Guidelines for handling oesophageal biopsies and resection specimens and their reporting

Nassif B N Ibrahim

Keywords: oesophagus; reporting guidelines

The importance of the role of the histopathologist in the management of patients with oesophageal disease cannot be overemphasised. Pathological examination of specimens from these patients provides:

- Essential diagnostic and prognostic information for optimal clinical management.
- Material for research and audit.
- A database for epidemiological studies.

A close liaison between the surgeon, gastroenterologist, and histopathologist is of paramount importance, particularly in the evaluation of dysplasia or early carcinoma in Barrett’s oesophagus and, generally, to maximise diagnostic yield in any situation. The extent and usefulness of pathological information that can be conveyed to the clinicians is determined by the adequacy of clinical information, biopsy sampling, handling, laboratory processing, and any special studies that may be required. It is also influenced by the awareness of the histopathologist of the normal anatomy and histology of the oesophagus. There should be regular meetings between the surgeon, gastroenterologist, and histopathologist to discuss clinical and pathological findings.

Collection and preservation of specimens

The endoscopist should ensure that a separate container is used for biopsies taken from different sites so that the precise location of each biopsy can be identified. Containers should be properly labelled, including the number and site of the biopsy (for example, biopsy No 2 at 33 cm). Interpretation of the biopsy is considerably enhanced if it is taken with a large forceps and oriented1 (with its mucosal surface, if it is identifiable, upwards on small squares of porous non-soluble paper tissue) and placed immediately in an appropriate fixative (usually buffered 10% formalin or 10% formal saline). Biopsies should then be processed, embedded, and cut correctly oriented, having their luminal surface on one side of the section and the submucosal surface on the other. This is particularly important for the assessment of epithelial dysplasia or stromal invasion. However, small fibreoptic biopsies and those for malignant disease usually do not require orientation.

The histopathologist should ensure that the clinicians are well aware of the importance of completing the request forms with all the patient’s identification data, the endoscopic appearances, and the site of the biopsy. Details of previous biopsies and their laboratory reference numbers should be provided.

It is advisable that all oesophageal resection specimens are sent to the laboratory fresh in a sealed plastic bag placed in a leak proof container. Histopathology departments should ensure that there are appropriate arrangements in place for dealing with specimens arriving outside their normal working hours (this is especially important for taking tissues for research).

Precancerous glandular and squamous epithelial changes

**BARRETT’S OESOPHAGUS AND BARRETT’S DYSPLASIA**

There is confusion regarding the definition of Barrett’s oesophagus. This is partly because of lack of clarity regarding important anatomical landmarks and partly because of disagreement on precise diagnostic criteria.2 In this article, the widely but not universally accepted definition of Barrett’s as a columnar lined oesophagus is used. Within this definition, three epithelial types are recognised:1 (1) gastric cardiac type; (2) gastric fundic type; and (3) intestinal metaplasia. The latter is a columnar type of epithelium with goblet cells, identical to gastric intestinal metaplasia of the incomplete type. Because the squamocolumnar junction is usually irregular and does not always coincide with the anatomical gastro-oesophageal junction, it has been argued by some workers that the distal 2–3 cm of the oesophagus may be lined normally by columnar cardiac or fundic type mucosa, and they reserve the term Barrett’s oesophagus only to cases showing “specialised” intestinal metaplasia.4,5 The latter is the most important but by no means specific marker for Barrett’s oesophagus. It may show a
flat or villous configuration. This goblet cell containing metaplastic columnar epithelium is considered abnormal regardless of its length, and its presence indicates an increased risk of malignancy when it occurs anywhere in the oesophagus. The diagnostic histological criteria for Barrett’s oesophagus in adults is also applied in children, although it should be pointed out that intestinal metaplasia appears to be an age related change, and it is less commonly found in children.

The histopathologist can only definitively identify Barrett’s/columnar lined oesophagus in an oesophageal biopsy when columnar mucosal epithelium is seen overlying submucosal oesophageal glands or gland ducts opening onto the surface epithelium. Otherwise, the findings should be regarded as consistent with Barrett’s oesophagus if the biopsy was taken from a site more than 3 cm proximal to the anatomical gastro-oesophageal junction.

In considering the length of the Barrett’s segment, two types are recognised: short segment Barrett’s oesophagus (SSBO), a columnar lined mucosa with evidence of intestinal metaplasia extending for less than 3 cm into the distal oesophagus; and long segment Barrett’s oesophagus (LSBO), a columnar lined mucosa extending for more than 3 cm into the distal oesophagus with or without intestinal metaplasia. However, more recently the term ultra-short segment Barrett’s oesophagus has been applied to cases in which intestinal metaplasia is confined to the gastro-oesophageal junction (at the cardia). There appears to be a close relation between carditis, intestinal metaplasia of the cardia, and Helicobacter pylori infection.

Dysplasia associated with Barrett’s oesophagus is defined as unequivocal neoplastic transformation that is distinguishable from reactive and regenerative changes, and is classified histologically according to the guidelines for inflammatory bowel disease described by Riddell et al. Based on the severity of the architectural and cytological changes, dysplasia is graded as follows.

**Low grade dysplasia**—There is increased nucleus to cytoplasm ratio, nuclear hyperchromasia, and pleomorphism, but the gland/crypt architecture is preserved; the cytological changes are present throughout the crypt and generally the nuclei maintain a basal orientation; mitoses can be seen in the upper half of the crypts and there is loss of mucus secretion. There may also be some crowding and stratification of the nuclei which is confined to the basal half of the cells.

**High grade dysplasia**—There is severe disruption of the gland/crypt architecture with more severe cytological and nuclear atypia and loss of nuclear polarity, and often there is crowding and stratification of the nuclei that extends into the luminal parts of the cells.

**Indefinite**—This grading is applied when distinction between neoplastic transformation and regenerative changes cannot be made, or in the presence of active inflammation or in case of misorientation of the biopsy material when only the crypt bases are seen.

Generally, there is good interobserver agreement on the diagnosis of high grade dysplasia/intramucosal carcinoma, but there is significant disagreement on low grade dysplasia, which reflects the difficulty of differentiating dysplasia from reparative epithelial changes associated with active inflammation or ulceration.

The management of patients with Barrett’s oesophagus associated with high grade dysplasia is somewhat controversial. Some have recommended early surgical resection, while others advocate repeated endoscopic surveillance with biopsies. Those who advocate early surgery point out that when endoscopic biopsies show high grade dysplasia, approximately 50% of resected specimens show evidence of invasive carcinoma. It is recommended that high grade dysplasia should be confirmed by immediate re-endoscopy with multiple biopsies, or the diagnosis is agreed upon by two histopathologists experienced in reporting oesophageal biopsies before considering oesophagectomy. There are, as yet, no available markers that differentiate, with confidence, high grade dysplasia from carcinoma or predict progression of dysplasia to frank malignancy. The field of molecular biology is rapidly expanding and there are promising research developments in this area which may prove in future to be of practical significance. Currently, however, these cannot be recommended in routine practice. Most published series reported the presence of dysplasia in over 80% of cases of Barrett’s oesophagus complicated by carcinoma. A five tier classification for the various histological diagnoses associated with Barrett’s oesophagus is shown in table 1 and a suggested clinicopathological collaborative plan for the management of Barrett’s oesophagus is shown in table 2.

**SQUAMOUS DYSPLASIA AND IN SITU MALIGNANCY**

Neoplastic transformation in the oesophageal squamous epithelium, which should be distinguished from reactive and regenerative changes, can be classified as follows.

- **Mild dysplasia**—Cytological and nuclear atypia confined to the lower third of the epithelium.
- **Low grade dysplasia**—Infiltration of dysplastic cells extending into one or two thirds of the epithelium.
- **High grade dysplasia**—Infiltration of dysplastic cells extending into the upper one third of the epithelium.
- **Intraepithelial carcinoma**—Infiltration of dysplastic cells extending into the basal third of the epithelium.
- **Invasive carcinoma**—Infiltration of dysplastic cells extending beyond the basement membrane.

**Table 1** Histological findings in endoscopic biopsies from Barrett’s oesophagus

<table>
<thead>
<tr>
<th>Grade</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Negative for dysplasia</td>
<td>There is no evidence of atypia in the biopsy.</td>
</tr>
<tr>
<td>Indefinite/low grade dysplasia</td>
<td>There is increased nucleus to cytoplasm ratio, nuclear hyperchromasia, and pleomorphism, but the gland/crypt architecture is preserved; the cytological changes are present throughout the crypt and generally the nuclei maintain a basal orientation; mitoses can be seen in the upper half of the crypts and there is loss of mucus secretion. There may also be some crowding and stratification of the nuclei which is confined to the basal half of the cells.</td>
</tr>
<tr>
<td>High grade dysplasia</td>
<td>There is severe disruption of the gland/crypt architecture with more severe cytological and nuclear atypia and loss of nuclear polarity, and often there is crowding and stratification of the nuclei that extends into the luminal parts of the cells.</td>
</tr>
<tr>
<td>Intraepithelial carcinoma</td>
<td>Infiltration of dysplastic cells extending into the upper one third of the epithelium.</td>
</tr>
<tr>
<td>Invasive carcinoma</td>
<td>Infiltration of dysplastic cells extending beyond the basement membrane.</td>
</tr>
</tbody>
</table>

**Table 2** Suggested clinicopathological collaborative plan for the management of Barrett’s oesophagus

<table>
<thead>
<tr>
<th>Category</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Negative for dysplasia</td>
<td>Yearly endoscopic surveillance.</td>
</tr>
<tr>
<td>2. Indefinite/low grade dysplasia</td>
<td>Repeat biopsy in 3–6 months until two consecutive negative biopsies or until dysplasia progresses.</td>
</tr>
<tr>
<td>3. High grade dysplasia</td>
<td>Confirm by immediate rebiopsy or the diagnosis is confirmed by another experienced pathologist; oesophagectomy may be considered.</td>
</tr>
</tbody>
</table>

*Intraepithelial adenocarcinoma denotes unequivocal breach of the gland basement membrane but not the muscularis mucosae.*
Handling oesophageal biopsies and resection specimens

Moderate dysplasia—Cytological and nuclear atypia confined to the lower two thirds of the epithelium.

Severe dysplasia—Cytological and nuclear atypia involves almost the full thickness of the epithelium.

Carcinoma in situ—The full thickness of the epithelium is abnormal, being composed of neoplastic cells.

Gross examination and laboratory investigation

**BIOPSY SPECIMENS**
- The number of biopsies in each container should be documented. Biopsies must be handled with care to avoid crush artefacts.
- Biopsies should be step sectioned at three different levels and stained routinely with haematoxylin and eosin. The unstained intervening sections should be kept for any special stains that might be required later.

**SURGICAL RESECTION SPECIMENS**
Fresh specimens should be carefully examined, and the outer (circumferential) resection margin painted with Indian ink or other marking dye. This is important for the assessment of completeness of excision and measurement of distance of tumour from the circumferential resection margin. The specimen should then be opened longitudinally, pinned to a cork board, and fixed by immersion in a fixative (usually buffered 10% formalin or 10% formol saline) for 48–72 hours to ensure adequate fixation and facilitate obtaining thin slices. It should be noted that after resection the oesophagus undergoes shrinkage, which affects the upper more than the lower margin, with tumour tissue changing little in length. Even if the oesophagus is immediately pinned and fixed after resection it shrinks by more than 10%, and if pinning and fixation are delayed it shrinks by more than 50%, which accounts for the discrepancy between surgeons’ and histopathologists’ measurements.

After fixation, it is advisable to have a photograph or diagrammatic representation of the specimen made to illustrate pathological findings and indicate sites of blocks selected for histological examination.

The gross description should include:

1. The length and maximum diameter of the oesophagus, and for oesophagectomy specimens, the length of the gastric segment along both the lesser and greater curve.
2. The presence of any abnormalities such as achalasia, cysts, webs, rings, diverticula, strictures, or ulcers, and these should be described, measured, and sampled adequately.
3. In tumour containing specimens: the presence or absence of Barrett’s mucosa or any other area of background mucosal abnormality should be accurately described, measured, and adequately sampled.
4. The gross appearance of the tumour (polypoid, fungating, flat, ulcerated), as well as its length and width.
5. Site of tumour and its relation to the gastro-oesophageal junction.
6. Distance of the tumour from the proximal and, in oesophagectomy specimens, distal surgical lines of excision.
7. Extent of invasion of the oesophageal wall.
8. Relation of the tumour to the overlying serosa/outer fibrofatty tissue.
9. Lymph nodes which are clearly replaced by tumour should be sampled. All other lymph nodes should be completely embedded.
10. Tumour should be adequately sampled; this is important for the assessment of various prognostic features and for the evaluation of tumour heterogeneity. It is recommended that the whole tumour is serially sectioned with a sharp knife. First, the bulk of the tumour should be sectioned transversely, to allow assessment of the circumferential resection margin, then the proximal and distal parts sectioned longitudinally, to allow demonstration of the junction between tumour and adjacent non-neoplastic mucosa (fig 1, A, B and C). Sections should be laid flat and examined to assess maximum tumour infiltration. Photographs may be taken at this stage as a permanent record. As a minimum, four blocks should be taken from the tumour, two to include maximum circumferential infiltration and two to include tumour and adjacent non-neoplastic mucosa; one proximally and the other distally. One transverse section should be taken from the oesophagus above the tumour and, depending on tumour site, one distal to the tumour (mainly to demonstrate submucosal vascular invasion). Sections should also be taken to include the proximal and distal resection margins, and if applicable, the gastro-oesophageal junction/presumed junction. Abnormal background mucosa should be adequately sampled.

If no obvious tumour is seen on macroscopic examination of oesophagectomy specimens carried out following a diagnosis of in situ squamous carcinoma or high grade dysplasia/carcinoma in Barrett’s mucosa, the entire segment of oesophagus lined by abnormal mucosa should be serially sectioned in longitudinal bands (fig 1D). Before sectioning the specimen, the demonstration of squamous dysplasia/in situ squamous carcinoma may be facilitated by the application of Lugol’s solution to the oesophageal mucosa, at a concentration of 1% for one to two minutes and observing the colour change (normal squamous epithelium stains dark brown while severe dysplasia/in situ carcinoma shows no colour change). Representative blocks of tissue (individually numbered and their sites marked on the specimen photograph or diagram), 25 mm in length and 3–4 mm width (painted along their proximal margin with Indian ink or other marking dye), should be taken for histology. If these fail to show malignancy, further blocks should be taken. As multifocal disease is often
seen in early/superficial squamous cell carcinomas, these specimens should be thoroughly sampled.

**HISTOLOGY**

**Non-neoplastic diseases**

Haematoxylin and eosin (H&E) sections are often sufficient to establish the diagnosis. However, in biopsies of columnar type mucosa sections should be stained with periodic-acid-Schiff (PAS) and alcian blue at pH 2.5 as an aid for the recognition of intestinal metaplasia in Barrett’s oesophagus. The columnar cells in Barrett’s epithelium differ from normal gastric surface cells in that they contain alcian blue positive mucus but stain less intensely than goblet cells. In rare cases where there is a suggestion of infective oesophagitis, particularly in immunocompromised patients, a search for viral inclusions and stains for acid fast bacilli and fungi should be carried out, and these may be supplemented by immunohistochemical techniques.

**Neoplastic diseases**

In oesophageal malignancy, the histopathology report should incorporate all data that are regarded as having prognostic significance. Residual disease at surgery, depth of invasion, and lymph node status are the most important, independent prognostic indicators, while other histopathological variables such as tumour his-
Handling oesophageal biopsies and resection specimens

tological type, tumour grade, vascular invasion, and even the status of microscopic resection margin, appear to lose their prognostic significance in multivariate analysis. However, it is recommended that detailed histopathological data (see below) should be recorded, as they may prove useful in predicting response to future therapeutic regimens and for epidemiological studies.

H&E sections are often sufficient to establish the diagnosis. Nevertheless, histochemical techniques (such as mucin stains, Grimelius technique, and so on) and immunohistochemical methods such as neuroendocrine markers (for example, NSE, PGP9.5, and chromogranin) can be useful in establishing the precise tumour type. Techniques such as DNA studies cannot yet be recommended for routine use.

The histopathology report should incorporate:

1. **Tumour type**—Tumours should be classified according to a widely used nomenclature such as the WHO classification.

2. **Tumour differentiation**—This should be recorded according to the predominant area as well differentiated, moderately differentiated, or poorly differentiated.

3. **Local invasion and the depth of infiltration of the wall**—The text of the histopathology report should clearly state whether the tumour is a high grade dysplasia/in situ carcinoma or an invasive malignancy. The depth of invasion (mucosa/muscularis mucosae, submucosa, muscularis propria, serosa/outer fibrofatty tissue, or adjacent structures) should be documented.

4. **Background mucosal abnormality**—It has been suggested that the presence of concomitant Barrett’s epithelium in distal oesophageal adenocarcinoma improves prognosis.

5. **Vascular/lymphatic invasion**—Studies have shown that vascular (venous/lymphatic) invasion is an important prognostic factor in univariate analysis.

6. **Lymph node metastasis and number of lymph nodes involved by tumour**—These are among the most important, and independent prognostic indicators. The site of lymph nodes involved by tumour (paraoesophageal, gastric lesser curve, gastric greater curve, and others if received separately and identified as to site) may be useful to document, though there is evidence to suggest that this may not be of prognostic significance.

7. **Other histological variables**—Features that may be recorded but appear to be of little independent prognostic significance include: pattern of advancing margin (pushing or infiltrating), lymphocytic reaction, and intramural metastasis. Infiltration of perinodal fibrofatty tissue by tumour was suggested to be an important prognostic factor in patients with curative resection for oesophageal carcinoma, but this needs to be supported by further studies.

8. **Adequacy of excision**—Carcinoma involving the circumferential, proximal, or distal resection margins and clearance (in mm) should be documented. The status of resection margins has been shown in univariate analysis to be an important prognostic factor. Dysplasia or Barrett’s metaplasia at the proximal or distal resection margin should also be documented.

**Adenocarcinoma at the gastro-oesophageal junction**

The distinction between adenocarcinoma associated with Barrett’s oesophagus and adenocarcinoma of the “gastric cardia” secondarily involving the oesophagus is difficult and may be impossible. Current evidence suggests that many carcinomas involving the gastro-oesophageal junction actually arise in Barrett’s epithelium. Short segment and ultrashort segment Barrett’s mucosa may be completely overrun by the adenocarcinoma at presentation. Adenocarcinoma at the gastro-oesophageal junction can be classified as follows.

1. **Barrett’s adenocarcinoma**—If adjacent Barrett’s epithelium with dysplasia is seen or if non-dysplastic Barrett’s epithelium is present adjacent to an adenocarcinoma where more than 50% of the tumour mass is present in the oesophagus.

2. **Gastric adenocarcinoma**—If dysplasia is seen in the gastric epithelium immediately adjacent to the tumour distally or more than 50% of the bulk of the tumour is present in the stomach and is not associated with dysplasia in Barrett’s epithelium.

3. **Oesophageal adenocarcinoma not otherwise specified**—Where origin from Barrett’s epithelium cannot be demonstrated and the tumour lies predominantly in the oesophagus and there is no associated gastric dysplasia. It can be assumed that tumour arose from Barrett’s mucosa and has been overgrown by it. In oesophagogastrectomy specimens containing tumours straddling the gastro-oesophageal junction where mucosal landmarks are no longer identifiable, the anatomical gastro-oesophageal junction should be used as the landmark. This is best recognised at the site of the “notch” or the peritoneal reflection at the junction of the oesophagus and greater curve.

**STAGING OF OESOPHAGEAL CANCER**

The TNM (tumour, node, metastasis) staging system is the most important prognostic indicator (table 3). However, this does not differentiate between tumours confined to the mucosa and those involving the submucosa, a distinction that has been shown to be of considerable prognostic significance. Several studies have demonstrated that oesophageal tumours which are confined to the epithelial lining (in situ carcinomas) are always curable, and invasive tumours that are confined to the mucosa are nearly always curable. However, submucosal oesophageal cancer (unlike its gastric counterpart) has been shown to be a relatively advanced disease which is associated with a significant risk of vascular invasion and
Table 3: TNM clinical classification

<table>
<thead>
<tr>
<th>T: Primary tumour</th>
<th>N: Regional lymph nodes</th>
<th>M: Distant metastasis</th>
</tr>
</thead>
<tbody>
<tr>
<td>T1 Tumour invades lamina propria or submucosa</td>
<td>NX Regional lymph node cannot be assessed</td>
<td>MX Distant metastasis cannot be assessed</td>
</tr>
<tr>
<td>T2 Tumour invades muscularis propria</td>
<td>N0 No regional lymph node metastasis</td>
<td>M0 No distant metastasis</td>
</tr>
<tr>
<td>T3 Tumour invades adventitia</td>
<td>N1 Regional lymph node metastasis</td>
<td>M1 Distant metastasis</td>
</tr>
<tr>
<td>T4 Tumour invades adjacent structures</td>
<td>N2 Regional lymph node metastasis</td>
<td>M1a Metastasis in cervical lymph nodes</td>
</tr>
<tr>
<td></td>
<td>N3 Regional lymph node metastasis</td>
<td>M1b Other distant metastasis</td>
</tr>
</tbody>
</table>

For tumours of the lower thoracic oesophagus

- M1a Metastasis in coeliac lymph nodes
- M1b Other distant metastasis

For tumours of the upper thoracic oesophagus

- M1a Metastasis in cervical lymph nodes
- M1b Other distant metastasis

For tumours of the mid-thoracic oesophagus

- M1a Not applicable
- M1b Non-regional lymph node or other distant metastasis

pTNM pathological classification

The pT, pN, and pM categories correspond to the T, N, and M categories.

Lymph node metastasis, particularly in the case of squamous cell carcinoma. Lymph node metastasis has been reported to occur in 30–50% of squamous tumours involving the submucosa, a figure which is similar to advanced carcinoma. It is recommended that if the term “superficial carcinoma” is used in histopathology reporting, it should be qualified by the depth of invasion and, if applicable, the lymph node status.

As TNM is the most widely used prognostic indicator, it is recommended that the histopathology report should include a conclusion that incorporates the pTNM staging system (TNM staying with histopathological evaluation). Additionally, the use of a proforma is recommended for uniformity in reporting, which is important for accurate comparison of results of treatments at different centres, is also recommended.

I wish to thank Dr David Day, Dr Howard Rigby, Professor Neil Shepherd, and Dr Bryan Warren for reading the manuscript and for helpful comments, and Dr Masaki Mori for advice on Lugol’s staining.

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Letters

Necropsy organ weights are largely useless

In 1993 the Royal College of Pathologists published Guidelines for Post Mortem Reports.1 The guidelines may well have led to improvements in necropsy practice but were unrefere-
cenced and their evidence base was unstated. They have been quoted as a gold standard for audit of necropsy reports,2 and reiterated in an editorial in this journal.1 Most of the guidelines are sensible but I question the recommendation, in adult necropsies, of routine weighing of organs. Excluding the heart, the weighing of which can provide important information (particularly when the ventricles are weighed separately), organ weights are of little or no value. The apparent weight of an organ depends on dissection technique and on the accuracy of the weighing balance. In common with other branches of pathology, a numerical result should always be accompanied by the normal range, corrected for the patient’s sex, age, and body size. I suspect very few of us comply with this basic rule. However, even if we were to provide reproducibly accurate and referenced weights, they would not be of use to our clinical colleagues, who are accus-
tomed to clinical evaluation of organs, supplemented by imaging techniques in which, at most, a single linear measurement is the reported desirability of histological examination in all necropsy cases. There is an increasingly vocal body of opinion within the profession which disagrees with this requirement and the same audit of necropsy reporting in East Anglia provided useful data in relation to the sensible use of histology in necropsy practice.3 Such evidence must be incorporated into any future national guidelines for necropsy practice in order that the guidelines reflect the views of most practising pathologists, rather than those of a few interested individuals who perform few, if any, necropsies.

Our principal aim should not be for more necropsies but for better quality necropsies, which are fully supported by a system of formal quality assurance. The Royal College of Pathologists has a major part to play in this process and in recent years has overlooked the necropsy in favour of diagnostic histopathology and cytopathology. The intermittent production of necropsy related guidelines has been useful but a complete reappraisal of all necropsy related matters is urgently required at a national level. Other countries have already addressed these issues4 and this is presumably one reason for the commissioning of a Royal College of Pathologists Working Party to examine the current status of necropsy in the United Kingdom.


We obtained the following comments on this letter:

From Dr R D Start

Dr Barker is correct in his statement that the Royal College of Pathologists’ publication Guidelines for Post Mortem Reports has led to improvements in necropsy reporting.1 This is clearly evident within recent reports of the National Confidential Enquiry into Peri-
operative Deaths (NCEPOD).2 The necropsy report guidelines currently provide the only national audit standard for the quality of necropsy reports and I would agree that an appropriate evidence base is desirable. This was one reason behind my suggestion for national practice guidelines for necropsy.1 I am surprised that the routine weighing of organs is in question. Normal organ weight ranges, corrected for patient sex, age, and body size, are available and can be used if necessary. Some variation may occur with dissection technique but this would be consistent for individual pathologists and could be addressed in any national practice guidelines. Frequently, whole bodies and organs are fundamental requirements for any mortuary and accuracy is simple to achieve. The suggestion that we stop this “ritualistic, pseudoscientific practice”3 in order to com-
municate relevant findings to our colleagues by way of other variables such as a single linear organ measurement is illogical and unac-
ceptable. Linear measurements of focal abnormalities complement gross necropsy findings and allow comparative audit of modern imaging techniques, many of which now give measurements in three dimensions. Sup-
plementary information can also be provided by the dissection of organs in the planes of examination typically seen in modern imag-
ing techniques.

Most clinicians, coroners, and (possibly more importantly) relatives are able to comprehend the concept that organs are abnormal when organ weights are put in the context of normal ranges, particularly if the organ is several times the average normal weight for an individual of similar size, sex, and weight. Although I am unable to provide a specific evidence base to support the use of organ weights, the reasoning behind not put-
viding them is difficult to comprehend, particularly when the most recent major regional audit of necropsy reporting in East Anglia (including Norwich!) found “all pathologists agreed the value of routine weighing of heart, lungs and brain.”5 I hesitate to suggest that organ weights may be an indirect measure of necropsy quality because through personal experience I have found not only that high quality necropsy reports can be generated from appalling necropsies but also that seemingly accurate organ weights can be determined without removal from cadavers.

More questionable than organ weight measurement is the reported desirability of histological examination in all necropsy cases. There is an increasing vocal body of opinion within the profession who disagree with this requirement.4 The same audit of necropsy reporting in East Anglia provided useful data in relation to the sensible use of histology in necropsy practice.3 Such evidence must be incorporated into any future national guidelines for necropsy practice in order that the guidelines reflect the views of most practising pathologists, rather than those of a few interested individuals who perform few, if any, necropsies.

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From Dr M J Goddard

In our paper,6 we included the measurements of major organ weights (heart, lungs, brain, liver, and kidneys) as one of the audit criteria. We do not consider this a gold standard but set this audit standard on the basis of the guidelines of the Royal College of Pathologi-
gists, together with a consensus view of the pathologists whose reports were used in the audit.

It is interesting to note that in the audit, this was the area of the internal examination which was done least well, with only 73% of reports recording all five weights in the initial audit, improving to 84% at the time of re-audit. This could be taken to suggest that at least a proportion of pathologists are unwilling to record data that they perceive as meaningless.

As one of the pathologists who reviewed the necropsy reports at the time of the audit, I would have to say that there were very few if any instances where the weight of the organ other than the heart contributed to the quality of the report. I would have to agree that from my own personal viewpoint, together with my impressions gleaned from the audit, the routine weighing of organs was use-
ful, provides no information to clini-
cal colleagues, and should cease.


Book reviews


The editors have done a marvellous job, more than fulfilling their stated aim of producing a volume describing the multidisci-
plinary state of modern pathology which will be of interest to a wide range of readers. The book is beautifully produced with excel-

tent colour photographs and line diagrams which clearly explain the practicalities de-
scribed in the text. I was particularly
impressed by the many tables and flow charts, which can be used as aids to decision making.

All aspects of pathology are covered and it is very easy to find the specific information one needs; the first page of each chapter has its own index, there are summaries of topics at the top of each page, and the general index is detailed and comprehensive. Each subject is clearly introduced with relevant background information. Practical details are easy to follow and alternative methods discussed. Attention is paid to the interpretation of results and to the use of quality controls. At the end of each chapter future directions are covered and there is a useful further reading list. The unifying theme to the book is the application of similar methods to different disciplines. Microscopy and molecular techniques are referred to frequently and it is obvious that a common language is developing between pathologists. This welcome situation will be helped considerably by this work, which should be on the bookshelf of every laboratory. The availability of a version on CD-ROM will increase its appeal.

D M BARNES


Immunologists may well wonder whether there is any need for another volume to complement the many excellent texts that already exist on clinical immunology. Gavin Spickett explains why he wrote the book, saying it was for selfish reasons in that the book that he would like to have had was not available when he was a trainee. Therefore the volume has to be assessed from two points of view: first, does it fill a need in the market, and second, does it meet Dr Spickett’s own requirements?

In my opinion the answer to both questions is yes. The volume is up to date, comprehensive, and easily referenced. The text includes information on both clinical and laboratory immunology and has useful appendices on quality and managerial issues which laboratory trainees will find particularly valuable. Skipping through the volume, I find it is generally very up to date, for example the section on X-linked lymphoproliferative disease identifies the cloned gene, information about which was only published in *Nature* in October 1998.

In a volume such as this, one’s attention tends to get drawn to areas of one’s own particular interest and not surprisingly not all contributions are similarly up to date. I found it surprising that the section on HIV infection did not mention combination antiretroviral therapy, particularly protease inhibitor therapy, or viral load testing in any detail, and also indicated that there was no value in sequential monitoring of CD4 counts once they had fallen below 0.05—information that is clearly incorrect following the advent of new treatments.

However, such criticisms are minor in what is otherwise an excellent volume, and I applaud Dr Spickett’s omission of fundamental immunology which has little or no place in such a book. I think that immunology trainees will find this volume invaluable, as will most of their trainers, but whether the market will be larger than the 150 or so individuals that this group comprises remains to be seen. I certainly cannot see many SHOs in general medicine carrying this book along with the many other small slim volumes that currently sit in RMO’s pockets, but I may be proved wrong on this too.

GRAHAM BIRD

Correction

Because of an error in the publishing process, Dr Ibrahim’s Best Practice article in the February issue (Guidelines for handling oesophageal biopsies and resection specimens and their reporting, vol 53, no 2, pp 89–94) has been numbered 155 instead of 156. Reprints of this article will be numbered correctly, as will references to the article in our in-house Best Practice advert that appears at the end of each issue. We apologise for this error.

Notice

St Mary’s Hospital Campus of ICSM, London

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