Trastuzumab has been shown to be quite effective in reducing suffering and mortality from breast cancer in both the metastatic and adjuvant settings. With short follow-up, remarkably consistent results across five adjuvant, prospective, randomized clinical trials suggest that trastuzumab may decrease the odds of distant recurrence and mortality by approximately one half and one third, respectively. Dr George Sledge, who discussed the first presentations of the adjuvant trials at the 2005 Annual Meeting of the American Society of Clinical Oncology, proclaimed these results “astonishing,” and we agree (oral communication, May 2005).

However, there is a dark side to trastuzumab that may limit its utility in some patients. Although in general, trastuzumab has been extremely well tolerated, a surprisingly high incidence of congestive heart failure (CHF) was observed in the early studies of metastatic disease, especially in patients who were treated concurrently with doxorubicin. With this knowledge, most subsequent trial designs, particularly those in the adjuvant setting, avoided concurrent anthracycline and trastuzumab therapy, and included careful baseline and serial monitoring of cardiac function. Early results from these trials suggest that approximately 5% of all patients treated with adjuvant trastuzumab, either with nonanthracycline chemotherapy such as the taxanes or vinorelbine, or after all chemotherapy is complete, will develop some objective evidence of systolic cardiac dysfunction. Approximately 1% of patients will develop symptomatic CHF. These rates are approximately four to five times higher than in control patients who did not receive trastuzumab, and the absolute difference in echocardiogram or scintigram-detected cardiac dysfunction between trastuzumab-treated and control patients seems to be approximately 3% to 4%.

These observations raise several questions. First, given the enormous benefit of trastuzumab, is pre-existing cardiac dysfunction, especially if it is asymptomatic, sufficient reason to withhold the drug? Are the classic risk factors for cardiac disease, such as hypertension, diabetes, and family history, important predictors of trastuzumab-induced CHF? Is trastuzumab-related CHF reversible if the agent is discontinued, and is it safe to reinitiate this potentially life-saving agent if a patient has developed cardiac dysfunction while receiving it previously? Are there ways to avoid or minimize trastuzumab-related cardiac dysfunction? Will the incidence of this complication increase with longer follow-up of women who receive adjuvant trastuzumab? Finally, what is the mechanism of this perplexing toxicity?

In the last few years, a substantial number of publications have addressed the cardiac toxicity of trastuzumab in both prospective clinical trials and from individual institutional experiences. In this issue of the Journal of Clinical Oncology, Guarneri et al report their results from a retrospective analysis of 218 metastatic breast cancer patients who received trastuzumab for at least 1 year from 1998 to 2003. In this series, 28% of patients had some type of cardiac event (CE), and 10.9% had grade 3 cardiotoxicity, with one cardiac death. Therefore, coupled with the larger, prospective trials, the results from this study provide additional insight into our questions.

Given that all of the clinical trials of trastuzumab have excluded women with obvious pre-existing heart failure, we cannot determine if this condition should preclude a patient’s receiving it; however, until data to the contrary are reported, we believe it should. What about those who apparently have healthy hearts but have received pretrastuzumab anthracyclines, which are incorporated into most adjuvant regimens? In the adjuvant, prospective, randomized clinical trials conducted by the National Surgical Adjuvant Breast and Bowel Project (NSABP B-31) and the Breast Cancer Intergroup of North America, 6.7% of women who had completed four cycles of doxorubicin and cyclophosphamide (AC) were deemed ineligible to proceed to trastuzumab because of left ventricular dysfunction. We agree that this condition of eligibility was prudent. One might wonder if these patients could have been treated with trastuzumab and monitored closely with discontinuation of the drug on evidence of additional decline in cardiac function. In this regard, a cautionary note comes from the NSABP, which has reported that patients in NSABP B-31 with marginal post-AC left ventricular ejection fractions (LVEF) but who were still eligible for trastuzumab had higher rates of subsequent cardiac dysfunction than those who started trastuzumab with normal cardiac parameters. So for now, the presence of LV dysfunction or heart failure symptoms should exclude the addition of trastuzumab.

The study by Guarneri et al may provide additional clues to our questions. Although the study was retrospective and limited by a small number of events, careful analysis of Table 1 suggests that several pre-existing factors may predict trastuzumab CEs. For example, it seems that although only 7% of women who received radiation to the right side developed a CE, 26% of those who received radiation to the left side were affected, although these differences did not reach statistical significance. Again considering the limited power, pre-existing diabetes, coronary artery disease, and valvular dysfunction were all associated with apparent higher rates of trastuzumab-related CEs. Curiously, hypertension and older age were not. In the NSABP/Intergroup analysis of adjuvant patients, age and hypertension were risk factors for cardiac dysfunction. Additional analysis of the NSABP trial data suggest that...
smoking, family history, history of taking hypoglycemic and/or lipid lowering medications, and radiation to the left side were not risk factors.2,5,13

The good news, both from Guarneri et al and previously reported studies, is that trastuzumab-related cardiac dysfunction usually seems to be reversible. In this study, 11 (79%) of 14 women who stopped trastuzumab therapy after developing symptomatic CHF recovered with appropriate cardiac treatments. Eighty-nine percent of patients with an asymptomatic decrease in LVEF on cardiac imaging recovered, with or without cardiac treatment, after stopping trastuzumab. These data are similar to other studies, and differ substantially from the early experience with doxorubicin-induced cardiac failure, from which few patients recovered.14,16 Remarkably, trastuzumab was not stopped in 17 patients who developed asymptomatic cardiac dysfunction, and 13 of these did well, most without concurrent cardiac treatment. However, such a strategy requires intensive monitoring of cardiac symptoms and function. This issue raises the secondary question of the definition of heart failure. For example, in the M.D. Anderson (Houston, TX) experience, nearly one third of patients were classified as having a CE, yet a much smaller percentage developed symptoms. Furthermore, four symptomatic patients, including the one patient who died of apparent CHF, had a normal LVEF. These apparent inconsistencies may potentially be explained in several ways. First, the LVEF by echocardiography can be measured by several methods, and each has an associated degree of variability. For this reason, guidelines and standards have been developed to reduce such variability.17 Second, although both nuclear and ultrasound methods can measure LVEF accurately, the approaches and the range of normal values differ slightly between these two techniques. Therefore, when one observes an individual patient, it is important to use the same technique throughout the period of monitoring. Third, although the LVEF is a measure used by most clinicians to communicate about systolic ventricular function, it is influenced by both preload (filling pressures) and afterload (blood pressure), two factors that may vary during the course of treatment in these patients. Fourth, and perhaps most important, heart failure must be defined by more than just LVEF. In fact, the common nomenclature known as the New York Heart Association classification is based on symptoms and functional capacity, not the LVEF. Some patients can remain compensated or asymptomatic despite a significant decrease in LVEF—in some of these patients, the symptoms can be unmasked with more rigorous activity or exercise. Symptoms can also occur in the absence of a decrease in LVEF (the syndrome of diastolic heart failure). This isolated diastolic heart failure is found in up to one third of patients with symptomatic heart failure, and its treatment differs from that of those with systolic heart failure. Heart failure symptoms are often nonspecific and may mimic those of pulmonary disease, so thoughtful integration of cardiac examination and ventricular imaging are critical. Careful and frequent monitoring of patients treated with trastuzumab should include history and physical examination in addition to noninvasive imaging of ventricular function. In summary, until additional evidence is accumulated from the randomized trials, we should err on the side of caution in the use of this agent when either clinical or echocardiographic/scintigraphic evidence of heart failure exists or develops. New methods for early noninvasive detection of cardiac dysfunction are under development and validation. These methods, including tissue Doppler echocardiography, may enable more sensitive detection of subtle changes in myocardial function before those detected by measurement of LVEF, but careful studies will be necessary to ensure that we do not withhold trastuzumab on the basis of a surrogate indication of cardiac dysfunction.18

In addition to avoiding trastuzumab in women with cardiac problems, several strategies have been adopted or proposed to minimize cardiac toxicity. After the initial observations, most investigators quickly eliminated concurrent doxorubicin and trastuzumab, and general guidelines state that the combination is contraindicated.1,19 However, in a few preliminary trials, trastuzumab has been given concurrently with less cardiotoxic anthracycline compounds, such as epirubicin or liposomal-encapsulated doxorubicin, and the results have been promising.20-22 Another strategy has been to shorten the duration of adjuvant trastuzumab therapy. In a prospective randomized trial conducted in Finland (HERFin), with short follow-up and few events, 12 weeks of trastuzumab given with post-AC docetaxel or vinorelbine seems to be as effective and perhaps safer than longer therapy.23 Investigators in the HERceptin Adjuvant trial chose to delay initiation of trastuzumab until after all chemotherapy was complete. However, incidence of cardiac events seems similar to that in other adjuvant trials of 12 months of trastuzumab.24 Although intriguing, these strategies must be considered investigational, and they are not recommended for routine clinical care.

In the M.D. Anderson experience, prior treatment with doxorubicin, surprisingly, was not a risk factor for CE. Nonetheless, another strategy to avoid CHF might be to eliminate anthracyclines altogether in the adjuvant setting. Because preclinical studies had suggested possible synergy between trastuzumab and two nonanthracycline chemotherapeutic agents (docetaxel and platinum salts), a prospective adjuvant trial conducted by the Breast Cancer International Research Group contained a third arm in which no anthracycline was given. In this arm, patients received docetaxel, carboplatin, and trastuzumab. As expected, cardiac events in the docetaxel, carboplatin, and trastuzumab arm were similar to those in the control group that received only AC followed by docetaxel but no trastuzumab. However, although not statistically significant, the reduction in breast cancer recurrences in this third arm did not seem to be as robust, relative to control, as in the group that received AC-doxetaxel plus trastuzumab.

Another possibility is that cardiac toxicity might be related to trastuzumab itself, and therefore it might not occur with other agents that target HER-2 in different ways. Recently reported results suggest that the pan–epidermal growth factor receptor family tyrosine kinase inhibitor, lapatinib, seems active against HER-2–positive breast cancer, and that this agent might have less cardiac toxicity than has been observed with trastuzumab.25,26 More mature follow-up is needed to determine if the efficacy and safety profiles are similar to those of trastuzumab.

Finally, what is the mechanism of trastuzumab-related cardiac dysfunction? Several preclinical models have suggested that the epidermal growth factor receptor family is an important component of cardiac development; in adults, the HER-2 receptor seems to participate in regulation of growth, function, repair, and survival of cardiomyocytes. Addition studies suggest that trastuzumab has a direct action on human cardiomyocytes.25-28 Therefore, the
combination of induction of cardiac injury with preceding doxorubicin followed by inhibition of cardiac repair mediated by erbB2 may result in the cardiac dysfunction associated with trastuzumab.

In summary, the stakes are high. It is essential that we continue to deliver the “astonishing” beneficial effects of trastuzumab therapy to those who will benefit, while avoiding major short- and long-term toxicities. These challenges exist in the context of tightening health care dollars, both in the United States and abroad. We eagerly await release of analyses of the large ongoing and future randomized trials that will permit additional individualization of this exciting approach.

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