Demonstration of oestrogen receptors in paraffin wax sections of breast carcinoma using the monoclonal antibody 1D5 and microwave oven processing

P Sannino, S Shousha

Abstract
This study aimed at assessing the usefulness of a new monoclonal antibody (1D5) for the demonstration of oestrogen receptors (ER) in paraffin wax sections, using brief microwave processing rather than proteolytic predigestion. Routinely processed paraffin wax sections of 50 cases of breast carcinoma with known ER concentrations, estimated by the standard dextran-coated charcoal (DCC) biochemical assay, were examined using the avidin-biotin complex-immunoperoxidase technique. The results were assessed semiquantitatively, using a five grade scoring system. Of the 50 cases examined, 37 were positive and six were negative by both DCC and immunohistology. Of the remaining seven cases, three (6%) were negative by DCC but positive with immunohistology, and four (8%) were positive with DCC and negative with immunohistology. The DCC results of the latter four cases were 14, 14, 14 and 16 fmol/mg protein which is at the lowest level of positivity, our cutoff point being less than 10 fmol.

The monoclonal antibody 1D5, as used in this study, can provide easily assessed reliable information about the ER status of breast carcinoma using routinely processed paraffin wax sections.

Methods
The study was carried out on 50 cases of breast carcinoma with known ER concentrations, estimated by the standard dextran-coated charcoal (DCC) biochemical assay. All cases had been fixed in formalin and imbedded in paraffin wax. Four micrometer thick sections were cut, mounted on APES (Sigma) treated slides, and left to dry overnight at 37°C. On the following day, sections were dewaxed in xylene for five minutes and rehydrated in graded alcohol. Sections were then placed in 10 mM citrate buffer, pH 6·0; microwaved twice, five minutes each, using a microwave oven at full power (650W); left to cool for 20 minutes; and then rinsed in distilled water. Endogenous peroxidase was blocked using 3% hydrogen peroxide in distilled water for five minutes. This was followed by rinsing in running tap water, then in TRIS buffered saline (TBS). Normal goat serum, at a dilution of one in five, was applied for 20 minutes and then tipped off. This was followed by applying the primary antibody 1D5 (Dako Ltd, High Wycombe, UK) in a concentration of one in 100 for 60 minutes; rinsing in TBS, applying the secondary antibody, goat anti-mouse/rabbit, one in 100, for 30 minutes; rinsing in TBS; and applying streptavidin-biotin complex (Dako Duet) for 30 minutes. After rinsing in TBS sections were incubated with 0·05% dianisobenzidine (Sigma, Poole, UK) in 0·01% hydrogen peroxide for three minutes; rinsed in tap water; counterstained in Mayer's haematoxylin for 25 seconds; blued in tap water; then dehydrated, cleared, and mounted with ralmount (BDH, Poole, UK).

Stored frozen tissue was available from three out of the 50 cases examined. All three cases were ER positive when assayed with the DCC technique. Frozen sections, 6 μm thick, were cut from the three cases and fixed in 4% buffered formalin for 15 minutes, followed by methanol, at −20°C, for five minutes, and then acetone, also at −20°C, for three min-
utes. Sections were then rinsed in TBS and stained with 1D5 antibody as described above, and with Abbott's ER-ICA monoclonal antibody as recommended by the manufacturers. Paraffin wax sections from the same three cases were stained with Abbott's monoclonal antibody ER-ICA using the previously described immunoperoxidase technique.4

Attempts were also made, using paraffin wax sections of a known ER positive case and 1D5 antibody, to avoid using the microwave step by incubating the sections overnight with 1D5. Reduced and increased times of microwave exposure were also tried on different sections from the same case.

Negative controls were sections of the cases under examination, treated in the same way except that the specific antibody step was omitted.

Positivity was indicated by the presence of dark brown nuclear staining. The results were assessed semiquantitatively using a five grade scoring system from − to +++++ based on the percentage of stained tumour cells (occasional, less than ± (score 1); less than ½ (score 2); and almost all (score 3), and the intensity of that staining (moderate (score 1) and strong (score 2)). A case was scored as − when no staining was seen in tumour cells; + indicated the presence of an occasional positive cell; ++ (total score 1) indicated cases where up to a third of the tumour cells were moderately stained (1 × 1); +++ (total score 2–4) indicated strong staining of up to two thirds of the tumour cells (1 × 2 or 2 × 2) or moderate staining in almost all tumour cells (3 × 1); +++++ (total score 6) was used when almost all the tumour cells were strongly stained (3 × 2). All cases were scored independently by both authors and the results compared.

### Results

Of the 50 cases examined, 37 were positive and six were negative by both DCC and immunohistology (figure). The results of the two techniques were thus similar in 86% of cases. Of the remaining seven cases, three (6%) were negative by DCC but positive with immunohistology, and four (8%) were positive with DCC and negative with immunohistology. The DCC results of the latter four cases were 10, 14, 16 and 16 fmol/mg protein, the lowest level of positivity, our cutoff point being <10 fmol. If the cutoff point was thus increased to 20 fmol/mg protein, as is the case in many other centres, the concordance between DCC and immunohistology in our study would rise to 94%.

The scoring results of both authors were almost identical. A difference of opinion occurred only in two cases, which were given a score of ++ by one author and +++ by the other. After discussion a score of +++ was agreed for both cases. Good correlation existed between the immunohistology score and the mean of DCC values for cases with similar score. Thus the DCC mean showed a gradual increase from 8 fmol/mg protein for immunohistology negative cases to a mean of 365 fmol for +++ cases (table). No cytoplasmic or background staining was seen, and preservation of tissue was good.

The three ER positive cases examined with four different techniques, 1D5 frozen and paraffin wax sections and ER-ICA frozen and paraffin wax sections, gave essentially similar results. The intensity of staining, however, was stronger with the two antibodies, in frozen sections, and preservation of tissue; hence the clarity of staining was better in the microwave than in the enzyme predigested paraffin wax sections.

Attempts at avoiding the use of the microwave oven or reducing the time of exposure were unsuccessful as they resulted in no staining or substantially reduced staining results. Excessive microwave exposure had deleterious effects on the tissue sections.

### Discussion

This study shows that the monoclonal antibody 1D5 can be used successfully for the demonstration of ER in paraffin wax sections of breast carcinoma, and probably in other
tissues also. No proteolytic enzyme predigestion of sections is required. This can sometimes affect the quality of the sections. However, a brief period of processing in a microwave oven is needed. This is a critical and essential step which has to be carried out in the precise manner described. The results obtained with 1D5 were more intense, but otherwise comparable with those obtained with ER-ICA antibody when used on paraffin wax sections.

Although the staining results were less intense than those obtained with frozen sections, paraffin wax sections have the obvious advantage of using routinely processed tissue and they provide sections of higher quality which are easier to interpret.

The semiquantitative scoring system used is simple and reproducible. Its results correlate well with the quantitative estimates obtained with DCC. The scores can provide an estimate of the level of ER in a given tumour and whether it is absent, low, moderate, high or very high which may be of value to oncologists.

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1 Scottish Cancer Trials Breast Group and ICRF Breast Unit, Guy's Hospital, London. Adjuvant ovarian ablation versus CMF chemotherapy in premenopausal women with pathological stage II breast carcinoma: the Scottish trial. Lancet 1993;341:1293-8.


Prevalence of Epstein–Barr virus in the cervix

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Abstract

Cervical smears from 327 women were examined using the polymerase chain reaction (PCR) targeted to a sequence in the Bam H1 W region of the Epstein–Barr virus (EBV) to determine the prevalence of the virus in the cervix. EBV was detected in 131 (40%) of the 327 women. Of the 235 women with normal cytology, 98 (42%) were positive. Of the 92 women with dyskaryotic smears, 33 (36%) were positive.

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Epstein-Barr virus (EBV) is a human herpes virus which is associated with a variety of neoplasias such as Burkitt's lymphoma and nasopharyngeal carcinoma. The reasons for this association are poorly understood, but the virus can transform cells bearing the EBV/C3d receptor and make them receptive to other oncogenic events.

In almost all healthy EBV antibody positive people the major site from which the virus is shed is the oropharynx, and viral replication seems to be restricted to squamous epithelial cells. With the recognition that the uterine cervix is another site where the virus is shed and can replicate, the role of EBV in cervical neoplasia requires elucidation. As an initial step it was decided to establish if EBV is commonly present in the cervix and whether there is any difference in its prevalence in cytologically normal and cytologically dyskaryotic cervices.

Methods

The study population of 327 was drawn from the Grampian call/recall cervical screening programme. It consisted of 235 women (mean age 33.4, range 17–67 years) with normal cervical smears attending family planning and general practitioner clinics, and 92 women (mean age 31.8, range 18–62 years) with dyskaryotic smears attending colposcopy clinics. Ninety of the latter group had confirmation of squamous epithelial abnormalities in biopsy specimens taken subsequent to the smears.

Cervical smears collected with a spatula were collected into phosphate buffered saline and stored at −20°C until DNA extraction. All DNA samples were subjected to an amplification reaction directed to a portion of the β globin gene to validate the quality of the extracted DNA.

The polymerase chain reaction (PCR) was targeted to a sequence in the Bam H1 W region of the EBV genome. Reactions were carried out in 50 μl volumes containing 50 mM K Cl, 10 mM TRIS- HCl (pH 8.3), 1 mM Mg Cl₂, 0.01% gelatine, 200 μM each dNTP (dATP, dGTP, dCTP and dTTP), 29 July 1993

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