A composite malignant tumour of the elderly female breast

D. M. WAYTE, J. B. STEWART, AND C. G. McKENZIE
From the Royal Army Medical College, Millbank, London, and the Military Hospital, Colchester, Essex

SYNOPSIS A composite malignant tumour arising in the breast of an elderly woman is described. The cystic tumour containing areas of squamous metaplasia, bone formation, adenocarcinoma, and osteosarcoma was surrounded by the typical changes of mammary dysplasia (fibroadenosis).

The classification and acceptance of such tumours is highly debatable. There is no one acceptable classification of breast sarcomas and hence the prognosis of such neoplasms, particularly those containing heterologous tissues, is poorly defined. Evidence is presented in support of such composite tumours as being definite entities which arise from the closely associated epithelial and mesenchymal components of the breast simultaneously.

Sarcoma of the human breast, particularly when arising contiguously with an adenocarcinoma, remains a perplexing and controversial problem for both the surgeon and the pathologist (Hill and Stout, 1942). The great rarity of such malignant tumours and the inability to predict possible behaviour from the microscopic appearance partly account for the aura of scepticism which attends the diagnosis of a combined sarcoma and adenocarcinoma (Tudhope, 1939). There is always the doubt that the sarcomatous element is simply an undifferentiated spindle cell variant of the carcinoma (Stewart, 1950; Willis, 1958). But it is clear that the difficulties concerning the interpretation and acceptance of a combined or composite tumour can be accounted for by neglect of two basic features of mammary structure and pathology.

In the first instance it is well known that there exists a unique relationship between the epithelial component and its closely investing 'specialized' connective tissue. In the mammary dysplasia of fibroadenosis and in the benign fibroadenomata such a relationship is clearly seen. It is therefore not unreasonable to accept that on rare occasions both components may assume malignant change jointly and thereby produce the composite tumour. This argument is reinforced by the knowledge that the great majority of such tumours and breast sarcomas are considered to arise from fibroadenomas (Curran and Dodge, 1962).

The second feature that is often forgotten in regard to breast pathology is the extraordinary tendency for sarcomas of the breast to undergo metaplasia resulting in the formation of heterologous tissues which include bone, cartilage, and their malignant counterparts (Jernstrom, Lindberg, and Meland, 1963; Gonzalez-Licea, Yardley, and Hartmann, 1967).

We report the present case because of the rarity of the so-called mixed or composite tumours of the breast and to outline some of the controversy that exists concerning breast sarcomas. The composite tumour reported here contained a large squamous-epithelium-lined cyst, areas of bone metaplasia, and an osteosarcoma in combination with an adenocarcinoma.

Clinical Report

A woman of 76 sought advice regarding a painless swelling in the left breast. She stated that it had been present for four months, and claimed that she had had no previous trouble with the breasts.

On examination she was a rather obese elderly
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Fig. 1  Clinical photograph of the breast tumour. The pigmented lesions on the abdominal wall are seborrhoeic keratoses.

Fig. 2  Key diagram of the opened cystic tumour.

Fig. 3  Key diagram of the larger of the two pedunculated polyps.

Fig. 4  The opened cystic tumour. The two pedunculated polyps (the larger, arrow 1; the smaller, arrow 2), the small white sessile growths (arrow 3), and the rigid, hard lining to the cyst (arrow 4) are clearly seen.

A woman. The left breast was the site of a multinodular mass about 12.5 x 7.6 cm situated behind the nipple and extending over to the medial quadrant (Fig. 1). Lymph nodes were not palpable in the axillae or supraclavicular fossae. The overlying skin was adherent to the swelling and showed a bluish discoloration. The mass was at the apex of a rather pendulous breast and was freely mobile on the chest wall.

She had gravitational ulcers on both ankles and a mild degree of uterine prolapse but was otherwise apparently fit. A chest radiograph showed slight cardiac enlargement and congestive changes in the lungs but no evidence of metastases.

OPERATION
A left simple mastectomy without axillary dissection was performed on 30 October 1968.

The postoperative course was uneventful, the wound healing satisfactorily, and the patient was discharged on the 25th postoperative day.

When seen for follow up at three and six months there was no evidence of recurrence and a radiograph showed no chest metastases.

Pathology

MACROSCOPIC APPEARANCE
The mass in the amputated breast proved to be a tense cystic tumour 10.0 x 7.5 x 5.0 cm. On cutting into the tumour 200 ml of dark brown viscous fluid escaped, revealing a unilocular cyst parted by a narrow semilunar fold on its posterior wall. The cyst failed to collapse, having a rigid wall which was extensively calcified. Two pedunculated polypoid nodules with greyish-white irregular surfaces projected into the cavity of the cyst; the larger, 3.5 x 1.5 x 1.3 cm, hanging from the roof, the smaller, 2.0 x 1.8 x 1.7 cm, arising from the medial floor (Figs. 2 and 3). Both nodules were bony hard in consistency and impossible to section with a knife. In addition five small, white sessile cauliflower growths were scattered on the cyst lining ranging in size from a few millimetres to 1.0 cm in diameter. The
remainder of the cyst lining was smooth, white and glistening (Fig. 4). Contiguous with the upper lateral pole of the cyst there was a solid area of fibrofatty breast tissue. Lymph nodes were not identified in the specimen.

Numerous blocks were prepared from the cyst wall, the intracystic projections, and surrounding breast parenchyma.

METHODOLOGY

Tissue was fixed in 10% neutral formol-saline. Decalcification of the bony areas was carried out in 5% nitric acid (stabilized with 0.1% urea). Sections were stained with haematoxylin and eosin. The following special stains were also utilized: Masson's trichrome, Gordon and Sweet's method for reticulin, alcian blue for mucopolysaccharides, and Hale's colloidal iron techniques for acid mucopolysaccharides (with and without hyaluronidase).

MICROSCOPIC APPEARANCE

The main feature of the histology of this unusual breast tumour was the varied mixture of tissue it contained.

The smaller of the intracystic bony nodules was composed entirely of malignant tumour histologically indistinguishable from an osteosarcoma of a well differentiated type (Fig. 5). The base of the nodule consisted of squamous epithelium with an abrupt transition to a cellular stroma composed of a mixture of spindle and polygonal cells with hyperchromatic nuclei. Mitotic figures were fairly numerous. Multinucleated giant cells of the type common in osteoclastoma were prominent (Fig. 6) and islands of osteoid and osseous material lay uniformly throughout the stroma (Fig. 7).

Sections of the larger, downwardly projecting nodule revealed a mosaic of benign and malignant tissues (Fig. 8). The base was again formed of regular non-keratinizing squamous epithelium merging abruptly into a stalk composed of mature bone with well formed trabeculae and a benign cellular fibrous stroma (Fig. 9). Embedded in this stalk were two foci of undoubted adenocarcinoma. The adenocarcinoma was composed of irregular ducts, some crowded 'back to back' and sharing walls lined by a single cuboidal or columnar epithelium and solid areas of pleomorphic cells with little intervening stroma (Fig. 10). In parts there was an intracystic papillary structure (Fig. 11). Mitotic figures were numerous. Surrounding this carcinoma were islands of mature squamous epithelium, a few small cysts with squamous epithelial linings, and the mature bone composing the main tissue of the nodule (Fig. 12). At the apex, however, there was an abrupt transition from benign-looking bone to malignant osteoformative tissue, the appearances being identical.
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Fig. 7 Osteogenic sarcoma. Osteoid, bone, and mitotic activity are clearly seen. Haematoxylin and eosin × 400.

Fig. 8 An area of the larger pedunculated polyp displaying squamous epithelium, bone, and adenocarcinoma. Haematoxylin and eosin × 40.

Fig. 9 Surface squamous epithelium and bone of the larger pedunculated polyp. Haematoxylin and eosin × 140.
with the sarcoma in the companion but smaller projection (Fig. 13).

Sections from the cyst wall revealed dense fibrous tissue lined by non-keratinizing squamous epithelium which was continuous with the bases of the tumour projections and which became hyperplastic and papillomatous to form the small cauliflower excrescences noted in the gross specimen (Fig. 14). Beneath the lining epithelium were thin plaques of calcification and a few fragments of benign osteoid.

The tissue immediately surrounding the cyst consisted of compressed atrophic parenchyma. The dense fibrofatty area adjacent to the cyst showed florid fibrocystic disease—dilated ducts lined by hyperplastic epithelium (mild epitheliosis), duct papillomatosis, apocrine metaplasia, fibrosis, and chronic inflammatory cell infiltration. There was no microscopic evidence to suggest that the cystic tumour had originated in a fibroadenoma. There was no malignant tissue in the breast parenchyma outside the cystic tumour.

Discussion

Early in foetal life the parenchyma of the breast develops as a result of downward growth of finger-like epithelial processes from the overlying ectoderm. As the epithelial tracts progress and proliferate each extension becomes closely invested by a specialized connective tissue. This special relationship between epithelium and mesenchyme can be seen during all the hormonal vicissitudes of mammary tissue and may also be demonstrated in the common benign pathological conditions of the breast.

Fibroadenosis with its multiplicity of microscopic features is undoubtedly the commonest pathological lesion of the human female breast. Occasionally cyst formation may predominate. The epithelial lining of such cysts most commonly undergoes pressure atrophy but on occasion proliferation, papillary formation, and even intracystic carcinoma can be seen. That a relationship exists between fibroadenosis and breast carcinoma is now accepted by most authorities (Davis, Simons, and Davis, 1964) but any such relationship between fibroadenosis and the less common breast sarcoma has not been defined. However, a distinct relationship between the fibroadenoma and sarcoma is definitely established (Curran and Dodge, 1962). A fibroadenoma, most commonly arising in the breast of the young adult woman, may be considered as a local area

Fig. 10 Infiltrating adenocarcinoma and an area of bone within the larger pedunculated polyp. 
Haematoxylin and eosin × 140.
of mammary dysplasia which is possibly hormone dependent. When seen in the older woman such growths are more likely to represent but one variant of a generalized picture of fibroadenosis and therefore separation between fibroadenoma and fibroadenosis is less clearly possible in this age group.

Giant fibroadenomas (benign cystosarcoma phyllodes) present most commonly in the elderly female. A history of recent rapid growth is often given by the patient and such a statement often results in consideration of a malignant process although on microscopic examination the majority of such tumours have a benign structure (Oberman, 1965a). The great percentage of giant fibroadenomas have a solid macroscopic appearance with only occasional clefts separating the constituent lobules but occasionally the tumour assumes a cystic form, the intra-canaliculic projections of stroma and epithelium remaining as polypoidal projections into the cystic cavity.

That the stroma of a fibroadenoma may undergo metaplasia to form heterologous tissue is well known (Robb and Macfarlane, 1958). Less well known is the fact that in rare instances the epithelial component may also undergo metaplasia to that of a squamous type (Salm, 1957; Willis, 1962).

Tumours of the canine (Fidler and Brodey, 1967) and feline breast (Schmidt and Langham, 1967) often show metaplasia of the stroma resulting in the formation of bone and cartilage (Allen, 1940; Willis, 1967). Such a phenomenon is rare in the human breast and when seen involves the stroma of benign or malignant fibroadenomas (Rottino and Howley, 1945; Willis, 1967). The histogenesis of osseous metaplasia in close approximation to epithelial tissues is poorly understood (Collins and Curran, 1959) but may be seen in the gastrointestinal and genitourinary tracts (Pang, 1958) and in certain tumours of the skin and salivary glands (Yates and Paget, 1952).

At these various sites it has been proposed that the epithelium acts as an 'organizer' in stimulating the surrounding mesenchymal cells to lay down bone. Experiments with transplanted urinary (Constance, 1954) and gallbladder epithelium have resulted in the formation of bone within the mesenchyme adjacent to the transplant (Huggins, 1931). In an attempt to relate this process to particular epithelial cells, Azzopardi and Smith (1959), in their study of salivary gland neoplasms, proposed that the myoepithelial cell was responsible for the production of the connective tissue mucins in which bone could be formed.

Occasionally giant intracanalicular fibroadenomas are found to be histologically malignant (malignant cystosarcoma phyllodes), the mesenchymal component being most commonly implicated to form a spindle cell sarcoma. Stewart (1950) pointed out that 'a very small number of these tumours develop bizarre struc-
Fig. 13. Osteoid formation within an area of osteosarcoma seen in the larger polyp. Note the mitotic figure. Haematoxylin and eosin × 400.

Fig. 14. The abundant squamous epithelium composing the small sessile 'cauliflower-like' excrescences. Note the bone formation and the vascular channels in the underlying cyst wall. Haematoxylin and eosin × 40.

atural patterns with the formation of atypical bone and cartilage' while Willis (1959) considers that such metaplastic tissue may be malignant and form osteosarcomas, chondrosarcomas, and osteoclastomas.

The classification of breast sarcoma is a confusing topic (Norris and Taylor, 1968). All authors accept the so-called 'pure' sarcomas as definite entities but it is in regard to the sarcomas containing an epithelial component where disagreement exists. Botham, McDonald, and Claggett (1958) considered that lesions containing an epithelial component intermixed with a rich cellular stroma should be excluded, while other workers (Oberman, 1965b; Lattes 1967) considered that carcinomas associated with malignant metaplasia of their surrounding stroma (composite tumours or combined tumours) should also be excluded from any series of breast
sarcoma. However, the majority of workers in this field accept the possibility of intermixture of components in mammary gland neoplasms, and, although there is no clear uniformity, the following categories of breast sarcoma are commonly listed:

'PURE' SARCOMA
These are (1) sarcomas not specifically related to mammary tissue, e.g., malignant lymphoma; (2) sarcomas arising from general breast mesenchyme, e.g., liposarcoma and haemangiosarcoma; (3) sarcomas not appearing to arise from fibroadenomas and having a uniform structure.

COMPOSITE TUMOURS
Sarcomas arising from a fibroadenoma, e.g., malignant cystosarcoma phylloides, are often composite tumours.

Certain authorities clearly separate sarcomas of the malignant cystosarcoma phylloides type from the diffuse pure sarcomas not seen to arise from a fibroadenoma, for the former rarely metastasizes, thereby differing from the more aggressive pure sarcoma. However, in a number of instances it may not be possible to demonstrate such an origin owing to the destructive and invasive nature of the neoplasm (Fawcett, 1967).

The great majority of sarcomas of the breast occur in elderly women. They are extremely rare (Botham et al., 1958; Hill and Stout, 1942) and in a 10-year period formed only 0·9% of all malignant breast tumours seen at one hospital (Curran and Dodge, 1962), and stromal sarcomas accounted for only 0·6% of 5,458 malignant breast tumours in the series of Kennedy and Biggart (1967).

In regard to the sarcomas which have undergone metaplasia, the case described by Jernstrom et al. (1963) was the only osteosarcoma in 3,309 malignant neoplasms of the breast collected in 18 years. It is clear, therefore, that osteosarcoma of the breast is a distinct rarity, and some authors (Lattes, 1967) do not accept such an entity and regard the examples of bone and cartilage formed by malignant tumours of the breast as probably carcinomas with osseous and cartilaginous metaplasia. Smith and Taylor (1969), in their recent series of 35 cases from the files of the Armed Forces Institute of Pathology, regard most of the heterologous bone and cartilage seen in breast tumours to arise from stromal metaplasia but also accept that certain adenocarcinoma cells may undergo direct transition to form bone- and cartilage-forming cells. Willis (1959) regards the majority of predominantly cartilaginous, bony and osteoclastic mammary tumours as metastatic variants of the 'cystosarcoma' group and advocates their segregation as a subgroup of mammary sarcoma because of their relatively frequent malignancy.

When an adenocarcinoma develops jointly with a metastatic breast sarcoma a most bizarre picture (Willis, 1967), as is seen in our present case, results. This intermixture of an epithelial and mesenchymal neoplasm must undoubtedly raise many difficulties in interpretation and in classification. It should be pointed out, however, that such composite or mixed malignant tumours are commonly seen in the canine and feline breast and metastases of both malignant elements have been described.

Hill and Stout (1942) postulated that there were three possible ways for a composite malignant tumour to develop: (1) malignant transformation of an epithelial and a mesenchymal component of a teratoma; (2) a primary carcinoma inciting the surrounding stroma to malignancy; (3) a malignant stroma inciting the adjacent epithelium.

Possibly a fourth postulate should be added to the above list which would explain the multiple changes seen in our present case. As indicated earlier in this discussion, a close association between epithelium and mesenchyme of the breast definitely exists and it seems not unreasonable to consider the possibility of an inciting agent inducing metaplasia and neoplasia of both components simultaneously.

The prognosis of mammary tumours containing osteoid tissue remains conjectural due to the small number of cases seen in any one series. Rottino and Howley (1945) pointed out the danger of incomplete removal by local enucleation techniques but considered that simple mastectomy was normally adequate because of the very low incidence of lymph node metastases.

References


