Oestrogen and progesterone receptor distribution in the cancerous breast

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Summary  To test the hypothesis that steroid hormone receptor expression is particularly pronounced in breast tumours when compared with non-neoplastic tissue, mastectomy samples were divided into 16 sectors. Multifocal tumours, of varying receptor phenotype were found in 4 patients and in addition different regions of large tumours also showed varying receptor contents. Remaining samples were found to consist of normal tissue, with fat, connective tissue and some benign breast disease. In the 9 patients with oestrogen receptor positive tumours (ER), ER content was invariably much greater in the tumours than in the remainder of the breast. Progesterone receptor (PR) content was not closely related to ER, and was lowest in the poorly differentiated tumours. This relation to differentiation was not seen in ER. The data support the view that ER concentration in ER positive tumours may reflect the transformed nature of neoplastic tissue.

The significance of the presence of oestrogen and progesterone receptors (ER & PR) in breast tumours has been emphasized by many authors because of its implications for the progression of the disease and its management (Hubay et al., 1984; Sebert & Lippman, 1982). It is less clear how the expression of receptors in tumour relates to the incidence in normal tissue. Some studies suggest that receptor content is greatest in malignant and proliferative benign breast disease (Allegra et al., 1979; Jacquemier et al., 1982) whereas in normal breast tissue ER is virtually undetectable (Jacquemier et al., 1982; Terenius et al., 1974). ER is raised in puberty and pregnancy but never to levels reached in carcinoma (Israel & Band, 1984). Even in ER positive breast cancer there is no significant binding in the histologically normal surrounding tissue (Johansson et al., 1970). Indeed Israel and Band (1984) developed the hypothesis that the presence of ER in malignant disease is a direct consequence of neoplastic transformation and oncogene expression. This hypothesis has recently received considerable support from the characterisation of oestrogen receptor cDNA, which shows structural similarities to known oncogenes (Green et al., 1986). To our knowledge, a systematic comparison between the receptor content of a tumour and the whole of the remainder of the breast has not been reported. Therefore this paper reports the results obtained from such an analysis of nine mastectomy specimens with ER positive tumours.

Materials and methods

Patients

In all cases a preoperative attempt was made to confirm the presence of malignancy either by aspiration cytology or by Trucut biopsy. Where malignancy was not confirmed, excision biopsies were performed for frozen section at the time of mastectomy. Ischaemic time, which might affect receptor values (Leighton et al., 1984; Teichner et al., 1985), was minimal in the operative procedure used. The breast was left attached to the axillary structures until fully mobilised, when it was removed between clamps and only following sampling for receptor analysis and histology was the required axillary procedure carried out. The removed breast was divided into 16 equal sectors (Figure 1) and specimens at least 1 cm² in size were removed from the centre of each sector. Grossly visible fat having been removed, the tissues were divided into two equal halves and used respectively for receptor assay and histological examination.

Specimens were collected from the posterior aspect of the breast and always from medial to lateral and cranio-caudally (i.e. A1, A2, . . . D4, cf. Figure 1). Collection was completed within ten minutes following clamping.

Receptor assay

Tissue specimens were placed immediately in liquid nitrogen, transported to the laboratory and stored in liquid nitrogen until processed.

Tissue preparation

Low speed tumour supernatants (2000 g) were prepared by the disemembration-centrifugation method (King et al., 1979). Briefly, frozen tissue was pulverised in a Braun Mickrodismembrator for 70 sec. The resultant powder was resuspended in 50 mm phosphate buffer (containing 30% glycerol, 1.5 mm EDTA and 10 mm monothioglycollate at pH 7.4), to give protein concentrations of ~2.5 mg ml⁻¹, mixed and allowed to stand for 15 min. After centrifugation (2000 g, 15 min) the supernatant was used for receptor analysis. For ER, duplicate aliquots of tissue supernatant (100 µl) were incubated with a single concentration of 3H-oestradiol (50 nm) for 18 h at 4°C. Duplicate tubes were incubated with the further addition of a 100 fold excess of diethylstilboestrol (DES). The incubation volume was the same in both sets of tubes. Free and bound steroid were separated by dextran coated charcoal (DCC) extraction. A suspension of DCC (200 µl of 0.25% charcoal 0.025% dextran in 10 mm Tris 1.5 mm EDTA buffer, pH 7.4) was

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added to each tube, mixed and incubated for 10 min. After centrifugation (5 min at 10,000g) portions of supernatant (200 μl) were added to 8 ml liquid scintillation cocktail (4 g PPO, 0.05 g POPOP in toluene). Steroid was extracted into the organic phase by vigorous shaking, and radioactivity measured in a Beckman LS7500 liquid scintillation spectrometer with correction for quenching in order to obtain dpm. Counts from the tubes containing competitor (non-specific binding) were subtracted from those of tubes without competitor (total binding) to give values for hormone specifically bound to receptor. Receptor concentration was calculated from these values by using the specific activity of the titrated steroid, and expressed as fmol mg⁻¹ protein. Values were means of duplicate estimations. For PR, similar methods were employed, with 3H-progesterone (100 nM) as the labelled ligand, and norethindrone (100 fold excess) as the unlabelled competitor. In PR analysis, cortisol (100 fold excess) was also added to displace the labelled ligand from transcortin. Protein estimations (Lowry et al., 1951) were carried out with bovine serum albumin as the standard.

Validation of the single saturating dose receptor assay

The single saturating dose (SSD) receptor assay has been validated in several ways: (i) sequential estimations of ER in a control uterine cytosol preparation gave an interassay coefficient of variation (COV) of 17% (n = 34); (ii) the intra-assay COV was 6% (n = 6); (iii) comparison between the SSD and the Scatchard methods of ER measurement gives an excellent correlation (r = 0.97; n = 50; P < 0.001; Puddefoot et al., 1986); (iv) sequential assay of PR in uterine cytosol preparations gave an interassay COV of 12.5% (n = 33). Shortage of suitable material and variable non-specific binding precluded comparison between the SSD and Scatchard methods in this case. However, quality control exercises in which tumours were also assayed by another laboratory gave good agreement between laboratories (Puddefoot et al., 1986).

Histopathology

Samples were fixed in buffered formalin and processed to paraffin in the usual way. Sections of 5 μm were examined in haematoxylin and eosin preparations.

Results

Data from 9 patients whose primary tumours were ER positive were obtained. Of these, two were premenopausal (cases 2 and 17, aged 38 and 49), one perimenopausal (case 6, aged 53), and the remaining six were postmenopausal subjects aged 46–75. Oestrogen receptor data and histological findings are illustrated in Figure 2. In all cases, the ER content was much higher in the ER positive tumours than in the nonmalignant specimens (see Table I). In one case (17) three distinct tumours were all ER positive/PR positive. In 4 patients, (cases 4, 5, 8 and 17) multifocal tumours were detected which on subsequent analysis were
Table I  Tumour ER content in nine ER positive tumours, maximum ER concentration in non-cancerous samples from the same patients. For comparison, PR concentrations for the same samples are also given. (cf. Figures 1, 2). Receptor concentration values are given as fmol mg\(^{-1}\) protein. Bloom and Richardson gradings are also given: w = well differentiated, m = moderate differentiation, p = poor differentiation.

| Case no. | 1  | 2  | 4  | 5  | 6  | 7  | 8  | 10 | 17 |
|----------|----|----|----|----|----|----|----|----|----|
| Age      | 70 | 38 | 60 | 64 | 53 | 59 | 75 | 46 | 49 |
| ER values: | Non-tumour tissue | 36 | 52 | 10 | 17 | 18 | 6  | 21 | 51 |
|          | Tumours             | 304| 93 | 37 | 97 | 196| 121| 495| 318| 42 |
| PR values: | Non-tumour tissue | 38 | 70 | 10 | 3  | 0  | 6  | 0  | 43 | 38 |
|          | Tumours             | 37 | 95 | 20 | 7  | 62 | 17 | 303| 9  | 34 |
| Differentiation | m  | m | m | p | w | p | w | p | w |

found to be distinct from each other by receptor content, relative position or by histological criteria. Additionally, receptor expression varied in the different regions of the large tumours in patients 1 and 10. In case 5, two distinct tumours were identified preoperatively on xeromammography. The nonmalignant tissue was by no means homogeneous, and contained areas of benign breast disease, as well as normal tissue. However, there were no differences between the receptor contents of the different types of nonmalignant tissue. PR data (Figure 3) shows no correlation between malignancy and receptor expression, or indeed between ER and PR.

Using the criteria of Bloom and Richardson (1957) three tumours were classified as well differentiated, three as moderately differentiated and three were poorly differentiated. In 4 cases, in-situ carcinoma was seen in the breast ducts. There was no correlation between tumour grade and ER, although the least differentiated tumours had the lowest PR values.

Discussion
Attempts have been made to correlate ER levels with various morphological and other criteria by which breast tumours are classified, but these have produced variable results (for recent review, see Seibert & Lippman, 1982). There may be several reasons why this should be. Most importantly, it is clear that the disease, as judged by hormone receptor content, takes several forms. Most obvious is the well documented recognition of at least four phenotypes for soluble (previously termed 'cytoplasmic', Walters, 1985) receptor status, i.e., ER positive/PR positive, ER positive/PR negative, ER negative/PR positive, ER negative/PR negative. Additional variants are recognised when nuclear-bound receptors are also considered. Further evidence for heterogeneity also emerges when receptor affinities are taken into account (Puddifoot et al., 1986). Moreover, data accumulated in the last few years has shown quite clearly that such heterogeneity is even to be found in the tumour tissue of individual patients. For example, serial samples from an individual tumour may show extremely different

![Figure 3a-i](image-url) Progesterone receptor (PR) content for the same samples as in Figure 2. Key also as in Figure 2. Values for PR content are expressed as fmol mg\(^{-1}\) protein. The PR distribution is not closely related to ER (cf. Figure 2). In particular, the tumour is not always relatively rich in PR compared with the remainder of the breast.
receptor contents (Tilley et al., 1978; Braunsberg, 1975; Silfversward et al., 1980; van Netten et al., 1985). Recently, too, the development of monoclonal antibodies to receptors has made possible the application of immunohistochemical techniques which in some cases show adjacent ER positive and ER negative tumour cells (King et al., 1985).

The present results confirm and extend these findings. One striking observation is that patients show extraordinary diversity in the nature of their disease. Not only do nearly all show benign breast disease as well as cancer (Figures 2, 3) but in addition some have multiple tumours. In case 17, all three were ER positive/PR positive, but other cases show multifocal tumours of more than one soluble receptor phenotype. For example, case 8 has both ER positive/PR positive and ER positive/PR negative tumours, while case 4 has an ER positive/PR negative and an ER negative/PR positive tumour. Case 5 combines two distinct ER positive/PR positive tumours with, remarkably, a second phenotype (ER negative/PR negative). This probably means that there is no simple relationship between receptor phenotype and oncogene expression. On the other hand, it is very clear that in ER positive tumours, the receptor concentration is generally vastly in excess of that found in the remainder of the tissue (see Table 1). The same relationship is not unerringly found for PR. In some cases PR content of PR positive tumours is higher than in the other tissue (e.g. cases 1 and 2, see Figure 3) although comparison with Figure 2 shows the effect is not so marked as for ER. In other cases (e.g. 10 and 17) highest PR values are found in non-tumour tissue. There would appear to be no relationship between ER status and tumour differentiation (Table 1), although it is possible that poorly differentiated tumours have low PR content.

It is obviously difficult to be sure that the material divided between histopathologist and biochemist is always quite homogeneous. In this connection some samples of relatively high receptor content were found in fat or connective tissue, (e.g. cases 1, 10, Figure 2). Here it may be suspected that the samples were not identical, and those taken for receptor analysis may have contained tumour cells in addition to normal tissue.

With this caveat in mind, however, it is nevertheless clear that the data supports the hypothesis that ER concentration in ER positive tumours reflects the transformed function of neoplastic tissue (Israel & Band, 1984). Following other work (Green et al., 1986), it also suggests that the expression of the oncogene involved may be related to ER functions.

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