Can breast MRI help in the management of women with breast cancer treated by neoadjuvant chemotherapy?

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Contrast-enhanced (CE) MRI was used to monitor breast cancer response to neoadjuvant chemotherapy. Patients underwent CE MRI before and after therapy, together with conventional assessment methods (CAM). CE MRI was carried out at 1.5 T in the coronal plain with 3D sequences before and after bolus injection. An expert panel determined chemotherapy response using both CE MRI and CAM. Histopathological response in the surgical specimen was then used to determine the sensitivity and specificity of CE MRI and CAM. In total, 67 patients with 69 breast cancers were studied (mean age of 46 years). Tumour characteristics showed a high-risk tumour population: median size 49 mm; histopathological grade 3 (55%); oestrogen receptor (ER) negative (48%). Histopathological response was as follows: – complete pathological response (pCR) 17%; partial response (pPR) 68%; no response (NR) 15%. Sensitivity of CAM for pCR or pPR was 98% (CI 91 – 100%) and specificity was 50% (CI 19 – 81%). CE MRI sensitivity was 100% (CI 94 – 100%), and specificity was 80% (CI 44 – 97%). The absolute agreement between assessment methods and histopathology was marginally higher for CE MRI than CAM (81 vs 68%; P = 0.09). In 71%, CE MRI increased diagnostic knowledge, although in 20% it was judged confusing or incorrect. The 2nd MRI study significantly increased diagnostic confidence, and in 19% could have changed the treatment plan. CE MRI consistently underestimated minimal residual disease. In conclusion, CE MRI of breast cancer proved more reliable for predicting histopathological response to neoadjuvant chemotherapy than conventional assessment methods.

Keywords: breast cancer; breast MRI; disease response; diagnostic impact; neoadjuvant chemotherapy

Breast cancer treatment has advanced over the past two decades, achieving a better outcome for women (Peto et al, 2000). One of several factors contributing to improved survival is chemotherapy, which is now widely used particularly for women under 50 years and those with high-grade ER negative or lymph node-positive tumours. When chemotherapy is given prior to definitive surgical treatment, there are various potential benefits (Valero et al, 2002):

- Primary tumour response provides evidence of responsiveness to the chosen chemotherapeutic agent and predicts local control (Chollet et al, 2002)
- A large tumour may become amenable to breast-conserving surgery (Makris et al, 1997; Fisher et al, 1998)
- An ineffective therapeutic agent may be changed to one with clinical effect, or surgery may be advanced, thereby saving cost and patient suffering (Sotiriou et al, 2002; Pusztai et al, 2003)

There is no evidence of benefit in overall or disease-free survival as a result of the use of chemotherapy in the neoadjuvant setting (Fisher et al, 1998; Green and Hortobagyi, 2002). Marginally statistically significant treatment-by-age interactions appear to be emerging for survival and disease-free survival, suggesting that younger patients may benefit from preoperative therapy, whereas the reverse may be true for older patients (Wolmark et al, 2001). It is logical to initiate systemic treatment early in those patients with a high risk of micro-metastatic disease and to monitor the response. Conventional tools to monitor response include measurement of maximum tumour diameter with callipers; loss of signs of inflammation and reduction in lymph node size. Serial core biopsy has been used to show changes in tissue morphology by standard histological criteria and more recently with molecular markers (Sotiriou et al, 2002; Pusztai et al, 2003). Response of the primary tumour or regional nodes may be monitored radiologically with mammography or ultrasound (Helvie et al, 1996; Vinnicombe et al, 1996; Huber et al, 2000); and newer imaging modalities (Mankoff et al, 1999; Tiling et al, 2001), such as contrast-enhanced (CE) MRI are being increasingly used. The literature contains many single institution reports of the use of breast MRI for monitoring tumour response or predicting tumour extent prior to surgery. Table 1 summarises the literature to date.
on the use of breast MRI to monitor neoadjuvant chemotherapy. It can be seen from the right-hand column of this table that the findings have varied very substantially. Breast MRI appears to hold promise for this purpose, and provides measurements that are more accurate than clinical or ultrasound assessment (Davis et al., 1996; Esserman et al., 1999). It is important to know whether predictions of residual tumour are accurate. Breast MRI has good sensitivity for invasive tumour but relatively poor specificity in untreated disease (Mussurakis et al., 1996). The limitations are more apparent for invasive lobular carcinoma and in situ (Kinkel and Hylton, 2001). Contrast-uptake characteristics have been shown to be altered by chemotherapy, which may potentially diminish sensitivity and impair specificity (Rieber et al., 1997). Contrast uptake characteristics have been used to give an early indication of tumour response using innovative functional MR and computer methods (Padhani, 2002; Delille et al., 2003). It is therefore an open question whether CE MRI would be helpful in monitoring response, or whether limitations emerging from studies undertaken concurrent with this one would make it unreliable.

In this study, we have investigated the impact of breast MRI on decision-making. Almost all of our patients underwent mastectomy after neoadjuvant chemotherapy. It is therefore possible to correlate the extent of tumour detectable on breast MRI following chemotherapy with histopathology of the mastectomy specimen. The aims of our study were:

1. To determine whether CE MRI is an accurate indicator of tumour response to neoadjuvant chemotherapy in comparison to the histopathology gold standard.
2. To assess whether CE MRI increases the diagnostic confidence above that achieved by conventional assessment methods (CAM) (i.e. clinical examination, ultrasound and mammography).
3. To determine whether CE MRI can accurately identify a subset of patients who might be suitable for breast-conserving surgery following neoadjuvant chemotherapy.

TABLE 1 Existing literature on the use of breast MRI to monitor response to chemotherapy

| Author                        | Year        | Case number | Measurement | Pathology response |
|------------------------------|-------------|-------------|-------------|--------------------|
| Knopp et al (1994)           | 1994        | 3 case studies |            | Promising prediction |
| Gilles et al (1994)          | 1994        | 18          |             | (15 out of 18) 83% |
| Abraham et al (1996)         | 1996        | 39          |             | MRI correctly predicted residual disease |
| Rieber et al (1997)          | 1997        | 13          |             | Underestimates residual tumour |
| Trecate et al (1998)         | 1998        | 27          |             | Valid tool for monitoring response |
| Tsuibo et al (1999)          | 1999        | 31          |             | Valuable for predicting response and margins |
| Weatherall et al (2001)      | 2001        | 22          | Cc0.93MR, 0.63 mam | Accurate assessment of residual tumour |
| Nakamura et al (2001)        | 2001        | 15          |             | Predict possibility of WLE. Mapping of dendritic tumours |
| Drew et al (2001)            | 2001        | 17          | Sensitivity 100% | Accurate method of predicting residual tumour |
| Eiserman et al (2001)        | 2001        | 33          | R0.89, C0.60 | Patterns of tumours identified |
| Partridge et al (2002)       | 2002        | 52          |             | MRI detected all residual disease |
| Balu-Maestro et al (2002)    | 2002        | 60 in 51 women |             | Valuable tool for size & multifocality prior to surgery |
| Rieber et al (2002)          | 2002        | 58          | Accuracy NR 83.3%, PR 82.4% | Good for assessing response to chemo, unreliable in CR cases where 66.7% had residual tumour |
| Cheung et al (2003)          | 2003        | 33          | Extraction flow product as marker of tumour response |
| Delille et al (2003)         | 2003        | 14          | EFPacc, accurate measure (P<0.02) |

**PATIENTS AND METHODS**

**Patients and chemotherapy**

The study had approval from the Local Research Ethics Committee for Cambridgeshire, and all women had given informed consent to participate. It is a retrospective review of 67 women with CE-MRI before and after neoadjuvant chemotherapy for breast cancer. The first 30 women were recruited between 1997 and 2001 to a prospective pilot study of CE MRI. The remaining 37 women were consecutively enroled once CE MRI monitoring became the standard of care at our centre in 2001. The decision to give neoadjuvant therapy was made by a multidisciplinary team, and in general included women less than 60 years with large, inflammatory or high-grade tumours. Chemotherapeutic regimens used in this study varied according to whether patients were treated in ongoing clinical trials (Appendix A). All cases had undergone mammography, ultrasound and core biopsy prior to the decision to treat with neoadjuvant chemotherapy, and the results of these tests were used by the team to determine which trials they should enter.

**Contrast enhanced MRI (CE MRI)**

The patients had two MRI studies, at the start of treatment and prior to surgery. The initial MRI study was usually undertaken prior to the first course of chemotherapy except in occasional cases where limitations of access to MRI resulted in imaging after the first treatment had been given. The 2nd MRI study was performed after all chemotherapy and prior to surgery. The studies were undertaken on a 1.5 T magnet from GE Signa Echospeed, which was upgraded in 2002 with Excite software. The MRI technique is similar to that described for the MARIBS study (Brown et al., 2000), with the exceptions that the dose of gadolinium was 0.16 mmol kg⁻¹ body weight, and no proton density study was undertaken in our study (Scanning protocol – Appendix B). MRI interpretation used a synthesis of dynamic and morphological...
features, including the pattern of contrast uptake and washout characteristics, and the studies were examined at the console to allow dynamic contrast enhancement curves at any point in the 3D volume of the dynamic acquisition. The studies were reported by radiologists and presented to the multidisciplinary team for decisions on patient management.

Conventional methods of assessment
All patients had a clinical assessment prior to each of six cycles of chemotherapy. This included caliper measurement of tumour size, degree of inflammation and size of palpable axillary nodes. All patients underwent mammography, ultrasound and core biopsy for diagnosis. Ultrasound was carried out after three and six cycles of chemotherapy. Conventional breast imaging comprised mammography, ultrasound and core biopsy according to the guidelines of the United Kingdom National Health Service Breast Screening Programme (NHSBSP) (Wilson et al, 2001).

Surgery
Surgery normally took place less than 4 weeks after the last course of chemotherapy. All patients underwent either mastectomy or wide local excision (WLE) with axillary lymph node dissection.

Histopathological response
Response to treatment was scored by standard UICC (Therasse et al, 2000) categories of Complete Response (CR), Partial Response (PR), Non-Response (NR), Disease Progression (DP). The reports were produced at the time of clinical care without particular knowledge of the MRI findings. It should be noted that after chemotherapy the criteria for grading cannot be applied, and so when grades are given here they are based on core biopsy (Harris et al, 2003).

There are no recognised standard criteria for grading of histological response following primary chemotherapy. In this study, mastectomy specimens were handled according to the NHSBSP guidelines (NHSBSP publication No. 3, 1993). In addition, in those cases where no macroscopic tumour could be identified, extensive sampling of the original tumour site (as identified by imaging methods and clinical examination prior to treatment) was performed.

On histological examination, where distinct in situ and invasive tumour was identified this was morphologically compared with tumour seen on the original core biopsy. The ratio of tumour cell mass to surrounding tissue was evaluated. If degenerative changes within tumour cells were present this was noted. The nature of the tissue surrounding tumour cells was noted, viz was it a cellular stroma similar to that seen in the original biopsy or was it inflamed, hypocellular scar tissue.

Thus:
(1) No histological response. Tumour resembling that in the original core biopsy was present and it extended within the breast tissue for the same or a greater distance than originally estimated by imaging and clinical examination. The nature of the tissue surrounding the tumour was cellular and did not resemble bland scar tissue.

(2) Complete histological response. Absence of invasive carcinoma cells/or <1 single, degenerate carcinoma cell/10 high-power fields. Absence of in situ carcinoma.

(3) Partial histological response. Tumour resembling that in the original core biopsy but which was considerably reduced in size (it is necessary to approximate size parameters used in clinical response staging). Presence of residual microscopic foci of invasive carcinoma but with reduced tumour cell/stroma ratio or residual pure in situ carcinoma.

Assessment of sensitivity, specificity of conventional assessment methods and CE MRI
Original records were used to score clinical response using CAM, CE MRI response and histopathological response, and from these sensitivity and specificity were calculated. Pathological response was regarded as the gold standard against which all evaluations of response were measured.

Assessment of diagnostic impact and confidence of knowledge
For the purposes of this study, all cases were reviewed by the multidisciplinary team and a research form completed to record the diagnostic impact. The review team consisted at minimum of a radiologist, a surgeon, an oncologist and a histopathologist, all expert in management of breast disease. The multidisciplinary review procedure is shown in Figure 1. The clinical notes were examined for the clinical findings recorded at the time of clinical care. The mammograms and ultrasound images with their original reports were presented as in a standard multidisciplinary team meeting. The MRI studies were presented to the clinical team using the images and reports taken at the time of clinical care of the patients. The multidisciplinary team were unaware of the histopathology findings when they interpreted the results of the MRI and other presurgical examinations. The pathology was presented from the histological report at the time of clinical care, and did not take particular account of MRI knowledge. The added value of MRI imaging was given a score 1…6 (Figure 2) by the team, taking into account the additional contribution made by the MRI study beyond that available with conventional breast imaging (Shuman et al, 1985a,b). A visual analogue scale was used to record confidence of knowledge with conventional breast imaging alone and with the addition of MRI (Blanchard et al, 1997). Confidence of knowledge included knowledge of the size, the contour, the extent and the suitability for any form of surgery. The two measures, added value and confidence of knowledge are complementary, since confidence of knowledge is an important aspect of value to a clinician. These are subjective clinical judgements.

Statistical analyses
Proportions in independent samples were compared using Fisher’s exact test or linear by linear $\chi^2$ test as appropriate. Parametric and nonparametric paired comparisons of mean and median values were performed using the paired $t$-test and McNemar test, respectively. The relationship between initial tumour size and subsequent tumour response was assessed using one-way analysis of variance. Statistical significance was accepted at the conventional $P<0.05$ threshold.

RESULTS
The study evaluated 69 cancers in 67 women including two bilateral cases. In all, 67 cancers were treated with mastectomy and two with WLE. The mean age of the patients at the time of the 1st MRI was 46.2 years (range 24 to 69 years). On initial radiological examination, the median size of the 69 tumours was 49.0 mm (range 9 – 100 mm). The core biopsy demonstrated that six (9%) tumours were grade 1, 25 (36%) grade 2 and 38 (55%) grade 3. These therefore represent relatively large, poor prognosis cancers in younger women. In total, 36 (52%) tumours were oestrogen receptor (ER) positive, and 33 (48%) ER negative, 21 out of 26 (81%) patients where the progesterone receptor (PR) status was recorded were PR negative. At definitive surgery, patients had a median of 13 nodes retrieved (range 0 – 42); 17 patients (25%) had more than four nodes involved, 23 (33%) had between one and
Clinical findings, and a further 16 (23%) lesions were close to the nipple–areolar complex. After the initial clinical and radiological evaluation, a total of 52 (75%) tumours were considered not suitable for breast conservation within the protocols of our unit, were known to be multifocal or diffuse lesions by their imaging or clinical findings, and a further 16 (23%) lesions were close to the nipple–areolar complex. After the initial clinical and radiological evaluation, a total of 52 (75%) tumours were considered not suitable for breast conservation within the protocols of our unit, where skin involvement by inflammation, very large size, multifocality and proximity to the nipple–areolar complex are all viewed as reasons for not using breast-conserving surgery. The median period between the 1st and 2nd MR imaging was 119 days (range 48–165 days). The median period between the 2nd MR and surgery was 24 days (range 2–77 days). The neoadjuvant chemotherapy included cases in trials, notably Anglo-Celtic 2, and cases off trial with a variety of regimens (Appendix A).

Estimation of response

Based on the gold standard histopathology findings, 12 lesions (17%) had complete response, 47 (68%) had partial response, 10 (15%) had no response following neoadjuvant chemotherapy (Table 2). There was no significant association between the grade of the tumour and the extent of disease response following chemotherapy. In total, 24% (nine out of 38) of grade 3 tumours had complete response to chemotherapy compared to 12% (three out of 25) of grade 2 tumours and 0% (zero out of six) of grade 1 tumours ($P=0.10$). Four out of 36 (11%) ER-positive tumours exhibited complete response at pathology compared to six out of 21 (29%) of tumours that were ER- and PR negative ($P=0.15$).

Lesions that did not respond or progressed after neoadjuvant chemotherapy were slightly, but nonsignificantly, larger than lesions that exhibited partial or complete response (mean size 56 vs 49 vs 48 mm; $P=0.56$). Tumour response was clearly related to the extent of node involvement. In all, 97% (28 out of 29) of lesions with no node involvement had complete or partial response compared to 87% (20 out of 23) of lesions with between one and four positive nodes and 65% (11/17) lesions with more than four involved nodes ($P<0.01$).

Conventional assessment methods and MRI in the estimation of tumour response

The clinical assessment of tumour response agreed with the pathology assessment in 47 out of 69 tumours (68%) (Table 2). CAM underestimated the degree of response in four cases, and overestimated the degree of response in 18 cases. In one case, a tumour which was categorised as completely responsive based on the clinical data had no response evident at pathological examination. CAM correctly identified 58 out of 59 lesions with complete or partial response based on pathology findings (sensitivity 98%; 95% confidence interval (CI) 91–100%). However, five out of 10 tumours with no response were incorrectly categorised as complete or partial response on the clinical assessment (specificity 50%; CI 19–81%).

There was agreement between MRI findings and the pathology findings in 56 out of 69 lesions (81%) (Table 3). The imaging findings underestimated the effect of chemotherapy in one case where disease appeared to have progressed on CE MRI, but where pathology results indicated no response. In a further 12 cases, MRI overestimated the response to therapy. The sensitivity (100%; CI 94–100%) and specificity (80%; CI 44–97%) of MRI for identifying complete or partially responsive tumours were higher than for the clinical assessment alone. The absolute agreement between MRI and pathology findings was marginally higher than the agreement between clinical assessment and pathology findings (81 vs 68%; $P=0.09$).

Diagnostic confidence and therapeutic impact after MRI

In the majority of cases, the 2nd MRI study was judged to increase diagnostic knowledge (>70% see Table 4). However, in a substantial proportion of cases (17% see Table 4) the 2nd MRI was judged to have reduced diagnostic knowledge. These 12 cases have been examined in greater detail – see Table 5A.

In the majority of patients the 2nd MRI study did not change the treatment plan but did increase diagnostic confidence (52%, Table 4). This confidence was clearly valued by clinicians. Again, in a substantial minority of cases (20%, Table 4) the MRI findings
were thought to be confusing or incorrect. In the majority of these cases, the problem stemmed from the MRI study underestimating the amount of residual tumour by suggesting a complete response when, in fact, there were residual invasive tumour cells or DCIS. These cases have been examined more closely and the findings set out in Table 5a. In a further 19% of cases (Table 4) the 2nd MRI either did, or had the potential to alter the treatment plan. These cases have been summarised in Table 5b. The reasons included a contralateral nonpalpable tumour, complete response that could have allowed breast conservation, demonstration of nonresponse, demonstration of a single focus where multifocality was suspected. The mean diagnostic confidence for all patients significantly increased from 5.8 pre-2nd MRI to 7.9 post-2nd MRI (P < 0.01; Table 4).

Surgical findings indicated that WLE would not have been possible following neoadjuvant chemotherapy in 47 out of 69 lesions (Table 4). In 35 out of these 47 cases, both conventional assessment and MRI correctly predicted the need for mastectomy. The remaining 12 cases were evenly split between cases where conventional assessment incorrectly predicted WLE was possible (4), cases where MRI incorrectly predicted WLE was possible (4), and cases were both presurgery evaluations incorrectly suggested that breast-conserving surgery was possible (4).

In all, 10 out of the 22 lesions (46%) where surgical findings demonstrated that WLE would have been feasible were correctly identified by both conventional assessment and MRI (Table 4). In five cases, MRI correctly contradicted the conventional assessment by suggesting that WLE was possible. However, in the remaining seven cases MRI suggested that WLE was possible when surgical findings indicated that it was not. The overall agreement with surgical findings regarding the feasibility of WLE was 74% (51 out of 69) for conventional assessment and 78% (54 out of 69) for MRI (P = 0.44). Cases where the 2nd MRI study disagreed with surgical findings regarding the possibility of WLE are summarised in more detail in Table 6.

DISCUSSION

To achieve the objectives of neoadjuvant timing in the use of chemotherapy, it is necessary to understand the available preoperative methods of assessing response. Assessment of response to permit conservation surgery and to judge the effectiveness of the chemotherapy and the biological responsiveness of tumours are both important. Clinical methods have been unreliable (El-Didi et al, 2000), and there are clear limitations in the power of mammography (Huber et al, 2000) or ultrasound (for this purpose, ultrasound is widely used, but very poorly researched) to provide accurate information. CE breast MRI has attractions for this purpose, since it has high sensitivity for
invasive tumour and is dependent on increase in vessel density and abnormal diffusion properties of tumour blood vessels for its mode of visualisation. It is essentially a tomographic technique, able to give attractive 3D images that can aid surgeons in planning their operative procedures (Figure 3).

There have been a number of studies testing CE breast MRI for this purpose, which have been summarised in Table 1. The conclusions of these studies have varied considerably and several of them have an up-beat approach that may give surgeons and oncologists false expectations of the capacity of the technique to be reliable in all the aspects needed for planning treatment. Our study is from a single multidisciplinary team in two hospitals, with clinical work of several experts in each discipline. We have used the contemporary reports and notes of the cases to assess the

| Case | Age, years | Clinical size, mm at start of treatment | Grade | ER | Quadrants involved | Spaced out cells DCIS present | cR | mR | pR | Comments |
|------|------------|----------------------------------------|-------|----|-------------------|-------------------------------|----|----|----|----------|
| (A)  |            |                                        |       |    |                    |                               |    |    |    |          |
| 1a, b| 46         | 50                                     | 1     | +  | Multifoc          | No                            | No | 1  | 2  | Underestimates residual tumour |
| 2a, b| 51         | Unmeasurable                           | 2     | +  | Unifocal          | No                            | No | 2  | 2  | Underestimates residual tumour |
| 6a, b| 47         | 60+                                    | 3     | +  | Subareolar        | No                            | Yes| 2  | 2  | Underestimates residual tumour ILC |
| 8b   | 41         | Unmeasurable                           | 3     |   | Diffuse           | No                            | Yes| 1  | 2  | Underestimates residual tumour |
| 13b  | 54         | 40                                     | 2     | +  | Unifocal          | No                            | Yes| 1  | 2  | Underestimates residual tumour |
| 25b  | 39         | 60+                                    | 3     |   | Diffuse           | No                            | Yes| 1  | 3  | Underestimates residual tumour |
| 29b  | 34         | 60+                                    | 3     |   | Subareolar        | No                            | Yes| 1  | 1  | No contrast – failed injection CR report |
| 30b  | 49         | 60+                                    | 2     | +  | Unifocal          | Yes                           | Yes| 2  | 1  | Underestimates residual tumour DCIS |
| 31b  | 51         | 60+                                    | 2     | +  | Multifoc          | No                            | Yes| 2  | 2  | Underestimates residual tumour |
| 32b  | 40         | 45                                     | 3     |   | Multifoc          | No                            | No | 2  | 2  | Underestimates residual tumour |
| 34b  | 43         | 30                                     | 3     |   | Unifocal          | No                            | Yes| 1  | 1  | Underestimates residual tumour Residual DCIS |
| 34b  | 34         | 60+                                    | 2     | +  | Multifoc          | No                            | Yes| 1  | 2  | Underestimates residual tumour ILC |
| 34b  | 47         | 60+                                    | 1     | +  | Multifoc          | No                            | Yes| 2  | 1  | Underestimates residual tumour |
| 63b  | 60         | Unmeasurable                           | 3     |   | Diffuse           | No                            | Yes| 2  | 3  | Underestimates residual tumour |
| 66b  | 47         | 45                                     | 3     |   | Subareolar        | No                            | Yes| 2  | 2  | Underestimates residual tumour |
| 78b  | 28         | 60+                                    | 2     |   | Unifocal          | Yes                           | No | 2  | 1  | Underestimates residual tumour |
| (B)  |            |                                        |       |    |                    |                               |    |    |    |          |
| 7    | 52         | Unmeasurable                           | 2     |   | Unifocal          | Yes                           | Yes| 1  | 2  | Could have endorsed WLE |
| 14   | 49         | 40                                     | 3     |   | Unifocal          | No                            | Yes| 2  | 2  | Better estimate of residual tumour |
| 21   | 47         | 60+                                    | 2     |   | Unifocal          | No                            | Yes| 3  | 2  | Could have endorsed WLE |
| 34   | 43         | 30                                     | 2     |   | Unifocal          | No                            | Yes| 2  | 1  | Could have endorsed WLE DCIS only |
| 37   | 57         | 54                                     | 1     | +  | Multifoc          | No                            | No | 1  | 1  | Could have endorsed WLE |
| 44   | 56         | 40                                     | 3     |   | Unifocal          | No                            | No | 1  | 2  | Confirmed residual tumour |
| 52   | 43         | 50                                     | 2     | +  | Subareolar        | No                            | Yes| 2  | 3  | Shown nonresponder | Skin thickening |
| 55   | 43         | 30                                     | 3     | +  | Unifocal          | No                            | Yes| 1  | 1  | Could have endorsed WLE |
| 59   | 52         | 40                                     | 3     | +  | Unifocal          | Yes                           | Yes| 1  | 1  | Could have endorsed WLE |
| 62bilat | 49        | 60+                                    | 2     | +  | Diffuse           | No                            | Yes| 2  | 2  | Contralateral tumour shown |
| 67   | 33         | 32                                     | 3     | +  | Multifoc          | No                            | No | 2  | 2  | Did not confirm multifocality suspected on ultrasound. Could have endorsed WLE |
| 71   | 37         | 55                                     | 2     |   | Subareol          | No                            | No | 1  | 1  | Could have endorsed WLE |

Grade Bloom & Richardson grades 1, 2 & 3 (Elston and Ellis, 1991) oestrogen receptor = ER positive (+), negative (–). Ductal carcinoma in situ = DCIS, invasive lobular carcinoma = ILC. Quadrants = unifocal, multifocal, diffuse, subareolar position. cR = clinical response, mR = MRI response, pR = pathological response 1 = complete response, 2 = partial response, 3 = static disease, 4 = disease progression. Wide local excision WLE. *Cases in which the 2nd MRI study reduced diagnostic knowledge n = 12. **Cases in which the MRI findings were thought to be confusing or incorrect n = 14.
impact, real or potential, on clinical care provided by a CE MRI study at the start and end of neoadjuvant chemotherapy. We have used the histopathological response as the gold standard and since most of our patients had mastectomy, we have the potential to judge whether WLE could have been achieved. By using the full multidisciplinary team in evaluating each case, we hope to have avoided the bias that might ensue from radiologists alone assessing their value. Diagnostic and therapeutic impact methodology has been used for assessment of other new diagnostic procedures (Shuman et al, 1985b) including MRI (Hollingworth et al, 2000) for other purposes. We have moulded this methodology to apply to the present clinical scenario, for example by including an extra category – ‘MRI results could have caused a measurable change in the original plan’.

A few patients had their MRI study after the first cycle of chemotherapy was given, but these cases have not been excluded because the end points in this analysis relate to the presurgical MRI findings. The range of length between the 2nd MRI and surgery is determined by difficulties with scheduling of both MRI studies and reconstructive surgery in our hospital. The range of times between 1st and 2nd MRI studies is determined by the different regimens and patient episodes such as depression of white count which may have delayed cycles of chemotherapy. Whether these variations are important to the end points of this analysis is a matter of speculation, but they relate to pressures that would apply to real-life use of MRI out of the study setting. Our patients received a variety of chemotherapeutic regimens but this should make our CE MRI evaluation generally applicable to actual clinical practice where chemotherapy choices vary. The analysis of the MRI study is not affected by the different chemotherapy regimens.

We have made the main comparison of sensitivity and specificity between clinical examination coupled with conventional breast imaging (mammography and breast ultrasound) and CE MRI. CE MRI had both better sensitivity and specificity than clinical measures in conjunction with conventional breast imaging for detecting complete or partial response. The sensitivity of conventional methods is high, but there are serious limitations in specificity. In all, 50% (five out of 10) of patients with no response on pathology were incorrectly classified as partial or complete responders on clinical assessment, mammography and ultrasound. The specificity of CE MRI was higher (eight out of 10), but, due to the small number of patients who did not respond to chemotherapy, this result did not reach statistical significance. Furthermore, CE MRI still gives erroneous results in a number of cases. This is demonstrated in Table 5a where it is seen that the commonest limitation of the value of CE MRI is to underestimate residual tumour when compared with final pathology. Several authors (Gilles et al, 1994; Drew et al, 2001; Weatherall et al, 2001; Partridge et al, 2002) have found very good prediction of residual tumour by breast MRI, whereas Rieber in two studies had findings similar to ours (Rieber et al, 1997, 2002). We wish to seek the explanation for this in greater detail, and will do so by a separate publication studying the dynamic analysis compared with detailed pathology. Many of the cases with small amounts of residual tumour have islands of invasive cells or of DCIS spread across the area previously occupied by the treated tumour. The contrast uptake of the CE MRI did not distinguish this minimal residual disease (MRD) from any background uptake within the glandular tissue of the breast. The chemotherapeutic agents attenuate the contrast uptake characteristics, rendering the curves on some occasions benign in appearance. Clinical assessment of response is based on the size of the palpable mass and the clinical evidence of reduction in lymphoedema or in lymph node size. CE MRI was shown here to be better than these clinical findings even when they are backed up by ultrasound.

The cases in which CE MRI gave added value are shown in Table 5b. These are demonstration of unsuspected contralateral tumour, endorsement of a decision for WLE, better demonstration of nonresponse by showing skin thickening, better estimate of residual tumour and proof of unifocal disease. In Table 6, we have scrutinised the performance of CE MRI in predicting the possibility of WLE after neoadjuvant chemotherapy. It was a surprise to us that so few cases were suitable for breast-conserving surgery from the start, and we do not have the power here to show whether MRI was helpful in a significant way when only the cases which from the outset could have been suitable are considered.

It has been shown that pathological response is predictive of survival (Kuerer et al, 1999; Amat et al, 2002; Chollet et al, 2002) and that outcome is predicted also by tumour grade and nodal involvement. It would be useful to know whether complete response on MRI is predictive of outcome, even when

### Table 6 Cases where the 2nd MRI study disagreed with surgical findings regarding the possibility of WLE

| Case no. | Age in years | Radiological size, mm | Core biopsy grade | ER | Initial assessment allows WLE | MRI2 indicates WLE possible | Final pathology indicates WLE possible | DCIS present in final pathology | Spaced out cells present in final pathology | Comments |
|----------|--------------|-----------------------|-------------------|----|-------------------------------|-----------------------------|-----------------------------------|-----------------------------------|------------------------------------------|----------|
| 1        | 46           | 60                    | 1                 | +  | No                            | No                          | Yes                               | No                                | No                        | 15 mm unifocal                        |
| 4        | 38           | 70                    | 2                 | +  | No                            | No                          | Yes                               | No                                | No                        | 22 mm unifocal                        |
| 14       | 49           | 34                    | 3                 | —  | Yes                           | No                          | No                                | No                                | No                        | 31 mm unifocal                        |
| 22       | 44           | 19                    | 3                 | —  | No                            | No                          | Yes                               | No                                | No                        | No residual tumour                   |
| 27       | 45           | 35                    | 3                 | +  | No                            | No                          | Yes                               | No                                | No                        | 25 mm unifocal                        |
| 31       | 51           | 58                    | 2                 | +  | No                            | Yes                         | Yes                               | No                                | No                        | No residual tumour                   |
| 62       | 49           | 39                    | 2                 | +  | No                            | No                          | Yes                               | No                                | No                        | 20 mm unifocal                        |
| 2        | 51           | 55                    | 2                 | +  | Yes                           | Yes                         | No                                | No                                | No                        | Multifocal                           |
| 6        | 47           | 53                    | 3                 | +  | Yes                           | No                          | Yes                               | No                                | No                        | Widespread residual                 |
| 10       | 41           | 51                    | 2                 | +  | No                            | Yes                         | No                                | No                                | No                        | 41 mm residual                       |
| 11       | 60           | 38                    | 3                 | —  | No                            | Yes                         | No                                | No                                | No                        | Mass+scattered foci                  |
| 13       | 54           | 34                    | 2                 | +  | Yes                           | Yes                         | No                                | No                                | No                        | Multifocal                           |
| 29       | 34           | 100                   | 3                 | —  | No                            | Yes                         | Yes                               | No                                | No                        | CR but inflammatory at outset         |
| 38       | 43           | 60                    | 2                 | +  | No                            | Yes                         | No                                | No                                | No                        | Widespread foci                      |
| 61       | 47           | 50                    | 1                 | +  | No                            | Yes                         | Yes                               | No                                | No                        | Multifocal                           |

### Table 5a

| Case no. | Age in years | Radiological size, mm | Core biopsy grade | ER | Initial assessment allows WLE | MRI2 indicates WLE possible | Final pathology indicates WLE possible | DCIS present in final pathology | Spaced out cells present in final pathology | Comments |
|----------|--------------|-----------------------|-------------------|----|-------------------------------|-----------------------------|-----------------------------------|-----------------------------------|------------------------------------------|----------|
| 13       | 54           | 34                    | 2                 | +  | Yes                           | Yes                         | No                                | No                                | No                        | Multifocal                           |
| 29       | 34           | 100                   | 3                 | —  | No                            | Yes                         | Yes                               | No                                | No                        | Multifocal                           |

### Table 5b

| Case no. | Age in years | Radiological size, mm | Core biopsy grade | ER | Initial assessment allows WLE | MRI2 indicates WLE possible | Final pathology indicates WLE possible | DCIS present in final pathology | Spaced out cells present in final pathology | Comments |
|----------|--------------|-----------------------|-------------------|----|-------------------------------|-----------------------------|-----------------------------------|-----------------------------------|------------------------------------------|----------|
| 13       | 54           | 34                    | 2                 | +  | Yes                           | Yes                         | No                                | No                                | No                        | Multifocal                           |
| 29       | 34           | 100                   | 3                 | —  | No                            | Yes                         | Yes                               | No                                | No                        | Multifocal                           |
it fails to identify isolated foci of malignant cells or residual DCIS. If response on MRI were to be predictive of survival or disease-free survival, then this would be a test that gave predictive information prior to surgery and full pathology. Many more cases than the present study would be required to answer this question.

Within this series, the grade 3 and ER negative tumours tended to have a slightly better probability of complete response than the lower grade, ER-positive tumours, however neither of these differences reached statistical significance. A factor to consider here is the under staging that occurs with core biopsy, where 70% of cases have been shown to be upstaged between core biopsy and definitive surgical histology (Harris et al, 2003). After chemotherapy, we do not have satisfactory grading, and so the grades discussed here are all from core biopsy. The response rate that has been recorded cannot be appraised since it is dependent on the case selection for use of neoadjuvant chemotherapy and the different regimens in use, neither of which have been made standard in this study. We are looking at the effectiveness of CE MRI as a tool for monitoring, not at response rates in themselves. Our study shows that our MRI findings had very serious limitations in predicting the possibility of undertaking wide local excision after neoadjuvant chemotherapy, and that the shortfall was in both directions and not predictable from any particular characteristics of the cases. This study also shows how inaccurate clinical assessment of tumour response is, even when enhanced by ultrasound or mammography.

Even though our study has shown definite limitations to the reliability of MRI, the clinical team found that it provided valuable additional information. The problems with CE MRI are sufficient to continue to seek improved methods of assessing tumour response. Functional MRI, PET scanning and molecular imaging are emerging techniques that may offer more accurate information.

CONCLUSION

Our study suggests that CE MRI may have better sensitivity and specificity for predicting complete or partial response than clinical examination with conventional imaging. Moreover, the absolute agreement between MRI and pathology findings was marginally higher than the agreement between clinical assessment and pathology findings. CE breast MRI is robust in predicting nonresponse and disease progression. Our clinicians valued the visual images that are obtained which show tumour response. They feel that they give additional information – probably by giving a better measure of size and providing images that help them to understand the permeative properties of the tumour.

In our series, MRI persistently underestimates residual malignancy where the tumour is not a single mass, and this may be a problem in some cases when it is used to predict the possibility of WLE.

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Appendix A
See Tables A1 and A2.

Table A1  Chemotherapy regimens in use

| Regimen                        | n   |
|-------------------------------|-----|
| Doxorubicin/cyclophosphamide six cycles | 41  |
| Doxorubicin/docetaxel six cycles | 21  |
| Epirubicin four cycles then CMF four cycles | 2   |
| Epirubicin four cycles         | 2   |
| Docetaxel/Herceptin            | 1   |
| Total                          | 67  |

Clinical trials (1997–2003): Anglo-Celtic 2 Study: doxorubicin/cyclophosphamide vs doxorubicin/docetaxel six cycles (Evans et al, 2002). _ Docetaxel/Herceptin expanded access phase 4 assessment (Howell et al, 2002). Standard off study treatment: doxorubicin/cyclophosphamide six cycles.
### Table A2  Cases included in the study

| Case | Age | Size | Infl | Gr | ER | PgR | Hist | Trial | cR | mR | pR |
|------|-----|------|------|----|----|-----|------|-------|----|----|----|
| 01   | 46  | 60   | n    | 1  | +  | IDC | AC2  | cCR   | mPR| pPR|
| 02   | 51  | 55   | n    | 2  | +  | IDC | AC2  | cPR   | mPR| pPR|
| 03   | 51  | 65   | n    | 3  | +  | IDC | AC2  | cCR   | mPR| pPR|
| 04   | 38  | 70   | n    | 2  | +  | IDC | AC2  | cPR   | mPR| pPR|
| 05   | 55  | 45   | n    | 1  | +  | IDC | AC2  | cPR   | mPR| pPR|
| 06   | 47  | 53   | n    | 3  | +  | IDC | AC2  | cPR   | mPR| pPR|
| 07   | 52  | 70   | n    | 2  | +  | 0   | IDC | AC2  | cCR   | mPR| pPR|
| 08   | 31  | 48   | n    | 2+ | +  | IDC | AC2  | cPR   | mPR| pPR|
| 09   | 41  | 70   | y    | 3  | +  | IDC | AC2  | cCR   | mPR| pPR|
| 10   | 41  | 51   | n    | 2  | +  | IDC | AC2  | cPR   | mPR| pPR|
| 11   | 56  | 38   | n    | 3  | +  | IDC | AC2  | cPR   | mPR| pPR|
| 12   | 61  | 65   | n    | 3  | +  | IDC | AC2  | cPR   | mPR| pPR|
| 13   | 54  | 34   | n    | 2  | +  | IDC | AC2  | cCR   | mPR| pPR|
| 14   | 34  | 69   | y    | 2  | +  | IDC | AC2  | cCR   | mPR| pPR|
| 15   | 49  | 34   | n    | 3  | +  | IDC | AC2  | cPR   | mPR| pPR|
| 16   | 34  | 53   | y    | 3  | +  | 0   | IDC | AC2  | cPR   | mPR| pPR|
| 17   | 32  | 32   | n    | 3  | +  | IDC | AC2  | cCR   | mPR| pPR|
| 18   | 35  | 50   | n    | 3  | +  | IDC | AC2  | cPR   | mPR| pPR|
| 19   | 35  | 50   | n    | 3  | +  | IDC | AC2  | cPR   | mPR| pPR|
| 20   | 40  | 34   | n    | 2  | +  | IDC | AC2  | cPR   | mPR| pPR|
| 21   | 30  | 34   | y    | 2  | +  | IDC | AC2  | cCR   | mPR| pPR|
| 22   | 30  | 34   | n    | 3  | +  | IDC | AC2  | cPR   | mPR| pPR|
| 23   | 45  | 19   | y    | 3  | +  | IDC | AC2  | cPR   | mPR| pPR|
| 24   | 49  | 40   | n    | 3  | +  | IDC | AC2  | cPR   | mPR| pPR|
| 25   | 50  | 46   | n    | 3  | +  | IDC | AC2  | cPR   | mPR| pPR|
| 26   | 49  | 39   | n    | 3  | +  | IDC | AC2  | cPR   | mPR| pPR|
| 27   | 43  | 35   | n    | 3  | +  | IDC | AC2  | cCR   | mPR| pPR|
| 28   | 36  | 35   | n    | 3  | +  | IDC | AC2  | cCR   | mPR| pPR|
| 29   | 34  | 100  | y    | 3  | +  | IDC | AC2  | cCR   | mPR| pPR|
| 30   | 49  | 54   | n    | 2  | +  | IDC | AC2  | cPR   | mPR| pPR|
| 31   | 51  | 58   | n    | 2  | +  | IDC | AC2  | cPR   | mPR| pPR|
| 32   | 40  | 44   | n    | 3  | +  | IDC | AC2  | cPR   | mPR| pPR|
| 33   | 40  | 44   | y    | 3  | +  | IDC | AC2  | cPR   | mPR| pPR|
| 34   | 43  | 30   | n    | 3  | +  | 0   | IDC | AC2  | cCR   | mPR| pPR|
| 35   | 59  | 66   | n    | 3  | +  | IDC | AC2  | cPR   | mPR| pPR|
| 36   | 49  | 49   | n    | 2  | +  | IDC | AC2  | cPR   | mPR| pPR|
| 37   | 51  | 50   | n    | 3  | +  | IDC | AC2  | cCR   | mPR| pPR|
| 38   | 43  | 100  | y    | 2  | +  | IDC | AC2  | cCR   | mPR| pPR|

**Key to column headings:**
- **Size:** radiological (mammography and ultrasound) size at diagnosis.
- **Infl:** inflammatory changes present, grade = Bloom and Richardson nuclear grade (Elston and Ellis, 1991) on core biopsy.
- **Gr:** grade (Bloom and Richardson nuclear grade) on core biopsy.
- **ER:** oestrogen receptor status on core biopsy.
- **PgR:** progesterone receptor status.
- **Hist:** histology, IDC = Invasive ductal carcinoma, ILC = Invasive lobular carcinoma.
- **Trial:** chemo-neoadjuvant chemotherapy regimen.
- **cR:** clinical response.
- **mR:** MRI response.
- **pR:** pathological response.
- **CR:** complete response.
- **PR:** partial response.
- **NR:** nonresponder.
- **DP:** disease progression.

IEC = Anglo-Celtic 2 trial.
Appendix B

See Table B1.

Table B1  MR pulse sequence protocol for a breast MR examination in use for this study

| Sequence type                                                                 | Repetition time TR (ms) | Echo time TE (MS) | Flip angle (°) | Matrix size       | Number of slices | Field of view (FOV) (MM) | Pixel size (mm x mm) | Number of averages | Acquisition time (mins) |
|--------------------------------------------------------------------------------|-------------------------|-------------------|----------------|-------------------|------------------|------------------------|----------------------|--------------------|------------------------|
| Precontrast, high-resolution 3D T1-weighted                                   | 8.9                     | 4.2               | 35             | 512 × 384         | 60               | 340                    | 0.66 × 0.89          | 1                  | 2                      |
| Dynamic contrast enhanced, 3D T1-weighted                                    | 8.8                     | 4.2               | 35             | 256 × 256b        | 480              | 340                    | 1.33 × 1.33b         | 1                  | 8 × 1.10               |
| Postcontrast, high-resolution 3D T1-weighted with fat suppression            | 18.9                    | 4.2               | 35             | 512 × 384         | 60               | 340                    | 0.66 × 0.89          | 1                  | 5                      |

"Contrast injection was 0.16 mmol kg⁻¹ body weight with an injection speed of 3 ml s⁻¹, given after the second precontrast dynamic sequence." The matrix size is optimised to ensure that the acquisition time of each 3D image is 70 s.