Detection of circulating epithelial cells in the blood of patients with breast cancer: comparison of three techniques

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This study compares the sensitivities and specificities of three techniques for the detection of circulating epithelial cells in the blood of patients with breast cancer. The number of circulating epithelial cells present in the blood of 40 patients with metastatic breast cancer and 20 healthy volunteers was determined by: immunomagnetic separation (IMS) and laser scanning cytometry (LSC), cell filtration and LSC and a multimarker real-time RT–PCR assay. Numbers of cytokeratin-positive cells identified and expression of three PCR markers were significantly higher in the blood of patients with breast cancer than in healthy volunteers. Using the upper 95% confidence interval of cells detected in controls to determine positive patient samples: 30% of patients with metastatic breast cancer were positive following cell filtration, 48% following IMS, and 60, 45 and 35% using real-time RT–PCR for cytokeratin 19, mammaglobin and prolactin-inducible peptide. Samples were significantly more likely to be positive for at least one PCR marker than by cell filtration (83 vs 30%, \(P<0.001\)) or IMS (83 vs 48%, \(P<0.001\)). The use of a multimarker real-time RT–PCR assay was therefore found to be the most sensitive technique for the detection of circulating epithelial cells in the blood of patients with breast cancer.

Keywords: breast cancer; micrometastases; circulating tumour cells

Over recent years, it has become apparent that circulating epithelial cells, assumed to represent metastatic tumour cells, can be detected in the blood of many patients with breast cancer (Ring et al, 2004). The detection of such cells could have significant clinical utility in risk stratification in early breast cancer, in early detection of relapse and in monitoring response to treatment. Cytometric techniques based on immunohistochemical analyses and nucleic acid-based approaches to cell detection have been described (Racila et al, 1998; Lambrechts et al, 1999; Smith et al, 2000; Aerts et al, 2001; Stathopoulou et al, 2002; Witzig et al, 2002). However, there is considerable variability in the reported sensitivities and specificities of existing techniques with putative carcinoma cells reported to be present in between 0 and 100% of blood samples from patients with metastatic breast cancer (Ring et al, 2004). Unfortunately, the design of published studies makes it difficult to compare directly the performance of current techniques and therefore to ascertain the optimal approach.

In the current study, we directly compare three techniques for the detection of circulating epithelial cells in the blood of patients with breast cancer and a healthy volunteer control population. The first technique combines positive immunomagnetic separation (IMS) with laser scanning cytometry (LSC), techniques that have both previously been applied to circulating epithelial cell detection (Brandt et al, 1998; Pachmann et al, 2001; Witzig et al, 2002). The second technique uses a novel approach to enrichment based on cell filtration as described by Vona et al (2000), and previously used in our own studies in combination with LSC (Zabaglo et al, 2003). The final method of analysis is a multiple marker real-time RT–PCR assay. The PCR assay was developed to detect cytokeratin 19 (CK19), mammaglobin and prolactin-inducible peptide (PIP) (Clark et al, 1999), which are expressed in breast epithelial cells but at low levels in normal blood components. CK19 was chosen as it is expressed in the majority of breast cancers (Bartek et al, 1986; Moll, 1994), and has been extensively used in studies in this area (Slade et al, 1999; Smith et al, 2000; Aerts et al, 2001; Stathopoulou et al, 2002). Mammaglobin gene expression has previously been used to detect circulating breast cancer micrometastases with no false positives in control populations (Zach et al, 1999; Silva et al, 2002). In a recent study, mammaglobins, and the third marker PIP, were found to be the most sensitive and specific markers for the detection of micrometastatic breast cancer in axillary lymph nodes (Mitas et al, 2003).

MATERIALS AND METHODS

Clinical samples

Ethical approval was granted for this study by the Royal Marsden Hospital Local Research and Ethics Committee, and written informed consent was obtained. Patients were eligible if they had metastatic breast cancer (stage IV according to AJCC criteria (Singletary et al, 2002)) and were not receiving cytotoxic chemotherapy at the time of enrolment (endocrine and bisphosphonate therapy were permitted). Healthy volunteers were selected from the staff of the Academic Department of Biochemistry and the Royal Marsden Hospital Breast Unit. One 18 ml blood
sample was taken from each subject via a peripheral vein. Venepuncture was with a 21- or 23-G needle, and the first 5 ml were discarded to minimise contamination by skin epithelial cells. Samples were taken into EDTA tubes (Greiner Bio-One Ltd, Stonehouse, UK) and processed within 30 min of venesection. All analyses were conducted blind of the patient/control status of the samples.

Each sample was split into 6 ml aliquots for analysis by one of the following three techniques:

**IMS followed by LSC:** The 6 ml blood sample was subjected to density gradient enrichment over an equal volume of Histopaque-1077 (Sigma-Aldrich Company Ltd, Poole, UK). The protocol used and PBS washes were according to the manufacturer’s instructions. The resulting cell pellet was resuspended in 100 µl of PBS and incubated with mouse BerEP4-FITC (20 µg ml⁻¹) antibody (Dako, Cambridge, UK) for 10–15 min at room temperature. Two washes were performed in IMS buffer (Ca²⁺- and Mg²⁺-free PBS, 0.5% bovine serum albumin and 2 mm EDTA) prior to incubation with goat anti-mouse IgG Microbeads (Miltenyi-Biotec Ltd, Surrey, UK) for 15 min at 4°C. After further washes in buffer, the cells were subjected to magnetic separation using a MiniMACS separation column (Miltenyi-Biotec Ltd, Surrey, UK). The positive fraction was centrifuged and resuspended in PBS and a cytospin prepared using a Hettich Universal 32R centrifuge (Hettich-Zentrifugen, Germany). The cytospin spot deposited on a microscope slide was fixed in 100% methanol for 10 min and air-dried.

For cytokeratin staining slides were rehydrated in PBS followed by TBP (0.5% Triton X-100 and 0.5% bovine serum albumin in PBS) and blocked with normal rabbit serum. Slides were then counterstained with 20 µg ml⁻¹ of the nuclear DNA fluorochrome PI (Promega, Southampton, UK) and 100 µg ml⁻¹ RNase A (Sigma-Aldrich Company Ltd, Poole, UK) for 30 min. Slides were mounted in Vectashield containing 5 µg ml⁻¹ PI (Vector Laboratories Ltd, Peterborough, UK). The MCF7 breast cancer cell line was used as a positive control, and an isotype mouse IgG1 antibody as a negative control.

Slides were analysed using a Laser Scanning Cytometer (CompuCyte Inc., Cambridge, MA, USA) with WinCyte PC-based software as we have described previously (Zabaglo et al, 2003). Briefly, the slides were scanned using the argon laser 488-nm line, and the cells separated on the basis of the DNA fluorescence area, maximal pixel intensity, integrated fluorescence and perimeter into cell clumps, doublets and single cells. Green fluorescence (FITC-labelled cytokeratin) against integrated red fluorescence was quantified for each of the three cell populations and cytokeratin-positive cells identified. After analysis, all the cytokeratin-positive cells were relocated and examined by eye; only cells fulfilling the criteria of the European ISHAGE Working Group (European ISHAGE Working Group, 1999) were accepted as positive.

**Cell filtration followed by LSC:** Blood samples, diluted 1:1 in PBS 2 mM EDTA, were filtered under gravity through a Poretics polycarbonate Track-Etch-type (PCTE) membrane (Genetic Research Instrumentation Ltd, Braintree, UK) with calibrated 8 µm cylindrical pores, as described previously (Zabaglo et al, 2003). The filters were then washed with PBS and fixed in 100% methanol prior to being removed from their holders and attached to a microscope slide. Slides were stained for cytokeratin and LSC analysis performed as for samples prepared by IMS.

**Real-time RT–PCR:** Samples were subjected to density gradient enrichment using Histopaque-1077 (Sigma-Aldrich Company Ltd, Poole, UK) as described for IMS. The cell pellets were stored in liquid nitrogen, and RNA extraction performed using TRIzol Reagent (Invitrogen Ltd, Paisley, UK) and RNA quantification using an Agilent 2100 Bioanalyzer (Agilent Technologies Ltd, West Lothian, UK). DNase treatment was carried out using an RNase-free DNase kit (Promega, Southampton, UK), according to the manufacturer’s instructions. The RNA pellets were dissolved in nuclease-free water and stored at −70°C prior to use.

Primers and probes were designed for CK19, mammaglobin, PIP and the ribosomal RNA RPL19 using Primer Express™ software (Applied Biosystems, Warrington, UK). Where possible, primers and probes were designed to span intron–exon junctions, and in the case of CK19 to capitalise on sequences of nonhomology between CK19 cDNA and its known pseudogenes (Ruud et al, 1999). BLAST sequence similarity searches (www.ncbi.nlm.nih.gov/BLAST) were performed to confirm that there was no significant homology between the sequences chosen and other genes in the database. The final sequences and labels used are shown in Table 1. The oligonucleotides were synthesised by Applied Biosystems (Warrington, UK).

Reverse transcription and real-time RT–PCR was carried out using the ABI Prism 7000HT operating system (Applied Biosystems, Warrington, UK). For each test gene, 10 µl of DNase-treated RNA (20 ng µl⁻¹) was added to 50 µl of 2 × AmpliTaq Gold® DNA polymerase master mix. 2.5 µl 40 × Multiscribe™ reverse transcriptase (both Applied Biosystems, Warrington, UK), forward and reverse primers and probe at the optimised concentrations, and nuclease-free water to a total volume of 100 µl. A measure of 25 µl of this mix was added to three wells such that each sample was run in triplicate with 50 ng of RNA template. All sample plates were run with positive controls (T47D breast cancer cell line RNA) and no-template negative controls. The following thermal cycling conditions were used: 48°C for 30 min; 95°C for 10 min; 40 cycles of 95°C for 15 s followed by 60°C for 60 s.

Mean threshold (Ct) values, the number of cycles after which the emitted fluorescence crossed the threshold fluorescence, were calculated for each sample for each of the test genes and the RPL19 normaliser. These were converted to mean normalised gene expression values using the ΔCt method.

Table 1 Primer and probe sequences for the target genes

| Target gene | Gene bank accession number | Forward primer sequence | Reverse primer sequence | Probe sequence | 5′ label | 3′ quencher |
|-------------|---------------------------|-------------------------|-------------------------|----------------|---------|------------|
| CK19        | NM0002276                 | TGGCGGACAGAAGATCTTAGGT | TCTCAAACCTTGGTCGGAAGTCA | ACCATGGAAGCTCCGAGT | 6-FAM TAMRA |
| Mammaglobin | AF015224                  | TGGCATATAGATGATCTGG     | TCATATATTATCTGCAAAAAC  | ACCAAGCAGGATGAAAT | 6-FAM MGB  |
| PIP         | J03460                    | TGGAGGCCTCCGTCTGTTG     | AGGAGAATTCGCCACAGTTTC  | CCCCAGTGTACCCTGTCGTC  | 6-FAM MGB  |
| RPL19       | NM000981                  | CCATGAGTACCTGAGGCCTGA  | CTCAGCGGAGTGGCAATG     | VIC™ TAMRA       |

CK19 = cytokeratin 19; PIP = prolactin-inducible peptide; 6-FAM, 6-carboxyfluorescein; TAMRA, 6-carboxy-tetramethylrhodamine; MGB, minor groove binder; nonfluorescent quencher.

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expression (MNE) using the following equation (Muller et al., 2002):

\[
MNE = \frac{(E_{\text{ref}} - C_t \text{ref mean})}{(E_{\text{target}} - C_t \text{target mean})}
\]

where MNE is the mean normalised expression, \( E_{\text{ref}} \) is the PCR amplification efficiency of the reference gene, \( E_{\text{target}} \) is the PCR amplification efficiency of the target gene, \( C_t \) ref is the threshold cycle of the PCR amplification of the reference gene and \( C_t \) target is the threshold cycle of PCR amplification of the target gene. When the PCR amplification curve had not passed the threshold fluorescence after 40 cycles, the marker was described as undetectable in that sample and the MNE designated zero.

Statistics

Differences between the number of circulating cell numbers between the control and patient populations were assessed using the Mann–Whitney U-test for unpaired non-normally distributed groups. Differences in cell numbers between the two cytometric techniques were analysed with the Wilcoxon signed-rank test for paired non-normally distributed groups. Correlations between the three techniques were assessed by calculating Spearman’s rank correlation coefficient. Differences in positivity rates between the three techniques were assessed using the McNemar test. The \( \chi^2 \) test was used to assess the relation between patient characteristics and rates of positive samples. Values of \( P < 0.05 \) were considered statistically significant. StatView 4.5 software (Cherwell Scientific, UK) was used throughout.

RESULTS

Baseline characteristics

Blood samples were taken from 40 patients with metastatic breast cancer and from 20 healthy volunteers. All analyses were performed blinded to the patient/control status of samples. The patients’ baseline characteristics are shown in Table 2.

Detection by LSC following IMS

In the 20 control subjects, the mean number of positive epithelial cells detected was 0.40 ml\(^{-1}\) using LSC following IMS. In the 40 patients with metastatic breast cancer, the mean number of positive cells detected was 6.55 ml\(^{-1}\) (\( P < 0.001 \), Mann–Whitney U-test). Positive cells were identified in the blood of four out of 20 (20%) healthy volunteers compared with 34 out of 40 (85%) patients with metastatic breast cancer (Figure 1). In common with a previous study, the upper 95% confidence interval of mean cell numbers in controls was used as a cutoff to define positive samples in patients with metastatic breast cancer (Aerts et al., 2001). Using such a threshold, 19 out of 40 (48%) samples from patients with metastatic breast cancer were positive. An alternative way to analyse this data is to set a threshold value for positive samples in the patient population, which corresponds to 100% specificity in the control population. Using this approach 15 out of 40 (38%) samples from patients with metastatic breast cancer were positive.

Detection by LSC following cell filtration

In the 20 control subjects, the mean number of positive epithelial cells detected was 0.42 ml\(^{-1}\) blood sample using LSC following cell filtration. In the 40 patients with metastatic breast cancer, the mean number of positive cells detected was 3.87 ml\(^{-1}\) (\( P = 0.0012 \), Mann–Whitney U-test). Positive cells were identified in the blood of five out of 20 (25%) healthy volunteers compared with 28 out of 40 (70%) patients with metastatic breast cancer (Figure 2). Using the upper 95% confidence interval of mean cell numbers in controls as a cutoff, 12 out of 40 (30%) samples from patients with metastatic breast cancer were positive. Alternatively ensuring 100% specificity in the control population, 10 out of 40 (25%) samples from patients with metastatic breast cancer were positive.
Detection by real-time RT–PCR for CK19, mammaglobin and PIP

The MNE values for the patients and healthy volunteer controls for the three PCR targets are shown in Table 3. Differences in MNE between the healthy volunteers and patients were statistically significant for all three markers: CK19 ($P<0.0001$), mammaglobin ($P=0.0011$) and PIP ($P=0.002$). There were no statistically significant differences between the $C_t$ values for RPL19 for the patients and controls (22.53 vs 22.47, $P=0.58$). The MNE for individual subjects for each of the three markers are shown in Figures 3–5.

The upper 95% confidence interval of MNE in control subjects was used as a cutoff to define positive samples in patients with metastatic breast cancer. Using this threshold, 24 out of 40 (60%) samples from patients with metastatic breast cancer were positive for CK19, 18 out of 40 (45%) were positive for mammaglobin and 14 out of 40 (35%) were positive for PIP. Using a threshold to ensure 100% specificity in the control population 22 out of 40 (55%) samples from patients with metastatic breast cancer were positive for CK19 compared with 18 out of 40 (45%) for mammaglobin and seven out of 40 (18%) for PIP.

Regarding all 40 blood samples from the patients and using the 95% confidence interval threshold: seven (18%) were negative for all of the genes, 16 (39%) positive for only one gene, 11 (28%) positive for two genes and six (15%) positive for all three genes. In the 20 healthy volunteers: two (10%) were positive for one gene, but none of them was positive for more than one gene (Table 4).

There were no significant differences between the number of samples positive for CK19 and mammaglobin, and PIP and mammaglobin. However, significantly more samples from patients with metastatic breast cancer were positive for CK19 than for PIP (60 vs 35%, $P=0.014$). Concordant samples were defined as those where the patient sample was reported as either positive or negative by both of the markers being used. As such, the concordance between CK19 and mammaglobin was 65%, between CK19 and PIP was 53%, and between PIP and mammaglobin was 50%.

Comparisons between the three techniques

The differences in mean cell numbers detected by LSC following IMS and cell filtration in patients with metastatic breast cancer did not reach statistical significance (Wilcoxon’s signed rank test, $P=0.08$). There was also no significant difference between the number of samples defined as positive by LSC following IMS and

Table 3  MNE of CK19, mammaglobin and PIP assessed in the blood of 40 patients with metastatic breast cancer and 20 healthy volunteers by real-time RT–PCR

| Test gene | MNE healthy volunteers ($n=20$) (range) | Healthy volunteers in whom test gene detectable ($n=20$) | MNE patients with metastatic breast cancer ($n=40$) (range) | Patients in whom test gene detectable ($n=40$) |
|-----------|----------------------------------------|-------------------------------------------------|-------------------------------------------------|----------------------------------|
| CK19      | 0.00000996 (0.0000049)                  | 7 (35%)                                          | 0.001780 (0–0.032)                               | 37 (93%)                         |
| Mammaglobin| 0.0000013 (0.0000025)                   | 1 (5%)                                           | 0.0000578 (0–0.01870)                            | 18 (45%)                         |
| PIP       | 0.0000043 (0.00000241)                  | 13 (65%)                                         | 0.0000128 (0.0000014–0.00000543)                 | 40 (100%)                        |

MNE = mean normalised expression; CK19 = cytokeratin 19; PIP = prolactin-inducible peptide; RT–PCR = reverse transcriptase–polymerase chain reaction.
cell filtration (48 vs 30%, \( P = 0.18 \), McNemar test). There were significant correlations between the numbers of cells detected by LSC following cell filtration and IMS both in patients with metastatic breast cancer and in controls (Spearman’s rank correlation coefficient, \( \rho = 0.432, P = 0.007 \) and \( \rho = 0.742, P = 0.0012 \), respectively).

Using the 95% confidence interval threshold in controls to define positive samples, patients with metastatic breast cancer were significantly more likely to be positive using the real-time RT–PCR assay for CK19 than using cell filtration followed by LSC (60 vs 30%, \( P = 0.006 \), McNemar test), but this was not the case comparing the CK19 RT–PCR assay with IMS followed by LSC (60 vs 48%, \( P = 0.06 \)). However, when all three PCR markers were used, samples from patients with metastatic breast cancer were significantly more likely to be positive for at least one of the markers than they were likely to be positive by cell filtration or IMS followed by LSC (83 vs 30%, \( P < 0.001 \) and 83 vs 48%, \( P < 0.001 \), respectively).

Using the threshold set at 100% specificity, samples from patients with metastatic breast cancer were also significantly more likely to be positive for at least one of the markers than to be positive by cell filtration or IMS followed by LSC (73 vs 25%, \( P < 0.001 \) and 73 vs 38%, \( P < 0.01 \), respectively).

Concordant samples were defined as those where the sample from a patient was reported as either positive or negative by both of the techniques being compared. As a result, the concordance between samples analysed by cell filtration and IMS was 65%. Of the 12 patients positive by cell filtration, nine were also positive by IMS. When IMS and real-time RT–PCR for CK19, mammaglobin and PIP were compared, the concordances were 65, 70 and 50%, respectively. The corresponding figures for cell filtration compared with the real-time RT–PCR assays were: 60, 60 and 50%.

The concordance between samples defined as positive as a result of expressing all three test genes and those defined as positive by cell filtration and IMS was 73 and 60%, respectively.

DISCUSSION

The detection of circulating tumour cells has considerable potential to influence the management of patients with breast cancer. However, using existing techniques there is considerable variability in the rates of positive samples in patients with breast cancer patients and healthy volunteers.

![Figure 4](image1)

**Figure 4** Mean normalised expression of mammaglobin detected in blood samples from patients with metastatic breast cancer and healthy volunteers when samples were analysed by real-time RT–PCR.

![Figure 5](image2)

**Figure 5** Mean normalised expression of PIP detected in blood samples from patients with metastatic breast cancer and healthy volunteers when samples were analysed by real-time RT-PCR. (PIP, prolactin-inducible peptide).

| Healthy volunteers (n = 20) | Patients (n = 40) |
|-----------------------------|------------------|
| **Number of genes positive** | **Level of specificity** | **Level of Specificity** |
|                             | 95%           | 100%           | 95%           | 100%           |
| None                       | 18 (90%)      | 20 (100%)      | 7 (18%)       | 11 (28%)       |
| \( \geq 1 \)                | 2 (10%)       | 0 (0%)         | 33 (83%)      | 29 (73%)       |
| All 3                      | 0 (0%)        | 0 (0%)         | 6 (15%)       | 4 (10%)        |

CK19 = cytokeratin 19; PIP = prolactin-inducible peptide; RT–PCR = reverse transcriptase–polymerase chain reaction. Positive samples defined using either the upper 95% confidence interval of levels in the healthy volunteers to describe positive samples or using a threshold that provided 100% specificity in healthy volunteers.
metastatic breast cancer and in control populations. Therefore, it is important to compare directly techniques, in order to ascertain the optimal approaches to use.

Using LSC following IMS and cell filtration more circulating epithelial cells were detected in the blood of patients with metastatic breast cancer than in control healthy volunteers. Similarly, the expression of CK19, mammaglobin and PIP was found to be significantly higher in the blood of patients with metastatic breast cancer than in controls. A cutoff for positivity was defined for all three techniques based on 95% confidence intervals of mean cell numbers in the control population. On this basis, 30% of the patient samples were positive by filtration, 48% by IMS enrichment and 35–60% using real-time RT–PCR, depending on the test gene used. Thus, positive samples using the cytometric techniques and real-time RT–PCR are within the previously observed ranges using such techniques to detect circulating epithelial cells in the blood of patients with metastatic breast cancer (Smith et al, 2000; Bosma et al, 2002; Baker et al, 2003; Stathampolou et al, 2003; Ring et al, 2004).

This study demonstrates the benefit of using more than one target for RT–PCR amplification. If using CK19, mammaglobin or PIP alone, then 60, 45 and 35% of samples would be expected to be positive. However, 82% of patients were positive for at least one of these markers. The relatively low concordance values for positivity between the different markers (50–65%) probably reflects not only differences in sensitivity but also that the different markers are identifying different populations of cells: in fact, of the 16 CK19-negative patients, five were PIP positive. This reinforces the benefit of using a multiple marker PCR assay. Similar results were reported by Taback and colleagues who used RT–PCR to assess the expression of four genes in the blood of 65 patients with breast cancer (stage I–IV) and 40 normal volunteers (Taback et al, 2001). Individual markers were detected in 11–37% of patients, but at least one marker was detected in the blood of 69% of patients. These two studies demonstrate that although no one marker may be ideal for the detection of circulating tumour cells, the use of a multiple marker assay may significantly improve sensitivity.

In a further study published by Baker et al, 13 out of 20 (65%) patients with metastatic breast cancer had overexpression in their blood of at least one of the six test genes (Baker et al, 2003). This study also provides evidence that enrichment using porous barrier density gradient centrifugation reduces background expression of CK19 and MUC1 compared with a Ficoll density gradient similar to that used in the current study. Therefore, it may prove to be possible to further improve on the real-time RT–PCR results reported here by modifying the enrichment step.

As far as comparisons between the techniques are concerned, the CK19 RT–PCR assay was found to be more sensitive than cell filtration followed by LSC, although there was no significant difference compared with IMS followed by LSC. However, when all three markers are taken into consideration, samples from patients with breast cancer are more likely to have a positive result with at least one of the markers than either of the two cytometric techniques (both \( P < 0.001 \)). These results show that when a multiple marker real-time RT–PCR assay is used, its sensitivity exceeds that of both an established and a novel cytometric technique. Previous studies using nonquantitative RT–PCR also reveal more positive results when patient samples are analysed by PCR than by immunocytochemical techniques (Schoenfeld et al, 1997; Lambrechts et al, 1999). However, these results may be compromised by lower specificity (Lambrechts et al, 1999). Slade and colleagues used a competitive PCR reaction to enable relative quantification of samples and to establish a cutoff point for positivity based on analyses of the blood of normal individuals (Slade et al, 1999). Using such an approach for CK19, it was found that 50% of samples from patients with metastatic breast cancer were positive by PCR compared with 42% of samples analysed using immunocytochemistry (Smith et al, 2000).

Aside from the relative sensitivities and specificities of the different techniques, the correlations between the techniques and their concordance are also of interest. Good correlations were observed between cell numbers detected by the two cytometric techniques and between real-time RT–PCR for CK19 and IMS enrichment. However, when a cutoff for positivity is established, the concordance between techniques was between 68 and 75%, meaning that although the sensitivity of techniques may be similar, the samples being called positive and negative are not necessarily the same. Other studies show qualitative concordance between RT–PCR-based assays and immunocytochemistry of 71–77% (Schoenfeld et al, 1997; Smith et al, 2000). In the case of the cytometric techniques, the limited concordance may reflect the differences in specimen handling. The differences between the real-time RT–PCR technique and the cytometric techniques may be further explained by the different markers detected and the determination of expression at an RNA as opposed to a protein level. An important corollary of this observation is that cytokeratin-positive cells identified by one technique should not be assumed to have the same significance as those detected by another technique: the populations of cells detected may be distinct and have different clinical, biological and prognostic significance.

The quantitative nature of the real-time RT–PCR assay makes it an ideal technique by which to establish a normal range of expression for markers in controls. This was a comparative study and hence the same approach was used in the cytometric techniques. Such an approach predetermines the specificity of the assay. In order to assess fully the sensitivity and specificity of these techniques, the cohort presented in this study should be regarded as a training set. The established cutoffs for positivity should now be used in an independent validation set, to establish fully the clinical performance of the technique. A second issue when considering RNA-based assays such as RT–PCR is that RNA is inherently labile and therefore blood should be analysed as soon as possible after sampling. This may not be practical if samples (as seems likely) need to be transferred to a central laboratory for analysis. This practical issue may limit the widespread clinical utility of the PCR approach, although recently blood RNA tubes have become available (PAXgene™, Qiagen Ltd, Crawley, UK), which may be able to overcome this problem.

These caveats aside, these data show that using a real-time RT–PCR assay there is evidence of circulating tumour cell dissemination in the blood of up to 80% of patients with metastatic breast cancer. Although such analysis of blood samples may enable the monitoring of responses to treatment, in the absence of further characterisation of the significance of the cells, the clinical utility of the approach may be limited. The ascertainment of circulating individual cell phenotype at relapse and during therapy could significantly enhance our understanding of mechanisms of resistance and facilitate targeting of therapy, but is only likely to be possible using cytometric techniques. Circulating tumour cell detection theoretically would have its greatest impact in the management of early breast cancer, where risk stratification may be improved and responses to adjuvant therapy and the detection of relapse monitored. However, not all patients even with metastatic disease have detectable circulating tumour cells, so whether a significant role will emerge will depend on applying such techniques in large prospective studies in patients with early breast cancer.

In summary, the data presented suggest that the use of a multiple marker real-time RT–PCR assay may be a more sensitive technique by which to detect circulating breast cancer cells than the use of a novel or an established cytometric technique. This does not necessarily mean that the PCR-based assay is superior to
the other techniques, as different information may be acquired by the different approaches. If the aim of the technique is to assess blood ‘tumour burden’, such as if one is monitoring responses to treatment, then real-time RT–PCR may be the superior approach. However, if it is desirable to visualise cells directly and perhaps perform further phenotypic or molecular characterisation, then a cytometric approach may be preferable.

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