ORIGINAL ARTICLE

HER2 amplification status in breast cancer: a comparison between immunohistochemical staining and fluorescence in situ hybridisation using manual and automated quantitative image analysis scoring techniques

C M Ellis, M J Dyson, T J Stephenson, E L Maltby

Aims: To compare the results of breast cancer sections with HercepTest\textsuperscript{TM} immunohistochemistry (IHC) scores ranging from 0 to 3+ with fluorescence in situ hybridisation (FISH) for HER2 amplification. The HER2 digital scoring application of the Micrometastasis Detection System (MDS\textsuperscript{TM}) was used, together with manual scoring of FISH and HercepTest, to determine whether this system provides an accurate alternative.

Methods: Paraffin wax embedded sections were stained using HercepTest and analysed by eye and automated quantitative image analysis. FISH was performed using the PathVysion\textsuperscript{TM} fluorescent probe and scored by eye and automated quantitative image analysis using MDS.

Results: Of 114 cases, 26\% were amplified by FISH, whereas only 18\% scored 3+; 32\% of IHC 2+ cases were amplified by FISH, and one showed borderline amplification. Six percent of IHC negative cases (0 or 1+) were amplified by FISH, and one showed borderline amplification. Of IHC 3+ cases, 10\% were non-amplified by FISH. Classification discrepancies were seen in 18\% of HercepTest cases scored by eye and using the MDS system. MDS was consistent with visual FISH scoring and correctly differentiated most ambiguous visual IHC scores.

Conclusions: FISH provides a more accurate and consistent scoring system for determining HER2 amplification than HercepTest. The MDS system provides a reliable, consistent alternative to visual IHC and FISH scoring. IHC is still a valuable technique to aid in identification of isolated or heterogeneous tumour populations for subsequent FISH analysis, and a combined FISH and HercepTest approach to all breast cancer cases may be the most efficient strategy.

Abbreviations: DDW, double distilled water; FISH, fluorescence in situ hybridisation; IHC, immunohistochemistry; MDS, Micrometastasis Detection System

The current literature suggests that fluorescence in situ hybridisation is more accurate and easier to interpret than immunohistochemistry. A study comparing HercepTest and FISH scores found that HercepTest resulted in a poor prognosis. Historically, testing for amplification of the HER2/neu gene has been performed using an immunohistochemical (IHC) staining technique, most often the HercepTest\textsuperscript{TM} (Dako, Glostrup, Denmark). The technique involves antibody specific staining of the encoded protein on the cytoplasmic membrane. A darker stain indicates the presence of more protein and therefore, theoretically, gene amplification should be present. Stained tissue sections are categorised as: 0, no staining; 1+, light staining; 2+, equivocal moderate staining; 3+, dark membrane staining (Fig 1). Patients with tumours showing a 3+ staining pattern may benefit from treatment with the drug Herceptin\textsuperscript{TM}. All patients are considered for conventional chemotherapy regardless of their HER2 status.

The accurate detection of HER2 amplification has now become extremely important in the treatment of breast cancer cases may be the most efficient strategy.
cancer because HER2 targeted treatment is available with the recombinant humanised antibody, Trastuzumab™ (Herceptin™; Genentech, South San Francisco, California, USA).9 Furthermore, HER2 amplification has been associated with a negative response to widely used hormonal drugs, such as tamoxifen.10 FISH has also been shown to be more predictive than the HercepTest with regard to the response to Herceptin.11–14

In our study, automated image analysis was performed on 114 breast cancer samples using the HER2 digital scoring application of the Micrometastasis Detection System (MDS™; Applied Imaging, San Jose, California, USA).

**METHODS**

**IHC**

Paraffin wax embedded sections (4 μm thick) were mounted on X-tra™ (Surgipath Medical Industries, Richmond, Illinois, USA) slides and stained with HercepTest using DakoCytomation standard protocols.

**FISH**

Unless stated otherwise, procedures were performed at room temperature. Paraffin wax embedded sections (4 μm thick) mounted on X-tra™ slides were dewaxed for 2 × 10 minutes in 50 ml xylene at room temperature, dehydrated for five minutes in 50 ml ethanol, then washed for between 20 and 23 minutes in 50 ml of 0.2N HCl. The slides were then washed in 50 ml double distilled water (DDW) for two minutes and transferred to 50 ml Zymed (South San Francisco, California, USA) heat pretreatment solution at 95˚C for 80 to 110 minutes. The slides were then washed in 50 ml of DDW for 2 × 3 minutes; 90 μl of Zymed digestion enzyme solution was then applied to a 22 × 50 coverslip and overlayed on to the slide. The slide was then incubated at 38˚C in a wet box for 50 to 70 minutes. Slides were then washed in 50 ml DDW for 3 × 2 minutes before being dehydrated through a 50 ml ethanol series: 70%, 95%, and 100% for two minutes each. The slides were then left to dry before PathVysion probe application. The sample DNA and

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**Figure 1** Images of invasive breast carcinoma captured on the MDS. (A–D) HercepTest immunohistochemical staining: (A) 0 (negative), (B) 1+, (C) 2+, and (D) 3+ (positive). (E, F) Fluorescence in situ hybridisation using the PathVysion probe. (E) A non-amplified ratio, (F) an amplified ratio.
probe DNA were co-denatured at 72°C for five minutes and then allowed to hybridise at 37°C overnight on a PTC-200 thermal cycler (MJ Research, Waltham, Massachusetts, USA). The slides were then washed in 50 ml of 0.4× saline sodium citrate/Tween 20 at 73°C for two minutes and then transferred to 50 ml 2× saline sodium citrate/Tween 20 at room temperature for 30 seconds. Ethanol series dehydration was performed as before and the slides were air dried in darkness. The slides were coverslipped with 20 μl of counterstain (20 μl mounting medium with DAPI in 1000 μl mounting medium for fluorescence; Vector Laboratories, Burlingame, California, USA).

**ANALYSIS**

Processed IHC and FISH slides were analysed by eye using published recommendations for scoring.1

The MDS is a computer controlled scanning microscope with capture capabilities using either brightfield (for IHC) or fluorescence filter wheels (for FISH). Quantitative analysis of captured IHC or FISH images can be performed using this software. The HER2 digital scoring application developed for the MDS was used for scoring the HercepTest and for quantifying the HER2 FISH amplification ratio. For the HercepTest, positive and negative batch control slides are analysed to train the system for the range of staining intensity. An image from a suitable invasive tumour location is captured from each sample slide from a given batch. Using the stored batch control intensity range, an automatic analysis is then performed. This process takes less than two minutes for each slide.

For FISH, a representative invasive tumour location is captured using Z stacking. This is an automated focus ability that takes captures of several focal planes and then combines them into a single image. Automatic analysis for the presence of HER2 gene amplification using Applied Imaging protocols is then performed.

**RESULTS**

Of the 114 cases that we analysed 30 were found to be amplified by FISH (ratio of HER2/17cen > 2.2), giving an amplification rate of 26%. This correlates with the published range of 20–30% of all breast cancer cases.11 Borderline amplification by FISH is classified as an amplification ratio between 1.8 and 2.2. When scoring the HercepTest by eye, 72 cases were determined to be either 0 (negative) or 1+ (probably negative), 22 as 2+, and 20 as 3+ (positive). Table 1 shows the full breakdown of data by FISH and HercepTest.

In two of the cases studied, two different tumour populations were seen in the same section. In each case, one minor tumour population was amplified by FISH and the other major tumour population was not amplified by FISH. In rare cases such as these the IHC slides provide a useful method of locating tumours. The IHC and FISH slides from a particular sample can be used in unison to facilitate the definition of separate tumour populations and to distinguish between invasive and ductal/lobular disease.

The IHC scoring facility on the MDS provides evidence for the accuracy of automated HercepTest scoring. When comparing HercepTest scores assessed by eye and MDS there was a difference in 21 (18%) of the 114 cases. All 14 IHC 2+ cases scored by eye were categorised by the MDS system as either 1+ or 3+, and 13 were also validated by FISH. Furthermore, of three 2+ cases scored by the MDS system and differentiated by eye, FISH validated the MDS in two cases. Table 2 shows the discrepancies seen when scoring the HercepTest and FISH by eye and with the MDS.

Scoring of FISH on HER2 amplified cases using the MDS was highly consistent and much less variable than when scoring was performed by eye. However, on non-amplified cases the MDS was slower, although it still retained its accuracy. Figure 2 shows the varying scores within amplified cases between human analysers and the MDS. The MDS scores were consistently lower than the visual scores, regardless of the analyser, but concordant FISH classification status was seen in every case.

**DISCUSSION**

There is evidence to support both the association between HER2 gene amplification and a positive response to the drug Trastuzumab, and the association between HER2 gene amplification and a negative response to drugs such as tamoxifen.1 The reverse scenario is equally concerning with regard to the potential side effects of Trastuzumab and the denial of conventional treatment. Therefore, appropriate prescription of these drugs is crucial in both human and financial terms.

The HercepTest false negative rate in our series was around 7% (percentage of 0 and 1+ cases; n=72) when FISH was used as a secondary test. We also found that the HercepTest false positive rate was around 10% (percentage of 3+ cases; n=20) when FISH was used as a secondary test. This confirms suggestions from previous data.

In our series, when using FISH, three more patients were eligible for treatment with Herceptin than might have been indicated by the HercepTest, and three patients were not eligible for treatment although the HercepTest found them positive. These HercepTest false negatives are not a reflection of the oversensitivity of the FISH test, but result from insensitivity of the IHC test, as illustrated by the accurate correlation with published data and the percentage of FISH amplified cases in total.

"In almost every case, immunohistochemical scoring by MDS agreed with the fluorescence in situ hybridisation result, suggesting that this method of scoring is more accurate than visual scoring."

The IHC scoring facility on the MDS is a quick, highly accurate, and consistent method of scoring immunooaasays in breast cancer samples. It also allows the operator the option to archive images and to relocate back to regions of interest on the slide. We found discrepancies between the HercepTest scored either visually or by the MDS system, and three cases (d, o, and s in table 2) would have been processed by FISH to confirm whether or not amplification exists. The MDS system provides a consistent scoring tool, which shows high levels of accuracy and is a time saving alternative to scoring by eye. In almost every case, IHC scoring by MDS agreed with the FISH result, suggesting that this method of scoring is more accurate than visual scoring. The three cases that showed discordance with FISH (i, j, and n in table 2) can be explained...
by either an increased copy number of chromosome 17 and hence a $3^+\text{st}$ staining pattern but a non-amplified ratio, or a breakdown of the protein product.

Scoring HER2 FISH with the MDS appears to be useful when scoring amplified and borderline cases. Again, the system can relocate to a region previously captured and allows archiving of cases for future reference and study. It also provides an analysis of the data. The results obtained are more consistent than when scored by eye, which is especially important when scoring borderline, ambiguous cases. When analysing amplified cases, the level of hybridisation to the HER2 gene must be counted by eye and in some cases only an estimate can be made because the probe signals cluster closely together (fig 1F). The Z stacking auto focus facility of the MDS allows the software to count the number of HER2 copies accurately.

With regard to FISH, the MDS system relies on a good digestion in the experimental preparation to define an individual cell for scoring. There also appear to be problems capturing small non-amplified cells because the software is designed to screen out smaller non-neoplastic cells. Therefore, in these cases it is much quicker for two scientists to score and check by eye than it is to score on the MDS.

The most important thing that can be learnt from our study is that FISH provides a more accurate and consistent scoring system for determining HER2 amplification than the HercepTest. However, it has been suggested that FISH is a more expensive and time consuming test. One recent study has now disputed this and has shown that the use of FISH alone is more cost effective than either the use of IHC alone or a combination of IHC and FISH.16

IHC staining still remains a valuable aid in the identification of isolated or heterogeneous tumour populations for subsequent FISH analysis. The use of the MDS has shown that the accuracy and consistency of an automated quantitative image analysis system enhances a strategy using a combined FISH and IHC approach on all breast cancer cases. This may be the most efficient, if not the most cost effective, strategy.

ACKNOWLEDGEMENTS

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### Table 2

<table>
<thead>
<tr>
<th>Sample</th>
<th>HercepTest score</th>
<th>FISH result</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>a</td>
<td>2</td>
<td>Amplified</td>
<td>Would analyse by FISH routinely but MDS accurate</td>
</tr>
<tr>
<td>b</td>
<td>2</td>
<td>Non-amplified</td>
<td>MDS accurate</td>
</tr>
<tr>
<td>c</td>
<td>0</td>
<td>Amplified</td>
<td>2 tumour populations seen: 1 amplified</td>
</tr>
<tr>
<td>d</td>
<td>3</td>
<td>Non-amplified</td>
<td>MDS accurate</td>
</tr>
<tr>
<td>e</td>
<td>2</td>
<td>Amplified</td>
<td>Would analyse by FISH routinely but MDS accurate</td>
</tr>
<tr>
<td>f</td>
<td>2</td>
<td>Non-amplified</td>
<td>MDS accurate</td>
</tr>
<tr>
<td>g</td>
<td>2</td>
<td>Non-amplified</td>
<td>MDS accurate</td>
</tr>
<tr>
<td>h</td>
<td>2</td>
<td>Non-amplified</td>
<td>MDS accurate</td>
</tr>
<tr>
<td>i</td>
<td>3</td>
<td>Non-amplified</td>
<td>MDS accurate</td>
</tr>
<tr>
<td>j</td>
<td>3</td>
<td>Non-amplified</td>
<td>MDS accurate</td>
</tr>
<tr>
<td>k</td>
<td>2</td>
<td>Non-amplified</td>
<td>MDS accurate</td>
</tr>
<tr>
<td>l</td>
<td>2</td>
<td>Non-amplified</td>
<td>MDS accurate</td>
</tr>
<tr>
<td>m</td>
<td>2</td>
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<td>MDS accurate</td>
</tr>
<tr>
<td>n</td>
<td>2</td>
<td>Amplified</td>
<td>MDS does not correspond to FISH</td>
</tr>
<tr>
<td>o</td>
<td>2</td>
<td>Amplified</td>
<td>2 tumour populations seen: 1 amplified</td>
</tr>
<tr>
<td>p</td>
<td>2</td>
<td>Non-amplified</td>
<td>MDS accurate</td>
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<tr>
<td>q</td>
<td>2</td>
<td>Amplified</td>
<td>Would analyse by FISH routinely but MDS accurate</td>
</tr>
<tr>
<td>r</td>
<td>2</td>
<td>Amplified</td>
<td>Would analyse by FISH routinely but MDS accurate</td>
</tr>
<tr>
<td>s</td>
<td>2</td>
<td>Amplified</td>
<td>Would not analyse by FISH routinely</td>
</tr>
<tr>
<td>t</td>
<td>2</td>
<td>Non-amplified</td>
<td>MDS accurate</td>
</tr>
<tr>
<td>u</td>
<td>1</td>
<td>Amplified</td>
<td>MDS accurate</td>
</tr>
</tbody>
</table>

FISH, fluorescence in situ hybridisation; MDS, Micrometastasis Detection System.

Figure 2 Comparison between the average of two human analysers and the MDS for 23 randomly selected samples when scoring amplification ratio by fluorescence in situ hybridisation.
REFERENCES


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