The problem of 'chronic mastitis' with epitheliosis

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SYNOPSIS

The incidence of breast cancer complicating chronic mastitis with epitheliosis has been reported to be as high as 45% and as low as 6%. At least part of this discrepancy appears to be due to difficulty in distinguishing between benign and malignant intraduct epithelial proliferation. A small series is presented here in which the incidence of invasive cancer after local excision of breast tissue showing chronic mastitis with epitheliosis was nil.

It is concluded that it is necessary to follow up larger series of cases longer before it can be said that severe epitheliosis and borderline lesions should in all cases be treated by anything more radical than local excision.

Lesions of breast tissue involving intraduct epithelial proliferation provide histopathologists with some of their most difficult problems. The borderline between proliferation of epithelium and intraduct carcinoma is ill defined and individual pathologists may vary in their assessment of any given case. Nevertheless they will be called upon to advise surgeons on the probable natural history of lesions falling into this group, and as the assessment of the risk of cancer is involved the clinicopathological responsibility is considerable.

A large literature has accumulated relating to the association between the condition variously known as chronic cystic mastitis (Foote and Stewart, 1945), fibroadenomatisosis (Kiaer, 1954), cystic disease, and cystic hyperplasia (Bonser, Dossett, and Jull, 1961) with breast cancer. These authors and many others agree that the condition which is referred to in this paper as 'chronic mastitis' is associated with an increased liability to breast cancer, particularly when there is intraduct epithelial proliferation.

Davis, Simons, and Davis (1964), after reviewing the world literature, estimated the incidence of breast cancer in all women with chronic mastitis as 2-64 times that of women in general. Authorities differ considerably as to the incidence of cancer when intraduct epithelial proliferation is a feature of chronic mastitis. Davis et al. (1964) found that only one out of 16 of their cases (6.25%), showing what they describe as 'solid epithelial hyperplasia' in an excision biopsy, subsequently developed breast cancer. But in the series of Kiaer (1954), who divided his 321 cases of fibroadenomatisosis into three grades according to the extent of the epithelial proliferation, no less than nine out of 20 women (45%) with grade 3 fibroadenomatisosis developed breast cancer.

Such marked variation in the recorded incidence of carcinoma complicating chronic mastitis with intraduct epithelial proliferation underlines the pathologist's difficulty in advising the surgeon and justifies further study.

TERMINOLOGY

From descriptions of the pathology of chronic mastitis (Foote and Stewart, 1945; Kiaer, 1954; Bonser et al., 1961) it is apparent that variations in terminology can lead to confusion. 'Epitheliosis' is used here to describe intraduct epithelial proliferation (Figs. 1, 2, 3 and 4) and it must be clearly distinguished from 'adenosis', which is a proliferation and branching of the lobular tree (Fig. 5) and is not regarded as being precancerous (Dawson, 1933). Epitheliosis may be accompanied by a variable amount of fibrovascular proliferation (Fig. 6), and where the latter predominates the appearances merge into papillomatosis of the duct characterized by broad fibrovascular papillae covered by a single or double layer of epithelium (Fig. 7).

It is important to distinguish epitheliosis from intraduct carcinoma. Carcinoma of the breast arises from the epithelium of the ducts or less frequently from the lobules. So long as the tumour is confined to the lumina of the ducts or their terminal parts within the lobules the terms 'intraduct carcinoma' (Muir, 1941) and 'lobular carcinoma in situ' (Foot, 1941)
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and Stewart, 1941) are used. Gowing (1968) recognizes three patterns of intraduct carcinoma—solid, cribriform, and papillary. A well known form of the solid intraduct carcinoma is the comedocarcinoma which is characterized by marked cellular atypia and necrosis (Fig. 9).

MATERIAL

Between 1946 and 1963, out of a series of over 1,000 cases of chronic mastitis referred to Westminster Hospital, 60 excision biopsies were reported as showing epitheliosis. During the same period 20 cases were diagnosed as intraduct carcinoma. Follow-up information, largely from the patients' own doctors, was obtained in 29 of the first group, and from the second group such information was available in 15, who were followed up as outpatients. The sections from these 44 cases were reviewed.

GRADING EPITHELIOSIS

A rough quantitative method of grading epitheliosis according to the number of ducts affected was adopted. The limitations of the technique are obvious, and it should be emphasized that cases showing cellular atypia were not classified as epitheliosis. The intraduct epithelial changes were finally graded as follows:

SLIGHT EPITHELIOSIS This is defined as epithelial proliferation partly or completely filling less than three or four ducts in any low-power field (Fig. 1).

SEVERE EPITHELIOSIS Epithelial proliferation which completely fills more than three or four ducts in any low-power field (Figs. 2, 3 and 4).

BORDERLINE MALIGNANCY Epithelial proliferation with some cellular atypia or other features where a definite diagnosis of intraduct carcinoma could not be made (Figs. 8a and b).

INRADUCT CARCINOMA A definite diagnosis of intraduct carcinoma was only made if there was

FIG. 1

FIG. 1. Slight epitheliosis. Three ducts are partly filled by proliferated epithelium. Haematoxylin and eosin × 36.

FIG. 2

FIG. 2. Severe epitheliosis. Numerous ducts are completely filled by epithelium. Note the presence of foamy cells amongst the epithelium in some of the ducts (arrows). Haematoxylin and eosin × 36.
FIG. 3. High-power view of one of the ducts in Fig. 2 showing variation in shape and intensity of staining of the nuclei of the cells filling the duct. Haematoxylin and eosin x 90.

FIG. 4. Duct from another field in the same section as shown in Figures 2 and 3. There is variation in shape and intensity of staining of the nuclei as in Fig. 3, but there is also an acinar pattern and some fibro-vascular proliferation. Haematoxylin and eosin × 90.

FIG. 5. Adenosis showing proliferation and branching of the lobular tree. In the area marked with the arrow, the appearances are becoming those of sclerosing adenosis. Haematoxylin and eosin × 36.
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marked cellular atypia often accompanied by appreciable mitotic activity and necrosis (Fig. 9).

RESULTS

My own review of the sections resulted in two of the cases originally diagnosed as epitheliosis being transferred to the malignant or borderline group, and five of those of the presumed intraduct carcinoma being reclassified as epitheliosis (Tables I and II). This fact underlines the variation in personal interpretation of these intraduct lesions which obviously has an important influence on treatment (Table I, cases 1 to 5).

Of the 30 patients with epitheliosis treated by local excision (including three originally diagnosed as intraduct carcinoma) none have returned with breast cancer, and all were alive and well when last heard of (Table I). This includes 10 patients with severe epitheliosis, eight of whom were followed up for over five years.

Table II shows that the patients with intraduct carcinoma and those on the borderline have done well, no deaths from breast cancer occurring in the group. All but three (Table II, cases 1, 2, and 6) had a simple or radical mastectomy. In case 7 mastectomy performed six years after intraduct carcinoma was found in an excision biopsy revealed invasive carcinoma.

DISCUSSION

Any study of a small number of cases must be of limited value, but the findings in the present series suggest that the short-term risk of cancer complicating all cases of chronic mastitis with epitheliosis is small. But how can the remarkable variation in the recorded incidence of carcinoma complicating...
Classed as a borderline lesion due to resemblance to the variety of intraduct carcinoma characterized by pale staining cells arranged with their long axes at right angles to the walls of the ducts. Haematoxylin and eosin × 90.

The presence of central necrosis resulted in classing as a borderline lesion. Haematoxylin and eosin × 90.

Comedocarcinoma showing necrosis, marked cellular pleomorphism, and mitotic activity. Haematoxylin and eosin × 90.
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FIG. 10. In this cribriform intraduct carcinoma the cytological features of malignancy are less obvious than in Figure 9. The malignant nature of the process is, however, shown by the invasive carcinoma in the surrounding breast tissue. Haematoxylin and eosin × 90.

FIG. 11. Lobular carcinoma in situ. Ductules filled with uniform cells showing occasional mitotic figures. Haematoxylin and eosin × 90.

FIG. 12. Degeneration of epitheliosis with formation of 'colostrum-like' foamy cells (see also Figure 2). Haematoxylin and eosin × 120.
chronic mastitis with epitheliosis be explained?

The paper of Davis et al (1964) contains few illustrations, and it is thus difficult to compare their results with those of Kiaer (1954) and my own results. Kiaer's monograph, on the other hand, is extensively illustrated, and if one assumes that what I have called 'severe epitheliosis' is similar to the grade 3 fibroadenomatosis of Kiaer, in whose series nine out of 20 cases developed breast cancer, there is an obvious discrepancy, even after allowing for the smaller number of cases in my series.

There are two possible explanations. The length of follow up is clearly important. Of Kiaer's 321 cases of fibroadenomatosis of all grades the average follow up was 17 years compared with seven years in the present series. Two of Kiaer's cases of grade 3 fibroadenomatosis did not develop breast cancer until 18 and 24 years after local excision. I agree with Kiaer that very long periods of time must elapse in assessing the malignant potential of epitheliosis.

The average follow up period of the 16 cases of Davis et al (1964) showing 'solid epithelial hyperplasia', of which only one developed breast cancer, was 14 years, which is comparable to that in Kiaer's series. This suggests that another possible factor in the high incidence of malignancy in Kiaer's series is

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**TABLE I**

<table>
<thead>
<tr>
<th>Grade of Epitheliosis</th>
<th>No. of Cases</th>
<th>Initial Treatment</th>
<th>Follow up and Further Treatment</th>
</tr>
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<tbody>
<tr>
<td>Slight</td>
<td>20</td>
<td>Local excision</td>
<td>At 5 yr, four alive and well</td>
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<tr>
<td></td>
<td></td>
<td></td>
<td>At 5–10 yr, 13 alive and well</td>
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<tr>
<td></td>
<td></td>
<td></td>
<td>Further excision same breast, chronic mastitis only two cases. Excision opposite breast, chronic mastitis only two cases. Further excision same breast, slight epitheliosis one case. Simple mastectomy opposite breast, chronic mastitis only one case. 10–15 yr, three alive and well. Excision opposite breast, chronic mastitis only one case.</td>
</tr>
<tr>
<td>Severe</td>
<td>Original diagnosis chronic mastitis with epitheliosis, 7</td>
<td>Case 1, local excision, Case 2, local excision</td>
<td>At 2 yr, alive and well, pregnant</td>
</tr>
<tr>
<td></td>
<td>Original diagnosis intraduct carcinoma, 5</td>
<td>Case 3, local excision and bilateral oophorectomy</td>
<td>At 3 yr, simple mastectomy showed severe epitheliosis.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Case 4, local excision, radiotherapy, and temporary menopause 1955</td>
<td>At 8 yr, alive and well</td>
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<td></td>
<td></td>
<td>Case 5, radical mastectomy</td>
<td>At 12 yr, alive and well, 1960 pregnancy terminated and sterilized</td>
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<td>At 7 yr, alive and well</td>
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1Average follow up of all cases was seven years.

**TABLE II**

<table>
<thead>
<tr>
<th>Grading</th>
<th>No. of cases</th>
<th>Initial Treatment</th>
<th>Length of Follow Up and Further Treatment</th>
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</thead>
<tbody>
<tr>
<td>Borderline</td>
<td>5</td>
<td>Case 1, local excision (original diagnosis epitheliosis)</td>
<td>4 yr, alive and well</td>
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<tr>
<td></td>
<td></td>
<td>Case 2, local excision and radiotherapy</td>
<td>10 yr, alive and well</td>
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<td></td>
<td></td>
<td>Case 3, simple mastectomy</td>
<td>6 yr, alive and well</td>
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<tr>
<td></td>
<td></td>
<td>Case 4, radical mastectomy</td>
<td>8 yr, alive and well</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Case 5, radical mastectomy</td>
<td>12 yr, diet of coronary thrombosis</td>
</tr>
<tr>
<td>Intraduct carcinoma</td>
<td>7 (2 with Paget's disease of the nipple)</td>
<td>Case 6 local excision (original diagnosis epitheliosis)</td>
<td>12 yr, alive and well</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Case 7, local excision 1953</td>
<td>Further local excision 1958, extensive IDC. Simple mastectomy 1959 revealed invasive carcinoma. Alive and well 14 years after original biopsy</td>
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<td></td>
<td></td>
<td>Case 8, simple mastectomy (with Paget's disease)</td>
<td>5 yr, alive and well</td>
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<td></td>
<td></td>
<td>Case 9, simple mastectomy (with Paget's disease)</td>
<td>9 yr, alive and well</td>
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<td></td>
<td>Case 10, simple mastectomy</td>
<td>7 yr, alive and well</td>
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<td></td>
<td>Case 11, radical mastectomy</td>
<td>5 yr, alive and well</td>
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<tr>
<td></td>
<td></td>
<td>Case 12, radical mastectomy</td>
<td>8 yr, alive and well</td>
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</tbody>
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Comedocarcinoma is easily recognized (Fig. 9) but in certain other forms of intraduct carcinoma (Fig. 10) and lobular carcinoma in situ (Fig. 11), the cells filling the ducts may be remarkably uniform and mitotic figures scanty. In the absence of invasion of the surrounding tissues the malignant nature of the epithelium may not be recognized, especially when the number of ducts affected is small. In case 7 (Table II) of the present series carcinoma was confined to the ducts for at least seven years, and in the case described by Godwin (1952) there was a latent period of 12 years before lobular carcinoma in situ became invasive. Clearly the incidence of invasive carcinoma in any series will be increased if cases with malignant intraduct lesions are included.

It is my own experience, however, that it is more common for lesions which are not actually malignant to be diagnosed as intraduct carcinoma (Table I). In epitheliosis the proliferation is believed to involve both the epithelium lining the ducts and the myoepithelium which lies just within the basement membrane (Kuzma, 1943; Biggs, 1947). The presence of these two types of cell within the ducts may give an impression of cellular pleomorphism and malignancy.

Muir (1941) and Dawson (1948) both agree that a sequence can be traced from epitheliosis to intraduct carcinoma, and that it is impossible to state when malignancy actually occurs. Epitheliosis is not inevitably progressive, and Dawson (1948) holds the view that many of the cysts seen in chronic mastitis are due to the degeneration of epitheliosis with the formation of 'colostrum-like' foam cells which subsequently disintegrate (Fig. 12). Muir (1941) also pointed out that the number of cases in which the progression of epitheliosis to intraduct carcinoma could be traced was small, and suggested that often there was a more direct development of malignancy from duct epithelium without a preceding phase of epithelial hyperplasia.

Although the results of surgery are excellent when breast cancer is confined to the ducts (Gillis, Dockerty, and Clagett, 1960), the case for performing mastectomy for epitheliosis seems as yet unproven. My own views are in accord with those of Foote and Stewart who as long ago as 1945 stated that 'the rationale of simple mastectomy in "chronic cystic mastitis", so far as we can ascertain, has not yet been clearly demonstrated. The performance of this operation for "chronic cystic mastitis" should be discarded until specific lesions are proved beyond doubt to be followed by cancer in a sufficiently high percentage of cases to warrant this procedure.' Information regarding the long-term potential of epitheliosis to become malignant will only be obtained by following up a large number of women treated by local excision only for a very long time. Until then borderline cases and those showing severe epitheliosis present a clinicopathological problem which must be shared by surgeon and pathologist. Management of individual cases may have to take into account other factors, such as the extent of the disease in both breasts and the effect of a mutilating operation, possibly on a young woman.

I wish to thank the surgeons of the Westminster and Gordon Hospitals who allowed me to study their cases, and the Department of Medical Photography, Westminster Medical School, for the photomicrographs. My special thanks are due to Professor A. D. Morgan and Dr D. H. MacKenzie for their advice and encouragement.

REFERENCES

Obstet., 110, 555.