregnancy
Breast cancer is the most common tumour occurring in women during their reproductive years, so the combination of breast cancer and pregnancy is not uncommon but does produce special management problems. It is extremely important to assess the potential for cure in an individual patient and the welfare of both patient and foetus need to be evaluated.

Definitive treatment of breast cancer during the first trimester of pregnancy may endanger the foetus and therefore therapeutic abortion may be recommended. The treatment of the malignancy will then proceed as in the non-pregnant patient. Individuals who decline termination should be treated by modified radical mastectomy. No adjuvant radiation therapy or chemotherapy should be given during pregnancy.

During the second trimester the breast cancer can be adequately treated surgically without pregnancy termination. Modified radical mastectomy is the treatment of choice without immediate radiation and chemotherapy.

During the third trimester, foetal maturity should be assessed by standard techniques. Consideration should be given to inducing
labour as soon as foetal viability is present. Initial treatment by modified radical mastectomy is appropriate and as soon as feasible following delivery. The patient should receive additional treatment as for the non-pregnant state. (See Table 3).

Table 3: Histopathologic Type

<table>
<thead>
<tr>
<th>Type</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Carcinoma, NOS (not otherwise specified)</td>
<td>Ductal</td>
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<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>Lobular</td>
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Acknowledgements:
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