A differential PCR assay for the detection of c-erbB 2 amplification used in a prospective study of breast cancer

B A Jennings, J E Hadfield, S D Worsley, A Girling, G Willis

Abstract

Aims—To establish a robust differential polymerase chain reaction (PCR) assay for the detection of c-erbB 2 amplification in breast cancer that can be used in a routine pathology laboratory. Once established, the assay was used in a prospective study of breast tumours to investigate the relation between c-erbB 2 amplification and both recognised prognostic features and short term clinical outcome.

Methods—The differential PCR was used for the co-amplification of c-erbB 2 and a reference gene from 48 tumour DNA samples and control DNA samples. The ratio of the two genes was determined by image analysis of the PCR products electrophoresed on a highly resolving agarose gel.

Results—The differential PCR assay was shown to be accurate and reproducible using the conditions outlined. Twenty six per cent of the breast cancer patients were shown to have c-erbB 2 amplification in their tumour biopsies. Twenty eight per cent of the patients died of their disease or had disease recurrence during the follow up period and 73% of these patients had amplification of c-erbB 2.

Conclusions—A significant association was found between c-erbB 2 amplification and early disease recurrence. This assay could be used to provide a marker for poor prognosis in breast cancer.


Keywords: differential PCR; c-erbB 2; breast cancer

The c-erbB 2 oncogene, also known as HER2 and neu, is located on chromosome 17 (q21–22) and encodes a 185 kDa transmembrane protein that is a member of the erbB family of receptor tyrosine kinases.1 2 Frequently, c-erbB 2 is amplified and overexpressed in breast cancer and both of these abnormalities have been found to correlate with both disease recurrence and reduced overall survival.3 4

This paper describes a simple, robust, and highly sensitive differential polymerase chain reaction (PCR) method for detecting amplification of c-erbB 2 in a routine pathology laboratory. Differential PCR is a semiquantitative assay for the co-amplification of a target gene and a reference gene in the same reaction tube.5 The level of amplification of the target gene is seen by the ratio between the intensity of the two PCR product bands visualised on a gel.

Other studies have examined a variety of differential PCR methods6–11 for the detection of c-erbB 2 amplification, which has been shown to correlate with p185-erbB immunostaining,12 13 but only a few studies have examined the prognostic use of the assay with clinical follow up.7 9 11

In this study, reliable measurement of c-erbB 2 amplification was achieved when the co-amplification of the two gene sequences (c-erbB 2 and β globin) was optimised. These gene targets in this study are on different chromosomes and so the results will reflect an increase in the c-erbB 2 copy number irrespective of whether a small region of the chromosome or the whole of chromosome 17 is duplicated. Chromosome aneuploidy, including loss and gain of chromosome 17, is seen frequently in breast cancer.

The two primer pairs were selected to be non-complementary at their 3' termini and for their similar GC content. In addition, the PCR amplification was stopped before the end of the exponential phase of the reaction (the plateau) was reached.

We have analysed DNA extracted from 42 breast tumour samples, most of which have been described previously.10 12 13 Short term clinical follow up of the breast cancer patients revealed that c-erbB 2 amplification was associated strongly with early relapse. These data demonstrate that this assay identifies a subset of breast cancer patients with poor short term prognosis.

Methods

Samples from 42 female patients treated for primary breast cancer by the same surgical team between 1993 and 1994 were included in this study. The mean age of the patients was 61.0 years, ranging from 35 to 85 years. No woman had received preoperative radiotherapy. Fresh tumour samples were obtained from both mastectomy and excision biopsy specimens and DNA was extracted as described previously.2

In addition to routine pathology examination, samples used for DNA extraction were examined histologically and were shown to consist of at least 70% tumour cells. The tumours ranged in size from 0.7 cm to 10 cm (mean 3.2 cm). The selection criterion for inclusion in the study was that an adequate amount of tumour was available for the extraction of DNA.

DNA was also extracted from the peripheral blood of 10 of the breast cancer patients and 28...
normal individuals. In addition, DNA was extracted from two cell lines with known alterations of c-erbB 2 copy numbers: MCF7 may be hemizygous for c-erbB2 and SKBR3 has up to eightfold c-erbB 2 amplification.7 These samples served as controls for the optimisation of the assay.

All DNA was diluted to the same concentration (50 ng/μl). A dilution series of the SKBR3 DNA in normal DNA was prepared to test the linearity of the measurement achieved by the differential PCR.

The primers used were as follows. For c-erbB 2: 5'-TGGAACGTGCTGTCA AGA-3' (sense primer) and 5'-ATGGAAT GCTCTACT CTGTCTCGTCAA-3' (antisense primer); these primers amplify a 91 base pair fragment from exon 3.

For β globin: 5'-ACACAACGTGCTTCCA CTAGC-3' (sense primer) and 5'-CAACATT CATCCACGTTCCACC-3' (antisense primer); these primers amplify a 110 base pair fragment from exon 1.

DNA amplification was carried out in duplicate for each sample using a Progene thermal cycler (Techne, Cambridge, UK). Each 50 μl reaction mixture contained 25 μl PCR master mix (Boehringer Mannheim, Lewes, East Sussex, UK), 5 μl of each primer (50 pmol), 2 μl of DNA (100 ng), and 13 μl of sterile distilled water. Two controls that contained all the reagents but no target DNA were included with each batch. The reaction mixtures were prepared and kept on ice until the heating block of the thermal cycler reached the denaturation temperature (94°C). Each reaction mixture was placed at 94°C for five minutes and then subjected to 35 amplification cycles; each cycle was 30 seconds at 94°C, 30 seconds at 50°C, and 30 seconds at 72°C. This was followed by a final extension at 72°C for seven minutes. Initially, the optimum number of PCR cycles was determined empirically by analysing the amplification products after 20 to 50 cycles, at five cycle increments.

Amplification products were separated by electrophoresis using a 3% metaphor agarose gel (Flowgen, Lichfield, Staffordshire, UK), stained with SYBR green DNA gel stain (Flowgen), and visualised by ultraviolet illumination. The sizes of the PCR products were compared with a molecular weight marker, pUC18 DNA digested with HaeIII (Sigma, Poole, Dorset, UK). The gel images were captured using a CCD camera linked to an image processing system (GDS 8000; UVP, Cambridge, UK). The intensity of the c-erbB 2 band and the β globin band was determined for each specimen, by means of Gelworks software (UVP). These results were expressed as the ratio: intensity of the c-erbB 2 band/intensity of the β globin band. The ratios determined for the tumour samples were converted into a measure of gene amplification using the ratios determined for the normal controls and cell line controls. The cut off point for amplification was the mean of the normal range plus two standard deviations (SD).

Univariate statistical analysis comparing clinical and laboratory findings was carried out using Fisher's exact test.

**Results**

The results relating to typical control DNA and breast tumour DNA samples are shown in fig 1. The 91 and 110 base pair PCR products were resolved clearly using 3% metaphor agarose. For all normal DNA controls the intensity of the higher molecular weight β globin band was greater than the intensity of the lower molecular weight c-erbB 2 band.

The study of the optimum numbers of PCR cycles showed that consistent differences in the c-erbB 2 and β globin PCR products could be observed from 25 cycles (the sensitivity limit of the assay) to 40 cycles (the PCR plateau).

A dilution series consisting of five twofold dilutions of SKBR3 DNA into normal DNA was used for differential PCR. The ratios of c-erbB 2 to β globin showed a linear relation with the number of copies of c-erbB 2 present (examples are shown in fig 1). This demonstrated the quantitative accuracy of this differential PCR method. Each sample was analysed at least twice and each replicate gave concordant results.

Forty two DNA samples from primary tumours and six DNA samples from nodal metastases were analysed for c-erbB 2 amplification. The distribution of c-erbB 2/β globin ratios was bimodal. The majority of samples had ratios similar to the normal controls (within the normal range of mean ±2 SD) and the remainder had from three to greater than eightfold c-erbB 2 amplification. Eleven primary tumours and lymph node metastases derived from two of these tumours had c-erbB 2 amplification. Thirty one primary tumours and lymph node metastases derived from four of these tumours had a normal c-erbB 2 copy number. Therefore, 11 of 42 (26%) of these breast cancer patients had c-erbB 2 amplification in their tumours. These results are shown in relation to the tumour types in table 1.

Clinical follow up information was available for 41 of 42 patients. The median duration of follow up was 28 months with a minimum of six months for a patient who died and a maximum of 42 months. Seven patients died of breast cancer and a further four had recurrent disease. One patient died of other causes and so was excluded from statistical analysis.
Eleven of 40 (28%) patients died of their disease or had disease recurrence and 8 of 11 (73%) of these patients had c-erbB 2 amplification. The latter was associated strongly with early disease recurrence (p = 0.0003).

Twenty three patients had histological evidence of lymph node metastases at presentation and eight of these patients (35%) died of their disease or had disease recurrence. Lymph node metastasis had an association with early disease recurrence but this did not reach statistical significance (p = 0.1). Six of seven (86%) patients with c-erbB 2 amplification and lymph node metastases died of their disease or had disease recurrence.

Discussion

We present a robust differential PCR assay for the detection of c-erbB 2 amplification in human DNA. Because a numerical result is generated by an image analysis system, this technique provides objective analysis of a molecular marker, making it a good candidate for development as a routine pathlogy test. The PCR primers and cycle numbers have been optimised to give reproducible results that are not subject to primer dimer or plateau artefacts. The sensitivity of this assay, with the use of BYDR disease recurrence. Lymph node metastasis had an association with early disease recurrence but this did not reach statistical significance (p = 0.1). Six of seven (86%) patients with c-erbB 2 amplification and lymph node metastases died of their disease or had disease recurrence.

Eleven of 40 (28%) patients died of their disease or had disease recurrence and 8 of 11 (73%) of these patients had c-erbB 2 amplification. The latter was associated strongly with early disease recurrence (p = 0.0003). This finding concurs with those of others, but not all, other studies that have used differential PCR methods in retrospective analyses with longer periods of clinical follow up than is presented here. In this study, the DNA analysed was from symptomatic patients, many of whom presented with relatively large and intermediate or high grade tumours. It would also be interesting to investigate the prognostic use of this assay in the often smaller and better differentiated tumours detected in the national breast screening programme.

Breast cancer is a heterogeneous disease. There may be many different mechanisms by which tumours grow, metastasise, and evade treatment response. Genetic markers that subclassify these tumours could help to identify those patients who would benefit most from adjuvant therapy.

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