Characterisation of a messenger RNA selectively expressed in human breast cancer

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Summary A complementary DNA library from MCF-7 cells was screened using 32P-cDNA derived from a breast carcinoma and from normal breast tissue. From 10^4 plaques (20% of library) we obtained a clone (Md2) which was differentially expressed in the carcinoma. The distribution of its corresponding transcript of 6–700 nucleotides was examined in normal and neoplastic cells, by filter and in situ hybridisation. We observed localisation of 32S-Md2 to the tumour cells of breast carcinomas and normal ductal or vascular elements or on normal ductal epithelia. M13 sequencing showed Md2 to be 250 nucleotides in length, of which 197 were homologous to the 3’-untranslated region and a short open reading frame of the pS2 gene (Masiakowski et al., 1982). Md2 mRNA was found principally in breast carcinoma cell lines and tumours, with low levels in benign breast disease and no expression in non-breast squamous cell lines. Approximately 43% (23/54) of carcinomas contained this mRNA (varying from + to ++++++ level); it was present in 20/38 (53%) of ER positive carcinomas compared to 3/16 (19%) of ER negative carcinomas. In 21 patients who had undergone primary endocrine therapy for recurrent disease expression of Md2 in the primary tumour correlated with the subsequent response to treatment (P=0.041) and was of similar predictive value as ER status. Both tests correctly predicted outcome in about 76% of cases.

The phenotypic changes which accompany the malignant transformation of normal cells reflect in all probability an underlying change in the genotype or of its expression. This is akin to the induction/repression of regulated genes (Caplan & Ordahl, 1978) during the differentiation process, the aberrant expression of which may also lead to the cancerous state (Wald et al., 1978). Comparisons of the transcribed genome using differential hybridisation techniques (St John & Davis, 1979) have been used to identify and isolate several genes that have altered transcriptional levels associated with human leukemias (Shiosaka & Saunders, 1982) and with gastric neoplasms (Shiosaka et al., 1987). Such studies could also provide clinically useful diagnostic and prognostic markers which would be of particular value in very heterogeneous cancers such as those of the breast. Two-thirds of human mammary carcinomas are oestrogen receptor (ER) rich and thus ER status has become a valuable predictor for response to endocrine therapy (Jensen & DeSombre, 1977), and may have some prognostic value (Coombes, 1987). However, about half of patients whose tumours express ER still fail to respond (Osborne et al., 1980) to anti-oestrogens, and this has prompted a search for oestrogen responsive elements which could serve as better indicators of the tumour responsiveness, and oestrogen-stimulated proteins have been described, principally using the MCF-7 cell line (Horwitz & McGuire, 1978; Butler et al., 1979; Edwards et al., 1980; Westley & Rochefort, 1980), but none of these have proved to be as useful as ER status.

More recently, differential hybridisation of MCF-7 cDNA libraries with reverse transcribed mRNA isolated from hormone treated and untreated cells has resulted in the cloning of several oestrogen responsive genes (Masiakowski et al., 1982; Prud’homme et al., 1985; May & Westley, 1986). At least one of these, the pS2 gene, has been the subject of intensive study by the same group (Jeltsch et al., 1987; Nunez et al., 1987; Rio et al., 1987).

Our approach has been to study differential gene expression in a human cancer, MCF-7, compared to normal breast, by screening a library of the cancer. Here we describe the isolation of a differentially expressed clone, Md2 (found to be homologous to pS2), and its initial characterisation, including its distribution in breast tissues and cell lines and its possible value as a predictor of response to endocrine therapy.

Materials and methods

Patients

Fifty-four samples were obtained from 53 patients with breast cancer. Forty-nine were samples obtained at the time of primary surgery, and five were obtained at the time of relapse from biopsies of recurrent soft tissue disease. (From one patient we obtained samples of both primary and recurrence.) No therapy was given after primary surgery before the development of recurrent breast cancer. A further five samples of fibroadenoma were also studied.

Patients’ ages ranged from 29 to 85. Of the 49 primary carcinomas, all but two (one lobular and one colloid carcinoma) were infiltrating ductal carcinomas and 12 were associated with ipsilateral lymph node involvement. Forty per cent of the patients were premenopausal and 60% were post-menopausal.

Samples were obtained from between 1980 and 1987 and stored in liquid nitrogen before study. In the intervening time, 21 patients had relapsed and received primary endocrine therapy (tamoxifen 16 cases; aromatise inhibitors five cases). All had been assessed for response according to International Union against Cancer criteria (Hayward et al., 1977). The most common sites of first relapse were bone, lung, local and liver.

Materials

Tissue culture medium and fetal calf serum (FCS) were obtained from Gibco (Paisley, UK). Other reagents were obtained from Sigma Chemical Co. (Poole, Dorset, UK) unless stated.

Cell culture

Human breast carcinoma cell lines (Engel & Young, 1978) and human squamous carcinoma cell lines (Easty et al., 1981) were used in this study. Breast cell lines were MCF-7 (two sources of this line were used: Dr M. Lippman, NCI, Bethesda, MD, USA, and the laboratory of origin, the Michigan Cancer Foundation), T47D (Dr H. Freake, Hammersmith Hospital, London, UK), MDA-MB-231 (Mason Research Institute, Rockville, MD, USA), ZR-75-1.

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Differentially expressed mRNA in Human Breast

Preparation of human breast organoids

Organoids (Stampfer et al., 1980) were prepared from reduction mammooplasty tissue. Briefly, the attached skin and excess fat were removed and the residual tissue was finely diced and digested at 37°C for 18-22 h with collagenase (Type 1A, at 0.5 mg ml⁻¹) in DMEM with 7% FCS, 10 mM Hepes and antibiotics. Epithelial tissue fragments were pelleted by centrifugation at 1,000 g for 5 min, then resuspended in fresh medium without collagenase and allowed to settle for 18 h at 4°C. After a further digestion of 2 h at 37°C with 2.5 mg ml⁻¹ collagenase, the organoids were recovered by repeated gravity sedimentation for 20-60 min at 4°C. Viability was estimated by trypan blue exclusion.

Some preparations were examined for breast morphology and the presence of contaminating blood vessels by immunocytochemistry. Monoclonal antibody LIRC-LON-59.2, which recognises a cell surface component of myoepithelial cells, and plus blood vessels in the human breast (R. Skilton et al., unpublished), was used to stain frozen sections of organoid preparations by the immunoperoxidase technique (Gusterson et al., 1985). Organoids were stored under liquid nitrogen.

cDNA library

A human breast carcinoma cell line (MCF-7) random primed cDNA library in λgt11, consisting of 5 × 10⁵ clones, was kindly provided by Professor P. Chambon (Institut Chimie Biologique, Strasbourg, France).

RNA extraction

Poly(A)+RNA was extracted from 200-500 μl of packed MCF-7 cells and organoids by lysis with 7 ml proteinase K buffer (300 μg ml⁻¹ proteinase K, 2% SDS, 10 mM vanadyl ribonucleoside complex (Gibco-BRL), 0.2 M NaCl, 1.5 mM MgCl₂, 0.2 M Tris/HCl, pH 7.5) and incubation at 45°C for 2 h. During the incubation the lysates were passed several times through a 0.6 mm diameter syringe needle to shear the DNA.

Lysates were cleared by centrifugation at 10,000 g for 10 min and poly(A)+RNA was extracted by affinity chromatography on oligo (dT)-cellulose (Maniatis et al., 1982). Its quality was checked by ability to produce translation products in vitro using the rabbit reticulocyte lysate system (Amersham Int. plc) in the presence of 14C-methionine (Amersham Int. plc) according to published methods (Davis et al., 1986), and labelled products were analysed by polyacrylamide gel electrophoresis (10% PAGE/SDS gels) (Laemmli, 1970).

All other RNA used in this study was extracted from cells and biopsy material by the guanidine isothiocyanate method (Chirgwin et al., 1979). The quality of this RNA quantified spectrophotometrically was verified by the integrity of the 28S and 18S ribosomal bands following agarose gel electrophoresis.

Screening

About 10⁵ clones from an MCF-7 cDNA library were screened using standard methods. Filters (Hybond N, Amersham, UK) were prehybridised at 42°C for 4-20 h in 50% (v/v) deionised formamide (Rose Chemicals, London, UK), 0.1% (w/v) SDS, 5 × Denhardt’s solution (50 × Denhardt’s solution: 1% (w/v) each of polyvinylpyrrolidone, bovine serum albumin and Ficoll 400), 5 × SSPE (20 × SSPE: 0.2 M EDTA, 3 M NaCl and 0.2 M Na₂HPO₄, pH 8.3) and denatured sonicated salmon sperm DNA (250 μg ml⁻¹). Duplicate filters were then hybridised under the above conditions for 18 h with the addition of 4 × 10⁶ c.p.m. ml⁻¹ 32P-cDNA, prepared from MCF-7 or organoid poly(A)+RNA by oligo dT primed reverse transcription (Huyhn et al., 1985). Following hybridisation, filters were washed with five changes of 2 × SSC, 0.1% SDS at 25°C, and two changes of 0.2 × SSC, 0.1% SDS at 55-60°C.

Subcloning into pBR322

cDNA inserts from λgt11 phage DNA were excised with EcoR1, ligated into pBR322 plasmid (Biolabs) in the EcoR1 site, and transformed in E. coli JM109. Plasmid DNA was isolated by the alkaline lysis method (Birnboim & Doly, 1979), and insert was removed by EcoR1 digestion, purified by preparative agarose gel electrophoresis, and labelled with 32P or 35S-dCTP (Amersham) by the random primer method (Feinberg & Vogelstein, 1983) to specific activities of 10⁸ c.p.m. μg⁻¹ and 1.5 × 10¹⁰ c.p.m. μg⁻¹ DNA respectively.

DNA sequencing

The Md2 cDNA was excised from λgt11 phage DNA with EcoR1 and inserted into the M13 vectors mp8 and mp9. Single-stranded templates were prepared from recombinant plaques, and subjected to dideoxy chain termination sequencing (Vieira & Messing, 1982).

Dot blot hybridisation and Northern analysis

As most biopsies were small and the RNA extracted was generally low, hybridisation was normally performed using total rather than poly(A)+RNA. Whenever possible dot blots were done using serial dilutions of formaldehyde or glyoxal denatured RNA ranging from 10–20 μg to 1.25 μg, spotted on to Biodyne A nylon membrane (Pall Filtration, Portsmouth, UK) using a Bio-dot manifold (Bio-Rad, UK). Northern analysis of total RNA (20 μg per lane) or poly(A)+RNA (2.5 μg per lane) was carried out following transfer from agarose/formaldehyde gels (Seed, 1982). Transcripts were sized using denatured RNA and DNA markers. Hybridisation was carried out as described for library screening above, except 32P-labelled Md2 cDNA at 0.5–1.0 × 10⁷ c.p.m. μg⁻¹ was employed as the probe, and filters were washed to a higher stringency (0.1 × SSC, 0.1% SDS at 60–65°C).

The autoradiograms were quantified by comparison with an Md2 standard and dots were given a value ranging from 0 (undetectable), + (just detectable above background) to ++++. The highest intensity represents 100 μg of hybridisable message per 20 μg total RNA.

DNA preparation and Southern blotting

Tumour samples and cells, stored in liquid nitrogen, were thawed to 25°C in 5 volumes of 10 mM NaCl, 1 mM EDTA, 10 mM Tris pH 8, and disrupted with a polytron. Sarkosyl, 0.1 volumes of 10% (w/v) solution, and proteinase K (to 200 μg ml⁻¹) were then added and the homogenate incubated, at agitation, at 25°C for 2 h. The solution was extracted twice with an equal volume of phenol (saturated with 1 M Tris pH 8) and twice with an equal volume of chloroform-isooamyl alcohol (24:1) and the DNA precipitated with ammonium acetate and isopropanol dissolved in 5–10 ml TE (10 mM Tris, 1 mM EDTA, pH 7.5) and re-precipitated with ethanol. The DNA was then dissolved in 10 mM TE and digested in 10 μg aliquots for 24 h with 80 μl EcoR1 or 60 μl HindIII or BamHI (NBL), ethanol precipitated, electrophoresed in 0.8% agarose gels (Maniatis et al., 1982) and transferred to Biodyne A nylon membranes (Schwalm, 1975). Hybridisation of Southern filters were carried out essentially as described above, using 32P-labelled Md2.
In situ hybridisation

The procedure was a modification of that described by Barrett-Lee et al. (1987) after Lawrence & Singer (1985). Frozen sections (5-7 μm) fixed in 4% paraformaldehyde in PBS, 5 mM MgCl₂, for 15 min and stored in 70% ethanol at 4°C, were rehydrated in PBS, 5 mM MgCl₂, for 10 min at 25°C, incubated with 50 μg/ml⁻¹ pronase in PBS, 5 mM MgCl₂, for 10 min at 25°C and briefly post-fixed (4% paraformaldehyde in PBS, 5 mM MgCl₂) for 5 min. To reduce non-specific adherence of probe, sections were immersed in 0.1 M triethanolamine buffer containing 0.25% (v/v) acetic anhydride for 10 min at 25°C. Slides were then transferred to 0.1 M glycine, 0.2 M Tris pH 7.4 for 10 min and then into 50% formamide, 2× SSC at 65°C for 15 min. Sections were hybridised to denatured random primed 35S-labelled Md2 cDNA (specific activity 1.5×10⁶ c.p.m. μg⁻¹) containing in a total volume of 10 μl, 10 μg of each of E. coli rRNA and sonicated salmon sperm DNA, 50% formamide 2× SSC, 2 mg/ml⁻¹ bovine serum albumin, 20 mM dithiothreitol, 10% dextran sulphate, 0.1× Denhardt's at 37°C for 4 h. Following extensive washing sections were sequentially dehydrated in 70, 80, 95 and 100% ethanol and air dried.

For cell lines, cells were grown on gelatin coated glass slides, fixed and treated as for tissue sections. In all experiments, parallel incubations were performed using labelled pUC8 fragments as a non-specific probe. Autoradiography was carried out using a 50% aqueous solution of K5 nuclear emulsion (Ilford Ltd, UK) at 43°C, followed by exposure for 4-10 days at 4°C. Counter staining was with Haematoxylin and Eosin.

Oestrogen receptor measurement

This was carried out using either the ligand binding dextran coated charcoal (DCC) technique (McGuire & De La Garza, 1973), with modifications outlined by McClelland et al. (1986); or by an immunocytochemical assay (ERICA) using the H222 monoclonal antibody kit (Abbott Laboratories, Chicago, USA). The staining procedure has been described in detail elsewhere (McClelland et al., 1986).

Results

Organoids

The viability of organoids estimated by trypan blue exclusion was greater than 80% in all preparations. Frozen sections were taken from some preparations and stained for myoepithelial cells and blood vessels by immunoperoxidase using the monoclonal antibody LICR-LON-59.2. The organoids in these sections showed well preserved morphology with intact layers of myoepithelial and epithelial cells similar to breast in situ (Figure 1). From the sections examined it was estimated that blood vessels constituted less than 5% of the tissue in the organoid preparations.

Products from the in vitro translation of organoid poly(A)+ RNA were analysed by SDS-PAGE alongside those from MCF-7. Both showed numerous polypeptides (many in common) as discrete bands up to about 90 kD and more weakly staining bands at higher molecular sizes (data not shown). Thus the poly(A)+ RNA from both MCF-7 and organoids were of comparable quality.

Differential screening

A screen involving about 20% of the phage from the λgt11 MCF-7 cDNA library yielded six clones which showed differential hybridisation to 32P-cDNA made from poly(A)+ RNA of normal breast organoids or MCF-7 cells. Organoid sections designated Md2, gave a very strong signal with 32P-MCF-7 cDNA but none with 32P-organoid cDNA. The cloned Md2 λgt11 phage was subjected to a differential screen with cDNA derived from mRNA of MCF-7 and organoids of different sources to those used in the primary screen, with the same result. The Md2 cDNA insert was subcloned into pBR322, and found to hybridise to about 0.067% of the clones in the library, suggesting a highly represented sequence.

Size of Md2 mRNA

Northern blot analysis of MCF-7 mRNA indicated hybridisation of the Md2 cDNA corresponding to a major mRNA species of approximately 0.6-0.7 kb (Figure 2). Sometimes a faint band of about 3 kb could be seen with MCF-7 cells with very much longer exposures. We also saw this using pS2 clone. No signal was seen with a primary breast ER negative carcinoma, a fibroadenoma and MDA-MB-231 cells.

Southern analysis

Southern blotting analysis with 32P-Md2 cDNA was performed on DNA from cell lines and breast tumours digested with EcoR1 (Figure 3), BamH1 and HindIII. This yielded discrete bands of 3.1 and 9.0 kb for EcoR1, 3.5 and 7.9 kb for BamH1 and 4.9 kb and >21 kb for HindIII (data not shown). No difference was observed between Md2 mRNA positive and negative tumours. We also found bands of 5.9 kb and 2.1 kb for EcoR1 digested DNA of MCF7, which could also be seen as a faint bands in the tracks of breast cell lines ZR-75-1, MDA-MB-231 and T47D, and an organoid preparation.

Sequence analysis of Md2 clone

The Md2 cDNA was sequence using the dideoxy chain termination technique. Figure 4 shows the nucleotide sequence between the EcoR1 linker insertion sites. There is a short open reading frame with a termination codon. Comparison of this sequence with that of pS2 (Jakowlew et al., 1984) showed complete homology for nucleotides 54-249 (Figure 4) with only two differences (underlined), one of which has also been reported by Prud'homme et al. (1985). Thus most of the Md2 corresponds to the 3'-untranslated region of pS2 mRNA. Nucleotides 1-53 are unrelated to pS2, and we can only surmise that this sequence became attached to the remainder, during the linker ligation to blunt ended cDNA in the construction of the library.
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**pS2 mRNA in cell lines and non-malignant tissues**

Md2/pS2 mRNA was undetectable by dot blot hybridisation in three organoid preparations. We found only low levels (+) of pS2 message in 4/5 (80%) biopsies histologically identified as fibroadenomas. This is illustrated in Figure 5. It was not expressed in normal lymphocytes, placenta, colon, skin, squamous carcinoma cell lines of various tissue origins (tongue, larynx, bronchus) or in breast tumour cell lines

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**Figure 2** Northern blot analysis. Total RNA (20 µg) was formaldehyde denatured, electrophoresed in a formaldehyde agarose gel and blotted on to Biodyne A nylon membranes and hybridised with 32P-Md2 cDNA probe. After washing, the filter was exposed to Hyperfilm (Amersham) for three days. The source of the RNA was: (1) primary breast carcinoma; (2) MCF-7; (3) fibroadenoma; (4) MDA-MB-231. Sizes of DNA markers are shown in kb. The single intense band corresponds to an mRNA size of 600-700 bases.

**Figure 3** Southern blot analysis of genomic DNA with 32P-Md2 cDNA. About 10 µg of DNA were digested with EcoRI, electrophoresed in an 0.8% agarose gel, blotted on to Biodyne nylon filter and hybridised with 32P-Md2 cDNA. Sizes of hybridising species are indicated. The varying intensities are due to different amounts loaded on to the gel and are not significant. DNA in lanes 1, 7, breast organoids; 2, 9, 10, 12, primary breast carcinomas; 3, T47D; 4, MCF7; 5, ZR-75-1; 6, MDA-MB-231; 8, 11, lymph node metastases.

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**Figure 4** Nucleotide sequence of Md2 determined by the dideoxy chain termination technique (Vieira et al., 1982). Number excludes the EcoRI linkers. A short open reading frame is indicated. Sequence homology to pS2 (Jakowlew et al., 1984) is 70%.

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**Figure 5** Northern blot of RNA from fibroadenomas, hybridised to 32P-Md2 cDNA. Experimental details as in legend to Figure 2. Lanes 2-4 had 20 µg RNA from three different fibroadenoma samples. Lane 1 had 4 µg MCF-7 RNA for comparison.

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specific hybridisation with Md2 when compared with controls.

Hybridisation of Md2 was also visualised in a number of cases of ER positive breast carcinoma (Figure 7). We found good agreement between this method and measurement using filter hybridisation. Specific hybridisation to the tumour cell component of the sections was noted while only background levels were observed in the stromal tissue. Ducts of histologically normal appearance in tumour sections did not show any significant reactivity.

**Distribution of pS2 mRNA in breast tumours: comparison with oestrogen receptor**

We examined pS2 expression (Figure 8) in a total of 54 tumours (49 primaries and five secondaries) by dot blot hybridisation, and found varying levels of message (+ to +++++) clearly detectable above background signal seen with poly A' RNA in 23 cases (43%). In four other cases very low (±) levels of pS2 were observed. Only 1/5 secondary carcinomas was positive (+).

The ER status of the tumours was determined by ERICA in all but eight cases (which had previously been assayed by the DCC method) and compared with pS2 expression. The results are shown in Table I: 38 (70%) were ER +ve and 16 (30%) ER -ve which agrees well with previous data on this distribution. In the ER +ve group 20/38 (53%) were pS2 +ve (+ level or more), whereas in the ER -ve cancers 3/16 (19%) were pS2 +ve. All these were primary infiltrating ductal carcinomas. There was no difference in Md2 hybridisation between pre- and post-menopausal patients, with 47% and 54% positivity, respectively. Nodal involvement was also uncorrelated.

From one patient we assayed both the primary tumour and a local recurrence obtained after tamoxifen and medroxyprogesterone acetate treatment. While both had ER +ve cells, only the primary expressed pS2 (+level).

In order to determine whether there was any correlation between pS2 expression in the primary tumour compared to ER status and the response of the patient to endocrine therapy, we performed the following analysis for 21 patients: for ER, a cut off point was taken of ≥15 fmol mg⁻¹ cytosol protein (DCC method) or >50% of cells stained (using ERICA) for assessment of positivity. For Md2, the cut off point was + or below. Outcome of therapy was assessed using UICC criteria (Hayward et al., 1977). The time to first relapse ranged from 8 months to 6 years with a median value of 36 months.

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Table I Relationship of oestrogen receptor content to pS2 status in 54 breast cancer biopsies

| ER status | Md2 hybridisation level* |
|-----------|-------------------------|
| Positive  | 16 2 7 5 4 4           |
| Negative  | 11 2 1 1 1 -            |

* Determined as described under Methods.
Differentially expressed mRNA in human breast

Figure 7 Demonstration of Md2 mRNA in frozen tissue sections by in situ hybridisation. Breast carcinoma cells were hybridised with Md2 (a, c) or with pUC8 (b, d). a, Infiltrating ductal carcinoma (ER positive) showing hybridisation of Md2 to tumour cells. Weaker grain density is found over stromal areas of section (original magnification x430). b, pUC8 control section of same tumour as shown in a giving representation of non-specific probe hybridisation (original magnification x430). c, Intraductal component of a breast carcinoma (ER positive) showing strong Md2 expression by tumour cells within ductal unit. Surrounding stroma remains relatively free of hybridised probe (original magnification x220). d, pUC8 control section of same tumour as shown in c giving representation of comparative levels on non-specific probe hybridisation (original magnification x220). Hybridisation in all cases was carried out with 35S-labelled probes of comparable specific activity at 37°C for 4 h; autoradiography was for 3-4 days. Sections were counterstained in Mayer’s Haematoxylin and Eosin.

Figure 8 Dot blot analysis of total RNA from several breast cancers showing hybridisation of 32P-labelled Md2 to serially diluted (10, 5, 1 μg) aliquots of each sample. Poly(A)+RNA was applied in lane 1, to assess the extent of non-specific hybridisation. Lanes 2-9 and 12 are primary breast cancer biopsies, lane 10 is MDA-MB-231 and lane 11 is MCF-7.

The results shown in Table II demonstrate that pS2 expression could correctly predict outcome of therapy in 76% of the patients (P = 0.041; Fisher’s exact test (one-tailed)), compared with 75% for ER. There did not appear to be any greater benefit in adding both predictors.

Discussion

We adopted the method of differential hybridisation (St John & Davis, 1979) in a search for differences in gene expression between normal and malignant breast, which could result in isolation of clones that may have clinical

| Table II Relationship of pS2 expression and oestrogen receptor expression to response to endocrine therapy |
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| **Md2 or ER status** | **Complete or partial response** | **Stable disease** | **Progressive disease** | **Correct prediction** |
| Md2 0/±/+ | 3 | 2 | 11 | 16/21 (76%) |
| Md2 + ++/+++/+++ | 3 | 0 | 2 | |
| ER negative | 1 | 1 | 10 | 15/20 (75%) |
| ER positive | 4 | 1 | 1 | |

*ER was not determined in one of the six patients in this group.

*The hypothesis being that the presence of the marker will be associated with successful therapy.
value as prognostic indicators. When compared by in vitro translation, mRNA from MCF-7 and organoids gave very similar products, and the differential screens also suggest that the two mRNA populations are very similar in the abundant to mid-abundance class. From a screen of about 10^6 plaques of an ampliﬁed jg11 MCF-7 library (equivalent to 20% of the original number of recombinants) we detected only six major differences, and have reported results with one of these clones, Md2.

Other groups have used different screening of cDNA libraries to clone speciﬁcally oestrogen regulated genes (Masiakowski et al., 1982; Prud’homme et al., 1985; May & Westley, 1986) and progesterin regulated genes in breast cancer (Chalbos et al., 1986). One oestrogen regulated sequence ﬁrst isolated by Masiakowski et al. (1982), subsequently by Prud’homme et al. (1985) and probably also by May & Westley (1986) is the pS2 gene. Sequence analysis of the Md2 clone showed that it essentially encoded most of the 3′-untranslated region homologous to pS2.

The pS2 gene is reported to be expressed predominantly in ER positive breast cancers (97% of pS2 mRNA positive cancers being ER positive (Rio et al., 1987)). We found it to be signiﬁcantly expressed in 3/16 (19%) ER negative tumours. Also pS2 mRNA has been reported as undetectable in benign breast tumours (Rio et al., 1987) whereas we found Md2 hybridisation, albeit at low levels in 4/5 fibroadenomas. These differences are probably due to the higher speciﬁc activity (10^6 c.p.m. µg^{-1}) of our probes (compared to 10^5 c.p.m. µg^{-1} (Rio et al., 1987)). Md2 hybridisation was also found in an ovarian carcinoma (data not shown). It is possible that the patient whose primary tumour but not the recurrence expressed Md2 (although ER status had remained unchanged) did so as a result of the tamoxifen therapy. Recently Rio et al. (1988) reported pS2 expression in normal human stomach, which we have also found (Bennett et al., 1989). They were unable to detect pS2 in any other normal tissue except salivary gland.

Southern blotting analysis of the Md2/pS2 gene using DNA from Md2 mRNA positive and negative breast tumours, digested with EcoR1 (Figure 3), BamH1 and HindIII gave similar results for Md2 positive and negative tumours. This indicates that the lack of Md2 expression is not due to an absence of the gene; similarly high pS2 expression is not due to gene ampliﬁcation. The slight differences in the intensities shown in Figure 3 were due to differences in loading as judged by the ethidium bromide staining before transfer.

The banding pattern was very similar to that reported for

the pS2 gene (Jeltsch et al., 1987). Results with BamH1 and HindIII digested DNA were similar to those seen using pS2 but the Md2 also revealed additional fragments, namely a 4.9 kb HindIII fragment and 7.9 kb BamH1 fragment. There were also two extra EcoR1 bands in the MCF-7 track (Figure 3), which could be seen as very faint bands in the tracks of the other breast cell lines and an organoid preparation but not in any tumour. The signiﬁcance of these additional bands, which were not reported for pS2, remains to be seen. They could be ascribed to hybridisation by nucleotides 1–53 of Md2. However, this portion does not contribute to the RNA hybridisation as only the pS2 mRNA band was observed.

As reported previously (Barrett-Lee et al., 1987) we found that results obtained using in situ hybridisation correlate well with the ﬁlter hybridisation method. We were able to observe good localisation of Md2 to cancer cells. It was not clear whether myoepithelial cells, which can still be seen in intraductal carcinomas, expressed Md2 to the same extent as the epithelial cancer cells, but no signiﬁcant reaction was seen in the stromal tissue or vascular elements. This technique will prove very useful in our intended studies of Md2 and other clones in needle aspirates, in which we hope to measure expression throughout the course of endocrine therapy.

The value of ER status in predicting response to endocrine therapy (Coombes, 1987) has been evident for some time. Here, we have shown that Md2 may be an equally reliable predictor of response. Like ER, its expression was signiﬁcantly related to outcome of therapy. It should also be noted that of the three responders who were put into the Md2 'negative' category (refer to Table II) only one had a zero level for Md2. Therefore it may be that a re-assessment of our cut-off point for Md2 will improve its predictive value. In view of its relative abundance (and hence ease of detection) compared to ER this mRNA could prove to be a valuable marker. As with most analyses of this kind it is important to keep sample numbers in perspective. We hope to obtain more clinical data from patients presently undergoing primary endocrine therapy to provide a larger group for statistical analysis, before coming to any firm conclusions on the usefulness of pS2 as a clinical marker.

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