Introduction

Targeted therapy for breast cancer was actually used, albeit unwittingly, as early as the late 1800s. Beatson first surgically removed the ovaries of women to treat metastatic breast cancer in 1896 [1], while Schinzinger recommended ovarian irradiation as adjuvant therapy in 1889 [2]. Ovarian ablation and subsequently diethylstilbestrol were initially used to treat breast cancer without knowing the target to which they were directed. More than 50 years later Jensen and Jacobson discovered the oestrogen receptor (ER) [3], which was subsequently understood to be the target or partial target of these endocrine approaches.

It is now known that women whose tumours are ER-negative and progesterone receptor (PgR)-negative have less than a 5% chance of responding to endocrine therapies, while those whose tumours are ER-positive and/or PgR-positive have a chance of response of somewhere between 40 and 70%. It was believed that having both receptors positive improved the response rate or benefit from adjuvant endocrine therapy for many years, but is now felt that the ER serves as the major predictive factor. The PgR is felt to give some prognostic value but not to be associated with better response to endocrine therapy in the adjuvant or the metastatic setting [4] (R. Peto, personal communication).

For a number of years there was controversy about whether tamoxifen might be effective in ER-negative as well as ER-positive patients. Even the large Oxford Overview [5] did not initially display this clearly for some long time, and it was in fact from other studies with carefully controlled biomarkers that the role of ER in preditory effectiveness became more clear [6,7]. This experience provides lessons for the future, in that large meta-analyses in which markers may have been measured by a variety of methods may not be the best way of sorting out these matters. Even today there is ongoing controversy regarding the role of the ER and response to chemotherapy. It is generally felt that highly ER-positive patients may respond less well to chemotherapy. Certainly high ER and PgR levels have an inverse link to proliferative indices such as Ki-67, which tend to predict more strongly for response to chemotherapy.

We now have the 21-gene OncotypeDX recurrence score that may be used as a continuous variable both to predict benefit from tamoxifen and to predict response from those treated with tamoxifen [8-11]. It has also been shown that the OncotypeDX recurrence score correlates more strongly with outcome in tamoxifen-treated patients than the Adjuvant online program and that the OncotypeDX recurrence score is also predictive of local recurrence in tamoxifen-treated patients [12]. The OncotypeDX assay is now being studied in the TAILORx study – in which women with a recurrence score <11 will receive hormonal therapy only and women with a recurrence score >25 will receive hormonal therapy and chemotherapy, while women with an intermediate recurrence score of 11 to 25 will be randomized to receive hormonal therapy alone with or without chemotherapy. The OncotypeDX recurrence score may serve as a prognostic and predictive factor; that is, predicting both the chance of recurrence and the potential efficacy of hormonal and/or chemotherapy. Being a combination of 21 genes, however, the variable is not really a target as such. Furthermore, the assay is constituted mainly of genes representing ER, proliferation and HER2.

The HER2 protein and gene proved not only to have prognostic value [13,14] but also to be a genuine target that could be targeted for a specific response. The first treatment directed toward the HER2 oncoprotein was trastuzumab (Herceptin), a humanized monoclonal anti-HER2 antibody that produced 15% response rates as a single agent in...
HER2-positive patients [15] and significantly added to progression-free survival and overall survival in women with metastatic disease treated with several types of chemotherapy [16]. Concordance between local and central laboratories in determining HER2 has been problematic, and some patients who have been declared HER2-negative on central review in large trials have nonetheless received benefit from Herceptin. This is being further explored. Although HER2 is clearly a *bona fide* target for anti-HER2 therapies such as Herceptin, lapatinib (Tykerb) and others, problems with its measurement and with the fact that only ~15% of those who are HER2-positive actually seem to benefit remain a challenge. Interestingly HER2 has an inverse relationship with the ER and PgR [17]. HER2 is also involved in endocrine resistance [18,19]. Endocrine resistance may in part be reversible by adding a HER2-targeted agent, as has been done in the TANDEM trial [20].

Many of us have attempted to use HER2 measurements to predict differential response to anthracyclines, and this seems to be reliable although not all data are consistent [21-23]. Some feel that HER2 is predictive in this setting only because it is located on the topoisomerase I(α) gene of chromosome 17, topoisomerase IIα being an enzyme that is directly involved in the mechanism of action of anthracyclines against human cancers. Topoisomerase I(α) gene amplification and HER2/neu gene amplification and protein overexpression are closely related, and each predict for responsiveness to anthracyclines in some randomized trials comparing anthracycline-containing versus nonanthracycline-containing regimens [21,24]. Furthermore, topoisomerase IIα protein overexpression also appears to predict for differential anthracycline benefit, although it seems less closely related to topoisomerase I(α) overexpression and/or HER2/neu [25].

A meta-analysis of phase III trials evaluating the predictive value of HER2 and topoisomerase I(α) in early breast cancer patients treated with cyclophosphamide, methotrexate and 5-FU (CMF) versus anthracycline-based adjuvant therapy did not show a strongly statistically significant predictive value [26], particularly when the large British NEAT and BR9601 studies were included (J. Bartlett, personal communication). Some investigators now believe that chromosome 17 polyomysy is a more appropriate predictive marker [27,28]. The predictive value of these particular gene and protein changes remains to be clarified but it seems unlikely that they actually represent targets in the same way as HER2 is a target of Herceptin.

Additionally, targeted anti-HER2 agents such as lapatinib were meant to be effective against HER1 and HER2 overexpressing tumours but have proved to affect only individuals with HER2-positive tumours. Newer drugs such as bevacizumab – an antivascular endothelial growth factor antibody – are effective, but one cannot find a target that selects subgroups of patients in whom it is more valuable [29]. Some targeted therapies, including the endocrine therapies and trastuzumab, are therefore clearly available and can be directed in effective, albeit less than perfect, ways towards specified and relatively well measured targets. Some of the new biologics or so-called targeted agents such as bevacizumab and lapatinib, however, are not as clearly directed towards their targets as might have been initially predicted. I would therefore conclude that tailored targeted therapy for all may be worthwhile but is not yet realistic.

**Competing interests**

The author declares that they have no competing interests.

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