An objective biochemical assessment of therapeutic response in metastatic breast cancer: a study with external review of clinical data

M.R. Williams1,*, A. Turkes2, D. Pearson1, K. Griffiths1 & R.W. Blamey1

1Nottingham City Hospital, Nottingham NE5 1PB and 2Tenovus Institute for Cancer Research, University Hospital of Wales, The Heath, Cardiff, UK.

Summary
A series of tumour related markers have been examined in 179 patients receiving primary endocrine therapy for metastatic breast cancer. Significant correlations between therapeutic response (UICC criteria after 6 months of treatment) and appropriate alterations in serum concentrations of carcinoembryonic antigen, ferritin, c-reactive protein, oosromocoid and the erythrocyte sedimentation rate, have been observed when changes in these markers were examined only at high serum concentrations. By combining these five markers a ‘therapeutic index’ of response has been devised which can be employed at an early stage of treatment in more than 90% of patients, giving an overall sensitivity specificity of 90%/78% for therapeutic response or disease stabilisation over a 6-month period. The design of an objective measurement of response, which is easy to perform, has the potential to replace the existing, largely subjective, UICC criteria for retrospective judgement of response, and may also be used to direct systemic endocrine therapy.

Many systemic treatments are now available for the palliation of disseminated breast cancer. In the absence of an immediate threat to life, endocrine therapy remains the initial treatment of choice, due to its relative lack of toxicity. An early and accurate assessment of response to endocrine therapy would be of value, in order to allow a timely alteration of treatment in unresponsive patients. At present, such a decision is based upon ill defined clinical criteria, although the International Union Against Cancer (UICC) have suggested guidelines for the retrospective judgement of response in clinical trials (Hayward et al., 1977).

Previous reports have explored the role of tumour markers in an attempt to measure therapeutic response objectively throughout treatment (Chu & Nemoto, 1973; Steward et al., 1974; Borthwick et al., 1977; Tormey et al., 1977a, b; Haagenson et al., 1978; Lamerz et al., 1980; Haagenson et al., 1980; Lee, 1983; Krieger et al., 1983; Coombes et al., 1983; Campbell et al., 1983; Heim et al., 1984; Hortobagyi et al., 1984). Few studies have examined the potential of a combination of markers recorded simultaneously, despite indications that this approach might prove rewarding (Coombes et al., 1977; Woo et al., 1978; Cowen et al., 1978; Cove et al., 1979; Caffer & Brandau, 1983).

This study describes an assessment of response to endocrine therapy which employs a combination of five tumour markers, and compares this biochemically based assessment with that achieved using the accepted clinical criteria after 6 months of treatment. The majority of tumour markers for advanced breast cancer have lacked specificity for the determination of response, as they have been employed at low concentrations which are frequently found in tumour-free women. The present study attempts to correct this by utilising markers only when concentrations exceed predefined high levels (above the ninetieth centile for control populations).

Patients and methods

One hundred and seventy-nine patients who presented to our clinic over a 2-year period were studied. All had advanced recurrent breast cancer and all received endocrine therapy as initial treatment for overt metastases. Treatment was prescribed according to several protocols and consisted of tamoxifen (Nolvadex, 20 mg b.d.) or megestrol acetate (Megace, 160 mg b.d.) in post-menopausal patients, and oophorectomy or a luteinising hormone-releasing hormone agonist (Zoladex, IC 118630) in premenopausal patients. Menopausal status was confirmed using luteinising hormone and follicle stimulating hormone levels in women who had previously undergone hysterectomy, or who presented within 5 years after natural cessation of menstruation.

The oestrogen receptor status (ER) of primary tumour tissue had been documented in 109 patients; 51% were ER positive (greater than 5 fmol mg⁻¹ cytosol protein). ER estimations were performed at the Tenovus Institute, Cardiff, using the dextran coated charcoal method (Nicholson et al., 1981).

All patients were staged before treatment in an advanced breast cancer clinic. All but six patients had received no previous endocrine treatment for overt metastases. Additional symptomatic treatments were prescribed throughout the study which included systemic analgesics, steroids and/or bronchodilators for lung metastases, and radiotherapy for skin ulceration or localised bone pain. Patients unassessable to hormonal therapy either by the type of presenting disease or the addition of local treatments were not included in the study. Sixteen patients were excluded from analysis as sequential serum samples were not available because of early death. The patients studied, therefore, represented the great majority of those attending one unit over a 2-year period, who received endocrine therapy for assessable metastatic disease.

Initial staging included a full clinical examination, documentation of the Karnofsky performance status (Karnofsky & Burchenal, 1948) and accurate measurement of local and regional disease, with photography when considered helpful for later evaluation of response. Radiological assessment consisted of lateral and antero-posterior views of the skull, chest, dorso-lumbar spine, pelvis and upper femora (limited skeletal survey). Painful sites of disease at other locations were X-rayed when clinically indicated. Isotope bone, liver and brain scans were requested only if clinically indicated.

Haematological assessment consisted of measurement of a full blood count, urea and electrolytes, serum calcium and albumin, the erythrocyte sedimentation rate and liver function tests (gamma-glutamyl transaminase, alanine transferase, alkaline phosphatase and bilirubin). In addition, several other ‘tumour related markers’ were assessed sequentially throughout active treatment (Table I). These included two oncofetal proteins, serum carcinoembryonic antigen (CEA) and ferritin; two acute phase proteins, c-reactive protein (CRP) and oosromocoid; and the hormone beta human chorionic gonadotrophin (B-HCG). Calcium excretion (CAE) and the hydroxyproline creatinine ratio (OHP/CR) were
Table 1 Tumour markers studied

| Serum (all patients)                      | Upper limit of normal adopted for study |
|-------------------------------------------|----------------------------------------|
| Carcinoembryonic antigen<sup>a</sup>      | 6 ng ml<sup>-1</sup>                   |
| Ferritin<sup>b</sup>                      | 220 µg l<sup>-1</sup>                  |
| C-Reactive protein<sup>c</sup>            | 10 mg l<sup>-1</sup>                   |
| Orosomucoid<sup>d</sup>                   | 1.2 g l<sup>-1</sup>                   |
| Erythrocyte sedimentation<sup>e</sup>     | 20 mm h<sup>-1</sup>                   |
| Gamma-glutamyl transpeptidase             | 50 U l<sup>-1</sup>                    |
| Alanine aminotransferase                  | 50 U l<sup>-1</sup>                    |
| Alkaline phosphatase                      | 300 U l<sup>-1</sup>                   |
| Beta human chorionic gonadotrophin        |                                        |
| Urine (bone metastases only)              |                                        |
| Hydroxyproline/creatinine ratio           |                                        |
| Calcium excretion                         |                                        |

<sup>a</sup>Tumour markers combined to form therapeutic index for response.

recorded throughout treatment in the urine of a subgroup of patients presenting with bone metastases (n = 152 for CAE, n = 94 for OHP/CR).

Controls

Marker levels were established in women with benign breast disease attending a diagnostic breast clinic (controls, n = 55 for analysis of CEA and ferritin; n = 25 for analysis of acute phase proteins). These women had a mean age of 49 years (range 28–85) and presented with histologically confirmed benign breast lumps, breast pain or cysts.

The pre-treatment concentration of each marker in stage IV disease was also compared with that found in 87 patients with untreated stage III breast cancer (see Results).

Follow-up

Patients were reviewed at 1–3 monthly intervals. Those with bone metastases provided a fasting, early morning urine sample for the measurement of CAE and the OHP/CR ratio. The Karnofsky performance status was recorded at each visit, venous blood was withdrawn for measurement of routine haematological parameters and additional serum was centrifuged, aliquoted and stored at −170°C for later evaluation of the remaining markers.

Limited skeletal radiology was repeated after 3–4 months and after 6 months of treatment. At the first clinical sign of objective progression patients were prescribed alternative treatments and therefore removed from study. All assessments of clinical progress were performed without referral to biochemical data, and both the initial documentation of disease status and subsequent assessments of clinical progress were performed by the same observer at each visit (M.R.W.).

Assessment of clinical response

UICC criteria have been strictly applied with the British Breast Group stipulation that any remission should be of at least 6 months' duration to classify as response (British Breast Group, 1974; Hayward et al., 1977). These criteria require a 50% reduction in measurable tumour or objective signs of response in evaluable, but non-measurable, sites of disease (e.g. lung or bone metastases).

Clinical response has been categorised as 'disease progression' (greater than 25% increase in the bidimensional product of measurable tumour or the development of new lesions), 'objective response' (greater than 50% reduction in the size of measurable tumour with no new lesions) or 'disease stabilisation' (no new lesions and any alteration in tumour size lying between these two extremes). In this study response of less than 6 months' duration has been classified as disease progression. External review of response was performed by Dr A. Howell, Christie Hospital, Manchester.

Assay techniques

Routine biochemical and haematological parameters were measured using standard techniques.

CEA was measured using a monoclonal radiometric assay (Tandem-R CEA, kindly provided by Hybritech UK Ltd). The intra- and inter-assay coefficients of variation were 4.6–7.6% and 6.9–7.2%, respectively.

Serum concentrations of ferritin were measured using a solid phase, two site radioimmunoassay. Standards were prepared in ferritin free serum and calibrated against a WHO ferritin reference preparation. Standards covered the range of 0–1000 µg l<sup>-1</sup>. The estimated intra- and inter-assay coefficients of variation, over the working range, were 3.7–5.9% and 4.6–6.0%, respectively.

Concentrations of the beta subunit of human chorionic gonadotrophin were measured by radioimmunoassay using an antiserum raised in rabbit against B-HCG. Standards were calibrated against the First International Reference Preparation (kindly provided by the National Institute of Biological Standards and Control, London) and were prepared to cover the range 0–640 IU l<sup>-1</sup>. The intra- and inter-assay coefficients of variation were 4.7–6.1% and 5.4–8.8%, respectively.

Serum concentrations of CRP were measured using a turbidimetric technique on a centrifugal fast analyser (Centrifichem Roche). Changes in optical density were monitored, using an on-line computer which calculated CRP concentrations (O'Callaghan et al., 1984).

Orosomucoid was measured by immuno-turbimetry using a centrifugal fast analyser (Centrifichem 400). Standards were prepared from a commercially available serum (Behring Standard Human Serum, ORDT 06/07). Between batch precision for the assay was 7% at a concentration of 0.65 g l<sup>-1</sup>.

Urinary OHP was estimated in duplicate using Hypronosticon Kits (Organon).

Analysis of results

Pre-treatment concentrations of each tumour marker were compared in individual patients with values recorded after 1–2, 3–4 and 5–7 months of treatment.

For each marker the correlation between alterations in concentration and clinical assessments of response was examined only in patients who presented with, or developed, concentrations exceeding predefined levels; the proportion of patients that this represented for each marker is shown in the results. The object of this was to examine markers in each individual only when concentrations were well above those found in the majority of women without advanced disease; concentrations above this level may then be assumed to be due to the tumour. Such concentrations were seen in only 2–8% of women in the control groups. The 'cut-off' chosen for analysis also depended upon the shape of the distribution plots for each marker.

Patients maintaining concentrations within the 'normal' range have been considered unassessable for the marker in question. Above this level concentrations were classified as increasing or decreasing when they altered by more than 1% during treatment.

For statistical analysis patients with rising marker concentrations were combined with those in whom concentrations remained stable. Patients with no change in clinical disease status over 6 months, were combined with those showing objective response. Statistical significance was assessed using the χ² test with Yates' correction.

Finally, results of five markers were combined to form a 'therapeutic index', in order to improve the discrimination of response after 3–4 months of treatment.

Results

Patient details and therapeutic response

One hundred and six patients presented with bone or lung metastases alone (n = 69 and n = 37, respectively). Thirty-
eight presented with both bone and lung metastases, and 35
presented with other sites of visceral involvement (mainly
hepatic). Twenty-three per cent of patients (41/179) were
found to be premenopausal at the presentation of advanced
disease.

The overall response rate was 26% (n = 47) using UICC
criteria after 6 months. In 17% (n = 31) disease remained
static for 6 months and in 56% (n = 101) disease progressed.
Disease either responded or remained static in 64% (36/56)
of ER positive patients and in 26% (14/53) of ER negative
patients.

Correlation between therapeutic response and alterations in
individual marker concentrations

In only five markers did appropriate alterations in concentra-
tion correlate with UICC response to a degree that could be
usefully employed in clinical practice (Table II). For the
remaining markers the correlation was weak and these latter
results are not discussed further in this report.

In total, 579 serum samples were obtained for the analysis
of tumour markers in patients with stage IV disease. These
included 179 samples obtained immediately before treatment,
132 obtained after 1 or 2 months (from 128 patients), 131
after 3 or 4 months (from 128 patients) and 137 obtained
after 5–7 months of treatment (from 114 patients).

Only the first recorded assay result has been considered for
analysis when these were duplicated in individual patients
during a single interval of treatment. Individual results for
the five markers, recorded simultaneously on the same serum
sample, were unavailable on several occasions; four pre-
treatment results and 10 follow-up results were not available
for ESR, three follow-up results for CRP, and one for both
orosomucoid and ferritin.

For each marker, the pre-treatment concentration in stage
IV disease was compared with that found both in disease-free
women attending a diagnostic breast clinic and in a series of
patients presenting with stage III disease, in order to establish
upper limits of normal.

Carcinoembryonic antigen Only one control patient present-
ed with a CEA concentration in excess of 6 ng ml\(^{-1}\),
compared with 18% (16/87) of those with stage III disease and
49% (88/179) of those presenting with stage IV disease
(Figure 1, median concentrations illustrated). Alterations in
CEA during treatment have been examined only when con-
centrations exceeded 6 ng ml\(^{-1}\).

Table II Alterations in individual serum marker concentrations during treatment versus therapeutic response

| Tumour marker | UICC response at 6 months | 1–2 months treatment | 3–4 months treatment | 5–7 months treatment |
|---------------|--------------------------|----------------------|----------------------|----------------------|
|               | Marker concentrations    |                      |                      |                      |
|               | pre-treatment versus     | Decrease | Stable | Increase | Decrease | Stable | Increase | Decrease | Stable | Increase |
| CEA Response  |                           | 15        | 2      | 3       | 22      | 1      | 1       | 22      | 0      | 3       |
| Static        |                           | 5         | 1      | 3       | 8       | 1      | 2       | 11      | 2      | 2       |
| Progression  |                           | 6         | 9      | 19      | 5       | 2      | 29      | 3       | 1      | 21      |
| Ferr. Response| (\(x^2 = 15.1\); l.d.f.)* | (\(x^2 = 33.8\); l.d.f.)* | (\(x^2 = 28.16\); l.d.f.)* |                      |                      |                      |
| Static        |                           | 11        | 4      | 1       | 13      | 2      | 0       | 18      | 1      | 2       |
| Progression  |                           | 5         | 0      | 1       | 5       | 1      | 1       | 8       | 0      | 0       |
| CRP Response  | (\(x^2 = 15.1\); l.d.f.)* | (\(x^2 = 23.4\); l.d.f.)* | (\(x^2 = 23.0\); l.d.f.)* |                      |                      |                      |
| Static        |                           | 21        | 0      | 1       | 22      | 0      | 0       | 24      | 0      | 1       |
| Progression  |                           | 9         | 0      | 2       | 13      | 0      | 3       | 14      | 1      | 3       |
| Oros. Response| (\(x^2 = 26.52\); l.d.f.)* | (\(x^2 = 30.82\); l.d.f.)* | (\(x^2 = 24.7\); l.d.f.)* |                      |                      |                      |
| Static        |                           | 26        | 2      | 2       | 25      | 1      | 2       | 29      | 3      | 1       |
| Progression  |                           | 11        | 1      | 0       | 13      | 1      | 3       | 15      | 1      | 2       |
| ESR Response  | (\(x^2 = 23.87\); l.d.f.)* | (\(x^2 = 26.4\); l.d.f.)* | (\(x^2 = 22.6\); l.d.f.)* |                      |                      |                      |
| Static        |                           | 23        | 2      | 5       | 27      | 0      | 6       | 31      | 1      | 4       |
| Progression  |                           | 11        | 3      | 1       | 15      | 0      | 1       | 15      | 2      | 3       |
| (\(x^2 = 12.24\); l.d.f.)* | (\(x^2 = 36.09\); l.d.f.)* | (\(x^2 = 15.21\); l.d.f.)* |                      |                      |                      |

Statistical significance assessed by combining 'response' with 'static', and 'stable' with 'increase' *\(p < 0.001\).
stage IV disease have been examined only above 220 μg l⁻¹.

One hundred and twenty-eight patients had ferritin levels recorded before and after 1 or 2 months of treatment. Fifty (39% of the total) were assessable using concentrations above 220 μg ml⁻¹. In one hundred and twenty-seven patients ferritin was measured before and after 3 or 4 months of treatment; 49 (39%) were biochemically assessable as above. In 114 patients ferritin was measured before and after 5–7 months of treatment, 48 (42%) were biochemically assessable.

Again, during each interval of treatment, a highly significant association existed between alterations in serum ferritin (above 220 μg ml⁻¹) and clinical assessments of response after 6 months (Table II: pre-treatment vs 1–2 months, \( \chi^2 = 15.1, 1 \) d.f.; vs 3–4 months, \( \chi^2 = 23.4, 1 \) d.f.; vs 5–7 months, \( \chi^2 = 23, 1 \) d.f.).

C-reactive protein Only one tumour-free patient presented with a CRP concentration in excess of 10 mg l⁻¹, compared with 13% (11/87) of those with stage III disease and 53% (94/179) of those presenting with stage IV disease (Figure 3). Thus, alterations in CRP have been examined only when concentrations exceeded 10 mg l⁻¹.

One hundred and twenty-seven patients with distant metastases had CRP concentrations measured before and after one or two months of treatment, of whom 70 patients (55% of the total) were assessable employing CRP concentrations in excess of 10 mg l⁻¹. One hundred and twenty-six patients had CRP concentrations measured before and after 3 or 4 months of treatment, 73 (58%) were assessable as above. In 114 patients CRP was measured before and after 5–7 months of treatment, 67 (59%) were biochemically assessable.

A highly significant association existed between therapeutic response (UICC) and alterations in CRP concentration during the three time intervals of treatment (Table II: pre-treatment vs 1–2 months, \( \chi^2 = 26.52, 1 \) d.f.; vs 3–4 months, \( \chi^2 = 30.82, 1 \) d.f.; vs 5–7 months, \( \chi^2 = 24.7, 1 \) d.f.).

Orosomucoid Only two control patients presented with an orosomucoid concentration in excess of 1.2 g l⁻¹, compared with 17% (15/87) of those with stage III disease and 61% (110/179) of those with stage IV disease (Figure 4). Alterations in concentration during treatment have been examined only when levels exceeded 1.2 g l⁻¹.

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months of treatment, 73 (58%) were assessable as above. In 114 patients CRP was measured before and after 5–7 months of treatment, 67 (59%) were biochemically assessable.

A highly significant association existed between alterations in marker concentrations during each interval of treatment, and therapeutic response assessed using UICC criteria after six months (Table II: pre-treatment vs 1–2 months, $\chi^2 = 23.87$, 1 d.f.; vs 3–4 months, $\chi^2 = 26.4$, 1 d.f.; vs 5–7 months, $\chi^2 = 22.6$, 1 d.f.)

**Erythrocyte sedimentation rate** The ESR was greater than 20 mm h$^{-1}$ in only two of 50 tumour-free patients attending a diagnostic breast clinic (not illustrated). In contrast, the ESR was greater than 20 mm h$^{-1}$ in 33% (26/79) of patients presenting with stage III disease and in 66% (115/175) of those presenting with stage IV disease (Figure 5). Alterations in the ESR have been examined only above 20 mm h$^{-1}$.

One hundred and twenty-one stage IV patients had ESR measurements before and after 1 or 2 months of treatment. Ninety-one (75% of the total) were assessable employing an ESR of greater than 20 mm h$^{-1}$. One hundred and twelve patients had the ESR measured before and after 3 or 4 months of treatment. 97 (87%) were assessable. In 111 patients the ESR was measured before and after 5–7 months of treatment, 91 (82%) were assessable employing an ESR greater than 20 mm h$^{-1}$.

Again, a highly significant association existed between alterations in the ESR during each interval of treatment and therapeutic response assessed after 6 months using UICC criteria (Table II: pre-treatment vs 1–2 months, $\chi^2 = 12.24$, 1 d.f.; vs 3–4 months, $\chi^2 = 36.09$, 1 d.f.; vs 5–7 months, $\chi^2 = 15.21$, 1 d.f.)

**Combining markers to assess therapeutic response after 3–4 months of treatment**

From the results of individuals tumour markers it can be seen that the accuracy with which each marker was able to assess response, was highest during the 3–4 month interval of treatment.

In 127 patients the results of all five markers were available during this interval of treatment. Patients were considered biochemically unassessable when the concentrations of all five markers remained below the minimum levels adopted for analysis. This occurred in 10 patients (8% of the total). Of the remaining 117 biochemically assessable patients, 37 responded to treatment (UICC criteria), in 20 disease remained static for 6 months and in 60 disease progressed.

A ‘therapeutic index’ has been devised by allocating points according to a rise or a fall in each marker concentration above defined levels, as shown in Table III.

Several factors were taken into consideration when constructing the index and different weighting was applied to appropriate alterations in each marker. Progressing disease was associated with high but stable marker concentrations (score +1). However, high, stable marker concentrations did not detect disease progression as accurately as increasing marker concentrations (score +2). The predictive value for disease progression of increasing concentrations of ferritin, the ESR and acute phase proteins (score +2) was higher than the predictive value for non-progressing disease of falling concentrations (score −1). Finally, appropriate alterations in CEA were equally effective to confirm progressing and non-progressing disease (score +2 and −2, respectively). When all five markers increased in concentration by more than 10%, a maximum index score of +10 was achieved. Conversely, when all five markers were abnormal at presentation and decreased by more than 10%, a minimum index score of −6 was achieved.

The distribution of index scores at 3–4 months is shown after subgrouping patients according to UICC assessments of response at 6 months (Figure 6). By employing a ‘cut-off’ lying

![Figure 5 ESR at presentation of stage III and IV disease: median concentrations illustrated.](image)

![Figure 6 Biochemical index scores after 3–4 months.](image)

**Table III Allocation of scores towards a biochemical index for response**

(values are index scores)

| Tumour marker concentrations during treatment | Remain within normal limits | Decrease by 10% | Remain stable | Increase by 10% |
|-----------------------------------------------|-----------------------------|-----------------|---------------|-----------------|
| CEA (>6 ng ml$^{-1}$)                          | 0                           | −2              | +1            | +2              |
| Ferritin (>220 µg l$^{-1}$)                    | 0                           | −1              | +1            | +2              |
| Orosomucoid (>1.2 g l$^{-1}$)                  | 0                           | −1              | +1            | +2              |
| CRP (>10 mg l$^{-1}$)                         | 0                           | −1              | +1            | +2              |
| ESR (>20 mm h$^{-1}$)                         | 0                           | −1              | +1            | +2              |
between index scores of 0 and +1, the sensitivity that a low
index score has for objective response (or disease stabilisation
for 6 months) is 89.5% (51/57), with a specificity of 78%
(47/60) (Table IV, group A; Figure 7).

When patients with 'marked biochemical change' are con-
sidered separately, (index scores greater than +2 or less than
−2), the sensitivity for response/stasis increases to 92% (34/37)
with a specificity of 100% (38/38) (Table IV, group B; Figure
8). Seventy-five patients (59% of the total) developed 'marked
biochemical' responses to treatment.

Discussion

Significant advances have been made in the treatment of
malignant diseases where specific markers are able to monitor
tumour burden. Two examples are the use of B-HCG assays in
choriocarcinoma (Begent & Bagshaw, 1982) and measurements
of B-HCG and alpha fetoprotein in patients receiving treat-
ment for testicular teratoma (Lange et al., 1976; Schultz et al.,
1978; Lange, 1982). These advances are dependent in part upon
the ability to assess changes in tumour bulk at an early stage,
which allows appropriate alteration of treatment.

This work seeks a marker of tumour burden for advanced
breast cancer so that similar principles can be applied. A single
marker with the desired specificity for response has not been
found, even when each marker was examined at high serum
concentrations. However, we have found that a combination of
five markers (at high serum concentration) is able accurately to
reflect therapeutic response of 6 months' duration. The five
assays are easily performed, and their combination is able to
discriminate between response groups at a relatively early stage
of treatment.

Each marker has been considered only at high serum
centresations, to limit interference from factors independent
of tumour burden. Many previous studies have examined
tumour markers in advanced breast cancer with the aim of
assessing response objectively during early treatment. The
majority of these reports can be criticised as the patients
studied were few in number, only single markers were
examined, marker concentrations remained within normal
limits in many patients analysed, and clinical response
criteria were often inadequate.

Several problems are encountered when employing clinical
criteria to assess response retrospectively. They require exten-
sive radiology and repeated measurements of tumour deposits,
making any assessment both difficult and time consuming. As a
result, clinical signs of response may be misinterpreted,
especially when assessments are made on plain X-rays.

Bone is the commonest site for distant metastases from
breast cancer and it is particularly in this situation that
discovery exists over the current criteria for response (Lip-
shitz & Hortobagyi, 1981; Coombes et al., 1983; Hortobagyi et
al., 1984). The development of sclerosis within lytic metastases
is regarded as a prerequisite for therapeutic response. This is
often slow to develop and may be absent despite unequivocal
evidence of response in other sites. Furthermore, the emergence
of blastic metastases may indicate either response (DeMartini
et al., 1983) or progression of disease. As radiological evidence
of response may take many months to appear, it is frequently of
limited value to guide treatment in practice.

In this study, patients with static disease (for a minimum
duration of 6 months) were combined with those showing
objective clinical evidence of response. Previous studies have
indicated that these patients fair as well as responders with
respect to survival (Howell et al., 1984; Williams et al., 1986).
Biochemical responses to treatment, assessed using this com-
bination of markers, were similar in patients showing partial
response or static disease. In contrast, patients with progressive
disease showed markedly different biochemical profiles. It is
submitted that the employment of clinical criteria alone is
inadequate to detect response in many patients found to have
stable disease on clinical grounds.

This biochemical index of response was devised retrospec-
tively. It will require prospective testing on a further series of

![Figure 8 Distribution of index scores in patients with 'marked biochemical change' after 3-4 months.](image_url)

![Figure 7 Distribution of index scores in all patients after 3-4 months.](image_url)

| Table IV Biochemical index scores after 3-4 months’ treatment |
|---------------------------------------------------------------|
| **UICC assessment** | **Group A** | **Group B** |
|                   | ≤0 | >0 | <−2 | >+2 |
| Response           | 35 | 2  | 23  | 0   |
| Static             | 16 | 4  | 11  | 3   |
| Progression        | 13 | 47 | 0   | 38  |
| *χ² = 51.54; 1 d.f.* |   |    |    |    |
| *χ² = 70.05; 1 d.f.* |   |    |    |    |

*χ² = 51.54; 1 d.f.*

*χ² = 70.05; 1 d.f.*

*χ² = 0.0012 (P<0.0001)
patients to assess its reproducibility. Where biochemical and clinical assessments of response are at variance, both criteria should be evaluated against other parameters such as survival. These studies are currently in progress and if the index is confirmed as correlating strongly with an assessment of response based on UICC criteria, then the biochemical assessment has several important advantages. These include its simplicity, objectivity and ability to assess response at an early stage and thus guide treatment. There are also economic advantages, as marker estimations amount to only a small proportion of the costs of limited skeletal surveys.

The UICC criteria were designed to be employed retrospectively and in the context of therapeutic trials only. They are often not adequate for ongoing assessments of therapeutic response during treatment. Provided this biochemical measurement of response proves reproducible, it might be possible to replace clinical criteria for response by this simple and truly objective alternative.

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