Comparison of new immunohistochemical assay for oestrogen receptor in paraffin wax embedded breast carcinoma tissue with quantitative enzyme immunoassay

G Saccani Jotti, S R D Johnston, J Salter, S Detre, M Dowssett

Abstract

Aim—To validate the use of a new mouse monoclonal antibody (1D5) directed against the N-terminal domain (A/B region) of the oestrogen receptor in an immunohistochemical assay (ER-IHA) for paraffin wax embedded tissue. Methods—Breast cancer specimens were surgically obtained from 119 previously untreated patients. For comparison, oestrogen receptor was measured from cytosol fractions using an established oestrogen receptor enzyme immunoassay (ER-EIA) method. Oestrogen receptor “H-scores” were obtained from the ER-IHA after antigen retrieval using microwave treatment. Where discrepancies occurred between the two methods, further immunohistochemistry was performed using the H222 antibody from the Abbott Laboratories ER-ICA kit. Results—The correlation between the two methods was non-linear, but despite this there was an 86% concordance between ER-EIA and ER-IHA using the 1D5 ER antibody. Fifty four per cent of tumours (64/119) were oestrogen receptor positive and 32% (38/119) were negative by both assays. A mismatch between the ER-EIA and ER-IHA occurred in 17 cases. Seven tumours were IHA positive but EIA+, but five of these were borderline negative by EIA, having values of >5 and <10 fmol/mg protein. Ten tumours were IHA negative and EIA+; four of these tumours were completely negative by IHA in the section studied. A further IHA assay, carried out on the 17 tumour mismatches with H222 antibody, showed that three tumours remained substantially discordant. These three tumours were strongly positive with the 1D5 antibody and negative with the H222 antibody. Two of these discordant tumours were of the rare ER negative and PgR positive phenotype and may contain oestrogen receptor that is of biological interest but which lacks the hormone binding epitope. Conclusions—The concordance between the classic enzyme immunoassay technique and the new immunohistochemical method on paraffin wax embedded sections was good. Moreover, the IHA technique using the 1D5 antibody against the N-terminal was easily reproducible. This technique may allow oestrogen receptor content to be determined in large cohorts of patients in whom archival tumour material is available.

Oestrogen receptor is the biochemical marker most frequently measured in breast cancer. The presence of oestrogen receptors in breast cancer is an indicator of good prognosis, and tumours rich in oestrogen receptor have a 60 to 70% chance of responding to first line endocrine treatment. As a result of this, the choice of treatment for individual patients may be influenced by oestrogen receptor content. New methods of measurement must therefore be subject to close scrutiny and optimisation.

The monoclonal antibody H222, which is directed against the hormone binding region close to the C-terminal end of the human oestrogen receptor, has been used to quantify the expression of oestrogen receptor in homogenates of breast cancer by immunochromical means using a sandwich enzyme immunoassay technique (ER-EIA). This enzyme immunoassay has increasingly superseded the ligand binding assay (ER-DCC) because of its convenience and superior reproducibility between and within laboratories. In recent years it has tended to be the yardstick against which evolving methodologies have been compared. When clinical specimens are too small for biochemical assay, the H222 antibody has been used to probe frozen tissue sections in an immunocytochemical assay (ER-ICA). This approach allows the normal and benign positive staining elements in tumours to be excluded from assessment, a potential source of false positive results in biochemical assays. ER-ICA assays can be cost effective and quick to perform, although until recently have been applicable only to frozen tissue sections.

The availability of a reliable immunohistochemical assay on paraffin wax embedded tumours (ER-IHA) would be of great value in studies of archival material. Antigen retrieval by enzyme predigestion of tissue sections has made oestrogen receptor determination technically possible in paraffin wax embedded breast cancers which have been fixed in formalin. The conditions required for antigen retrieval depend on the fixative, fixation period, and the processing temperature. The
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use of the H222 antibody in this context has met with varying degrees of reported success, and inconsistencies between and within laboratories have been common. The lack of consistency could be due to differing pronase activity between batches, although with optimisation of predigestion techniques, individual laboratories have got the H222 antibody to work well and consistently.

Because of these problems, however, we investigated microwave pretreatment as an alternative and more reliable method of antigen retrieval, in addition to using a new anti-oestrogen receptor antibody (clone 1D5), directed against the N-terminal region of the oestrogen receptor (Dako), which is considerably cheaper than H222.

Methods

The study was carried out on 119 untreated primary breast carcinomas. Mastectomy or excision biopsy specimens were delivered fresh immediately after resection, and were processed immediately.

A portion of tumour, about 200 mg in size, was snap frozen in liquid nitrogen immediately after dissection from the breast, and stored at −80°C before analysis for oestrogen receptor using enzyme immunoassay. The remainder of the tumour was fixed for about 24 hours in 10% buffered formalin. The tissues were embedded in paraffin wax following a routine processing procedure which did not exceed 60°C. As far as possible, the portion of tumour taken for ER-EIA was immediately adjacent to the paraffin wax embedded tumour from which sections were cut.

Sections of 3 μm were cut from the paraffin wax embedded tumours on to slides coated with APES (3-aminopropyltriethoxysilane; Sigma). The sections were air dried overnight in an oven at 37°C, and one section stained with haematoxylin and eosin for light microscopic assessment.

OESTROGEN RECEPTOR ENZYME IMMUNOASSAY (ER-EIA)

Before cytosol preparation, fat, blood, and necrotic tissue were discarded from each sample and a piece of tumour between 0.1 and 0.2 g was taken. The tissue was finely chopped, weighed, and transferred to the Teflon container of a microdismembrator (Braun Medical Ltd.). The tissue was pulversed for one minute with liquid nitrogen for one minute. The powdered tumour was reconstituted 1:8 weight:volume in iced TRIS/molybdate buffer (5 mM sodium molybdate, 10 mM monothioglycerol, 1 mM dipotassium chloride EDTA, 3 mM sodium azide, and 10 mM TRIS, pH 7.4). The homogenate was centrifuged at 4°C for 20 minutes at 2000 × g after which the cytosol fraction was removed and diluted 1 in 5 in TRIS/molybdate buffer for protein assay using the Bio-Rad method with a bovine gamma globulin standard.

An aliquot of the cytosol was diluted to give a protein concentration of 1 to 2 mg/ml. The oestrogen receptor concentrations in the diluted cytosols were determined using the ER-EIA kit from Abbott Diagnostics according to the manufacturer’s instructions and values ≥ 10 fmol/mg protein were regarded as positive.

1DS ER IMMUNOHISTOCHEMICAL METHOD (ER-IIH-1DS)

The paraffin wax embedded sections were warmed on a hot plate and then dewaxed in xylene and rehydrated through a descending series of alcohols. Unless otherwise stated all the following reagents were obtained from Dako Laboratories and incubations carried out at room temperature. The sections were placed in a plastic dish filled with 10 mM citrate buffer, pH 6.0, and microwaved (750 watts) on full power for two five minute intervals, ensuring that the sections did not dry out at any time. The sections were then washed in tap water and phosphate buffered saline (PBS, pH 7.4) and incubated in 10% hydrogen peroxide/PBS for 15 minutes, and in a 1 in 5 dilution of normal rabbit serum for five minutes. The normal rabbit serum was tapped off and the sections were then incubated for two hours with a 1 in 100 dilution of monoclonal anti-human ER antibody 1D5. The sections were washed and incubated with a 1 in 100 dilution of biotinylated rabbit anti-mouse immunoglobulin for 45 minutes. After washing in PBS, the sections were incubated with a 1 in 200 dilution of horseradish peroxidase conjugated streptavidin complex for one hour. A solution of 0.05% 3,3’ diaminobenzidine (Sigma) dissolved in dimethyl formamide, plus 100 μl of 30 volumes hydrogen peroxide/100 ml PBS, was used to develop the peroxidase activity in the sections for 10 minutes. The sections were then washed in running water, counterstained lightly with Mayer’s haematoxylin, blued in tap water, dehydrated, cleared, and mounted. Previously identified strongly oestrogen receptor positive paraffin wax embedded tumours were used as controls, with the negative control being achieved by omission of the primary antibody.

H222 ER IMMUNOHISTOCHEMICAL METHOD (ER-IIA– H222)

Serial sections of tissue adjacent to those stained with 1D5 were taken for staining with H222. The method was the same as for the 1D5 series described above, with two exceptions. The sections were not microwaved, but instead, following incubation with H2O2, were digested with a 0.03% solution of XXV protease enzyme (Sigma) in PBS for nine minutes at 40°C and rinsed in PBS at 4°C. After incubation with a 1 in 5 dilution of normal rabbit serum for five minutes, sections were incubated overnight with rat primary monoclonal anti-human oestrogen receptor antibody H222 (Abbott Diagnostics ER-ICA Kit). The other difference in the immunohistochemical protocol was that the incubation time with the streptavidin biotinylated horseradish peroxidase complex was half an hour shorter.
SCORING ASSESSMENT

The sections were assessed without knowledge of the clinical and biochemical data. In scoring the immunohistochemically stained sections 10 fields were chosen at random at ×400 magnification (objective × eyepiece) and a minimum of 500 cells were counted. The intensity of staining in carcinoma cells was assessed semiquantitatively using a scoring index of 0 to 3, corresponding to negative, weak, intermediate, and strong positive staining intensity. The percentage of cells stained at each intensity was estimated. The product of the scoring index and the percentage gave an overall H score ranging from 0 to 300.12 Stroma, normal, and benign epithelial tissue were excluded from this assessment. A tumour was designated oestrogen receptor positive if the H score was ≥20.

Table 1 Concordance between enzyme immunoassay (ER-EIA) and immunohistochemical assay with 1D5 antibody (ER-IHA-1D5) in 119 untreated primary breast cancers

<table>
<thead>
<tr>
<th></th>
<th>EIA+</th>
<th>EIA−</th>
<th>Totals</th>
</tr>
</thead>
<tbody>
<tr>
<td>IHA+</td>
<td>64 (54%)</td>
<td>7 (6%)</td>
<td>71</td>
</tr>
<tr>
<td>IHA−</td>
<td>10 (8%)</td>
<td>38 (32%)</td>
<td>48</td>
</tr>
<tr>
<td>Totals</td>
<td>74</td>
<td>45</td>
<td>119</td>
</tr>
</tbody>
</table>

χ² = 55.6; degrees of freedom = 1; p < 0.001.

Results

The age of the patients ranged from 31 to 78 years (median 56 years). The mean value for the ER-EIA assay was 60.0 fmol/mg protein (median 25 fmol/mg protein). The immunohistochemical staining with the 1D5 antibody produced uniform nuclear staining (fig 1). The mean H score for the ER-IHA-1D5 assay was 60.5 (median 48).

There was an 86% concordance between ER-EIA and ER-IHA using the anti-1D5 antibody (table 1; χ² = 55.6; p < 0.001); 54% (64/119) of the tumours were oestrogen receptor positive and 32% (38/119) were oestrogen receptor negative by both assays. However, the quantitative results from the ER-EIA and the semiquantitative scores from the ER-IHA do not seem to be linearly related (fig 2).

A mismatch between the EIA and IHA was observed in 17 cases (fig 3). Seven of these tumours were EIA positive but IHA positive, although five out of seven had EIA values between 5 and 10 fmol/mg protein. The other 10 discordant tumours were EIA positive and IHA negative; four of these tumours were completely negative by IHA in the section studied and the EIA values in these four instances were between 10 and 18 fmol/mg protein, which may also be considered borderline. Half of the remaining six EIA positive IHA negative tumours, however, had EIA values greater than 50 fmol/mg protein (table 2).

A repeat ER-IHA-1D5 assay on the 17 discordant tumours showed that the H scores obtained initially were entirely reproducible. Identical sections of tumour were stained in a separate assay with the H222 antibody. A set of five positive and five negative concordant controls from the original series were also examined by both assays (table 2). As well as straightforward check of the data, this allowed us to address the possibility that some or all of these discrepancies were due to the antibodies targeting different ends of the oestrogen receptor.

In some tumours the IHA staining signal obtained with the 1D5 antibody was of greater intensity than that obtained with the H222 antibody. The results obtained using the two antibodies on the mismatched and control tumours are summarised in table 2. Excellent agreement was observed with both antibodies on the sets of positive and negative control tumours. Of the 10 tumours which were EIA positive but IHA negative, the H222 staining agreed with the 1D5 staining in
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Figure 3  Scattergram for tumours with EIA values of $<120 \text{fmol/mg}$ protein and IHA scores of $<120$. Positive cut off for ER-EIA is 10 fmol/mg protein, and for ER-IHA is H score 20. Discrepancies are shown as follows: ▲ = EIA positive/IHA negative cases ($n = 10$); ▲ = EIA negative/IHA positive cases ($n = 7$).

Table 2  ER-EIA results compared with ER-IHA testing with both 1D5 and H222 antibodies in 17 cases where EIA and IHA originally mismatched

<table>
<thead>
<tr>
<th>EIA Abbott fmol/mg protein</th>
<th>IHA-1D5 H score</th>
<th>IHA-H-222 H score</th>
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<tbody>
<tr>
<td>EIA negative/IHA-1D5 positive tumour mismatches:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>29</td>
<td>12</td>
</tr>
<tr>
<td>5</td>
<td>24</td>
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<td>6</td>
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<td>0</td>
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<td>0</td>
</tr>
<tr>
<td>8</td>
<td>105</td>
<td>3</td>
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<tr>
<td>EIA positive/IHA 1D5 negative tumour mismatches:</td>
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<tr>
<td>98</td>
<td>18</td>
<td>31</td>
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<td>66</td>
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<td>8</td>
<td>9 (Benign positive)</td>
<td>0</td>
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<tr>
<td>18</td>
<td>0 (Benign positive)</td>
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<tr>
<td>16</td>
<td>11</td>
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<tr>
<td>12</td>
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<tr>
<td>4</td>
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<td>0</td>
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<tr>
<td>ER positive control tumours:</td>
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<tr>
<td>268</td>
<td>133</td>
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<tr>
<td>175</td>
<td>150</td>
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<td>146</td>
<td>100</td>
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<td>306</td>
<td>112</td>
<td>110</td>
</tr>
<tr>
<td>500</td>
<td>185</td>
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<td>ER negative control tumours:</td>
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| Ten control tumours (five strongly positive and five negative) are included.

Figure 4  Correlation between H scores from ER-IHA-1D5 and ER-IHA-H222 assays in 17 tumours which were discrepant between EIA and IHA-1D5 and in five positive and five negative concordant control tumours.

Discussion

Previous studies have shown a good correlation between the ER-ICA method on frozen sections and the established ER-DCC or ER-EIA assays. Overall, the median concordance of the published studies is 86%. A reliable paraffin wax embedded ER-IHA method would permit retrospective studies on archival material from large cohorts of patients subsequently treated with endocrine treatment in whom data on response, disease free interval, or survival are now known. In many clinical studies on adjuvant endocrine treatment, less than 30% of patients have oestrogen receptor status available as frozen tissue.

The detection of oestrogen receptor, however, has been hampered by the loss of immunoreactivity during tissue processing of paraffin wax embedded material. Antigen retrieval methods, such as proteolytic enzyme digestion, together with amplification of the signal with the more sensitive avidin biotin peroxidase complex, have improved oestrogen receptor detection. Different fixation methods may also determine whether oestrogen receptor can be detected reliably. Direct comparison of ER-IHA on paraffin wax sections with ER-ICA on frozen sections has shown a concordance of between 82% and 100%, although these studies suggest that paraffin wax assays are less sensitive. In previously published series comparing ER-IHA on paraffin wax embedded material with ER-DCC, the concordance has varied between 65 and 96%. There has been only one series comparing paraffin wax assays with ER-EIA which showed a concordance of 95% in 55 tumours. In these comparisons with biochemical assays the oestrogen receptor content showed less concordance for tumours with oestrogen receptor values of $<100$.
fmol/mg protein. With the ER-ICA approach there is also considerable variation in the cutoff level quoted in published series for the semi-quantification of oestrogen receptor expression in tissue sections, ranging from more than one cell positive to >30% cells staining positive.15 Clearly, these differing cut offs may explain at least in part the variability in concordance from 68 to 93% when comparing ER-DCC or ER-EIA with ER-ICA.

In our study we used microwave treatment for antigen retrieval together with a new monoclonal antibody (1D5) directed against the N-terminal of the oestrogen receptor protein. The incidence of oestrogen receptor positivity for this series of 119 primary untreated breast cancers is in line with other published results. Using a cut off of 10 fmol/mg protein for the ER-EIA, we assessed the validity of the ER-IHA in paraffin wax embedded material with 1D5 using the H score method for semi-quantitation with a positive cut off of 20. The concordance rate of 86% is similar to that in studies comparing biochemical assays with ER-ICA by H222 on frozen sections.

However, as fig 2 shows, the quantitative correlation between the two methods is not linear. For tumours with ER-EIA values of >100 fmol/mg protein, all tissue sections had ER-IHA H scores of >80 which implies excellent concordance between the two techniques for tumours rich in oestrogen receptor. For tumours with less than 100 fmol/mg on ER-EIA, there was much greater variation in the H score (fig 3). This is probably mainly due to there not being a linear correlation between protein concentration and the intensity of the colour developed.

Using the cut offs described, there were 17 tumours in which the ER-IHA and ER-EIA mismatched. By repeating the assay in these tumours with both 1D5 and H222, four of these tumours remained completely negative by IHA in the section studied. The EIA values in these four tumours was between 10 and 18 fmol/mg protein, which is borderline, and in this situation the IHA result may have been due to sampling of an oestrogen receptor negative area. Half of the remaining 6 EIA positive IHA negative tumours had EIA values greater than 50 fmol/mg protein, and in these tumours the H222 score was between 10 and 31. Therefore, several of the apparent discrepancies between ER-IHA and ER-EIA may be explained by borderline results. A further potential explanation for these EIA positive IHA negative discrepancies could be oestrogen receptor positive benign elements present in the tumour which in a tumour homogenate would give an EIA positive result.

Three tumours remained discordant following reassessment. These tumours, which had EIA values of 0, 6, and 8 fmol/mg protein, were strongly positive with 1D5 and negative with the H222 antibody. The ER-EIA technique uses a sandwich method with the D547 antibody (against the central part of the receptor) in the solid phase and the H222 antibody (against the hormone binding domain) labelled to the peroxidase enzyme.

Thus, both the ER-EIA and ER-IHA-H222 assays require an intact hormone binding domain to detect oestrogen receptor protein. There are a substantial number of reports of oestrogen receptor molecular variants which lack one part or other of the full molecule.

Receptor with a mutated or absent ligand binding domain might be negative by these two assays, but positive by the ER-IHA-1D5 directed against the N-terminal of the receptor. Such a receptor variant has been suggested as existing following the discovery of variant forms of messenger RNA (mRNA) in human breast cancers in which the exon-5 region, coding for part of the ligand binding domain, has been spliced out during transcription.22 These variant forms of mRNA are particularly abundant in the rare tumour phenotype which is ER negative by conventional assay, but which expresses high levels of the oestrogen-dependent protein PgR. It is interesting to note, therefore, that two of the three remaining discordant tumours in our series which are only positive by ER-IHA-1D5 are of the rare ER negative PgR positive phenotype. Thus, this antibody may be of potential use in investigating variant forms of oestrogen receptor protein which are not detected by conventional assays.

In conclusion, the ER-IHA which we have described using the 1D5 antibody shows good concordance with the established biochemical ER-EIA assay. Discrepancies may occur among tumours with borderline oestrogen receptor ER-EIA values. The use of separate monoclonal antibodies directed against different parts of the receptor may also be of value in the investigation of variant forms of oestrogen receptor.

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