Short Communication

Histochemical detection of oestrogen receptors in breast carcinoma: A successful technique

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The heterogenous nature of breast tumours (Hawkins et al., 1977; Silfversward et al., 1980) complicates the application of oestrogen receptor assay results to the management of breast cancer. Much effort has therefore been expended on the development of histochemical assays to detect receptors in tissue sections, initially with oestrogen-protein conjugates (e.g. Pershchuk et al., 1979; Lee, 1978 and Walker et al., 1980) and more recently with antibodies against the receptor protein, both polyclonal (e.g. Lope-Pihie et al., 1985; Tamura et al., 1984) and monoclonal (Greene et al., 1980; Coffer & King, 1981).

Few of these methods have been universally accepted or validated in other centres (e.g. McCarthy et al., 1980; Chamness et al., 1980). In our experience, too, both published (Penney & Hawkins, 1982) and unpublished, we have previously been unable to correlate histochemical and biochemical assays. We therefore feel it is important to report when a histochemical technique does correlate with our established biochemical assay.

One of the Greene-Jensen antibodies generated against oestrogen receptor from MCF-7 breast cancer cells is now marketed in the form of a kit (ER-ICA) by Abbott Laboratories Ltd, and in our hands, the results, in agreement with other reports, are very promising.

Thirty-four breast cancers were received on ice from operating theatres in the Edinburgh and Fife regions within ~1 h of excision. A portion of tumour was cut from the face of the tissue and fixed in formal-saline for routine histopathology, a second portion (~350 mg) was used immediately for biochemical assay of oestrogen receptor activity by a standard DCC assay used in these laboratories since November 1973 with one minor modification (Hawkins et al., 1981) and a third smaller portion (50–100 mg) was frozen in liquid nitrogen until assayed histochemically (within 12 weeks). For the latter, ~4 µm frozen sections were cut, fixed, incubated with either anti-receptor antibody (rat IgG; ‘test’) or a control antibody (normal rat IgG) and stained by a peroxidase-antiperoxidase technique, according to the manufacturer’s instructions (Abbott Laboratories, The Business Centre, Molly Millar’s Lane, Wokingham, Berks RG11 2QZ), with particular care being taken to avoid drying-out of the sections. In brief, the tissue sections were fixed by immersion in formaldehyde (3.7%) in PBS for 10 min, in methanol at −20°C for 2 min and in acetone at −20°C for 1 min, prior to incubations at room temperature with normal goat serum (15 min), rat antibody (30 min), ‘bridging’, goat anti-rat antibody (30 min), peroxidase-rat anti-peroxidase complex (30 min), and staining with diaminobenzidine plus hydrogen peroxide (6 min). When the peroxidase-staining reagents (DAB and phosphate buffer – H₂O₂) ran out, the last few specimens were stained using our own preparation of DAB solution (1 mg ml⁻¹ in tris-HCl buffer with 0.2% H₂O₂) containing imidazole (0.01 M). In each run, at least one known, high receptor-positive specimen was also processed (either the ‘control’ cell slide provided by the manufacturer or a breast cancer from our routine assays). Judging from these control specimens, there was a little variation from run to run. No striking effect of incorporating imidazole was apparent, although staining intensity was slightly increased.

Staining, when present, was confined to the epithelial cell population and was virtually all nuclear. The sections, it must be noted, did show some cellular distortion, possibly due to the triple fixation step employed. Within most sections, there were both stained and unstained cells, plus some variation in the intensity of staining amongst the positive cells. In order to quantify staining, each specimen was scored independently by each of 3 observers for (a) the cellularity (% of specimen occupied by tumour cells), (b) the proportion (%) of cells staining and (c) the average intensity of staining (assessed on an arbitrary scale of −, +, ++, or +++).

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The results for the 34 breast cancers are shown in Figure 1, where the histochemical assay result (staining intensity) is plotted against the biochemical assay result (fmol receptor sites mg\(^{-1}\) protein). Of the tissues, 27 (i.e. 79\%) were biochemically receptor-positive and 7 (21\%) were receptor negative (i.e. <5 fmol sites mg\(^{-1}\) protein). Biochemically, 25 (i.e. 74\%) showed some staining whilst the remaining 9 tissues, all with biochemically determined receptor concentrations of <10 fmol mg\(^{-1}\) protein, showed no detectable trace of staining. There was a strong correlation between either the intensity of staining (Figure 1) or the percentage of cells staining (Figure 2) and biochemical receptor site concentration, the correlation coefficients being +0.85 and +0.87 respectively (Spearman’s Rank Correlation Test, \(n=34\)).

In this small series of tumours, biochemically-determined receptor concentration on a protein basis (fmol sites mg\(^{-1}\) protein) was not significantly related to the cellularity of the specimen assayed (27 receptor-positive tissues only, correlation coefficient = 0.18, NS), though on a wet weight basis, receptor concentration (fmol mg\(^{-1}\) wet tumour) was related to tumour cellularity (27 receptor-positive tissues only, correlation coefficient = 0.40, \(P<0.05\)) as we have previously shown (Hawkins et al., 1977, Masters et al., 1978, Hawkins et al., 1981) and as might be expected from the localisation of receptors in the epithelial cells of the tumour.

![Graph showing correlation between histochemical and biochemical receptor sites](image1)

**Figure 1** The correlation between the histochemical staining for oestrogen receptors ('staining intensity') and receptor concentration as determined by a standard biochemical assay in 34 breast cancers. Staining intensity represents the mean of assessments by 3 independent observers on an arbitrary scale of 0, +, ++ and ++++. \(r=+0.85\).

![Graph showing correlation between cells staining histochemically and receptor sites](image2)

**Figure 2** The correlation between the percentage of cells staining histochemically for oestrogen receptors and receptor concentration as determined biochemically in 34 breast cancers. Percentage of cells staining represents the mean of assessments by 3 independent observers. \(r=+0.87\).

In the present study, although the tumour cellularity differed between the portion selected for histochemical assay and that selected for biochemical assay, the differences were slight in 33 out of 34 tumours and severe in only one case (7\% vs. 37\%). In order to facilitate comparison of histochemical and biochemical results, a 'staining index' (representing the total staining in the specimen) was calculated as follows:

\[
\text{Staining index} = \text{Staining intensity} \times \frac{\% \text{ cells staining}}{100} \times \frac{\% \text{ cellularity}}{100}
\]

(This was calculated using the cellularity of the biochemical specimen to minimise discrepancies due to the use of non-identical portions of tissue in the two assays). Staining index, too, was strongly correlated with biochemically determined receptor site concentration (correlation coefficient = +0.87, \(P<0.001\)) as shown in Figure 3.

These results strongly suggest that the histochemical assay described detects accurately the classical oestrogen receptor as determined by biochemical binding assays. Although others (King et al., 1985; Thorpe et al., 1985; Pertschuk et al., 1985; McLellan & Coombes, 1985; Harper et al., 1985) have previously reported such a correlation, we felt it important to report our confirmatory findings after our previous disappointing experiences with other assays of this type. From a clinical point of view, although it may be difficult
to quantify histochemical assays precisely, the assay, in our hands, starts to detect receptor activity at concentrations around 20–40 fmol biochemical sites mg⁻¹ protein; this cut-off is fairly close to that used routinely in this department for treatment decisions (20 fmol mg⁻¹ protein) and is very 'safe' as judged by other reports in relation to treatment of advanced disease (Cant et al., 1985) or adjuvant therapy (Rose et al., 1985; Stewart & Prescott 1985). That the assay is as good as, or better than, the biochemical procedure in identifying responders to endocrine treatment in advanced disease has already been demonstrated directly by McLelland and Coombes (1985).

We consider these preliminary results to be successful and believe that the technique shows great potential for (i) assay of small specimens, (ii) identification of tumours with a heterogeneous population of cells with respect to receptor status and (iii) application to cytological aspirates.

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