Targeting HER2-Driven Cancers in Non-Breast Cancer Malignancies

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Recent results from several large breast cancer clinical trials have demonstrated the value of targeting the cell surface receptor HER2 (erbB2) with multiple HER2-targeted agents [1-3]. This includes the use of combined antibodies (trastuzumab plus pertuzumab) as first line therapy in metastatic breast cancer [1], combined antibody and small molecule tyrosine kinase inhibitor (trastuzumab plus lapatinib) in early breast cancer [2] and antibody-drug conjugate (trastuzumab-DM1) in resistant disease [3]. The success of these approaches may now, in turn, suggest consideration of the use of such strategies in other cancer types in which single agent HER2 targeted therapy has some activity but is considered too limited for current use.

The cell surface receptor HER2 (or ErbB2) is a member of the EGFR family that drives the growth of a significant percentage of breast cancers [4]. It is amplified in approximately 20% of breast cancers and outcome for patients with these tumors has been linked to poor survival [5]. Breast cancers are now routinely assayed for the presence of HER2 amplification and such cases frequently benefit from treatment with the humanised antibody trastuzumab (herceptin) or the small molecule tyrosine kinase inhibitor lapatinib which are targeted against the HER2 receptor [6]. Several strategies have now been developed to improve outcome in HER2 positive breast cancer and also overcome resistance to trastuzumab. Recently published trials have now confirmed that combinations of HER2 targeted drugs such as trastuzumab + pertuzumab [1] or trastuzumab + lapatinib [2] are more effective in HER2 breast cancers than the single agents. The use of the antibody-drug conjugate trastuzumab-DM1 which combines the activity of trastuzumab with targeted delivery of the anti-microtubule agent DM1, is also effective in trastuzumab-resistant disease [3].

While these strategies have been pioneered within breast cancer, there is increasing interest in the possibility that other tumor types with amplification or overexpression of HER2 may also be responsive to HER2-targeted therapies and benefit from these approaches. Cancer types such as gastric cancer, ovarian cancer and lung cancer can express high levels of HER2. The TOGA phase III clinical trial demonstrated improved survival when trastuzumab was added to chemotherapy versus chemotherapy alone in treatment of HER2-positive advanced gastric or gastro-oesophageal junction cancer [7]. Preclinical studies in ovarian and lung cancer xenografts have demonstrated that combined therapies of trastuzumab and pertuzumab are effective against xenograft models of these diseases [8,9]. Therefore, while trastuzumab has not had marked efficacy as a single agent in these less sensitive diseases, these new HER2 combination strategies may prove more clinically effective.

However, several considerations will need to be taken into account in extending these approaches to other cancer types. Since these other diseases generally have lower percentages of HER2 amplification, the level of antitumor activity obtained with HER2-directed strategies will need to be sufficiently beneficial to justify the costs of diagnostic testing to identify potentially responsive tumors. The case of identification of these subgroups will also need to be considered and it may be that histological subgroups with known higher percentages of HER2 amplification should be investigated first. Furthermore, it would be valuable to identify additional biomarkers alongside HER2 that helps identify the sensitive population who are most likely to benefit. There are already biomarkers downstream of HER2 such as PTEN and SPROUTY [10,11] which can help identify breast cancers that are likely to be more resistant to trastuzumab and these may have value in high HER2 expressing non-breast cancers to help predict sensitivity.

If the twin hurdles of sufficient antitumour efficacy and cost-effectiveness of diagnostic testing can be overcome, then these useful HER2-targeted strategies may find wider applicability beyond just breast cancer and add to the growing use of targeted therapies in these other disease types.

References

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