Title
Do we need a new end point in clinical trials today?

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While emphasizing the mutual agreement about these broad issues, we would like to address three minor clarifications in response to points raised by Drs. Skeel and Ganz.

First, use of separate or companion protocols for quality of life assessment in Southwest Oncology Group trials is an administrative device currently mandated by the National Cancer Institute. It primarily serves as a mechanism for distinguishing credits for cancer control research from those for therapeutic research. We have no reason to believe that this approach will “limit clinician involvement in implementation of the research and the data collection process,” a concern noted by Dr. Ganz. To the contrary, Drs. Ian Thompson and Stephen Smalley, investigators for two Southwest Oncology Group therapeutic trials, have contributed to all aspects of the design of these companion protocols—for example, development of treatment-specific items for the questionnaire, discussions regarding expected effects on different components of quality of life over time, and identification of meaningful times for assessment.

An earlier experience with quality of life assessment as an add-on end point in a breast cancer therapeutic trial resulted in poor data collection compliance. We believe this was due to group inexperience with collection of patient-based quality of life data and with insufficient quality control procedures. Physician interest and a substantial investment in data manager training are key variables in successful quality of life assessment in a cooperative group study.

Second, given sufficient physician involvement, we would be interested in assessing quality of life in phase II trials. Limited resources and greater physician interest in such assessment for phase III trials led to our decision to begin with comparative trials. We agree with Dr. Skeel that studying quality of life in patients on a combined-modality regimen for head and neck cancer at the phase II level could yield informative data on trade-offs relating to treatment response and toxicity prior to the design of the phase III trial.

Response

We read with great interest the remarks of Drs. Roland Skeel (1) and Patricia Ganz on quality of life measurement in clinical trials. We are heartened by the considerable agreement emerging in the literature with respect to the following major issues addressed in our review (2):

(a) supplementation of the physician’s report with the patient’s report of quality of life;
(b) development of a component-based view of quality of life, with components measured separately (as opposed to a more global construct) and modules tailored to different protocols (e.g., treatment-specific symptoms); and
(c) consideration of the quality control issues that affect successful implementation in a clinical trial setting.
Third, we also agree with Dr. Skeel (1) that a modular approach is best suited to the varying nature of clinical trial protocols. We do, however, think that we have selected existing instruments appropriate for measuring physical functioning, emotional functioning, social functioning, general symptoms, and global quality of life in a number of different protocols. Only the treatment-specific items change with the protocol, and even these items can be used across trials that examine the same disease site (e.g., the two current prostate cancer trials).

We recognize the imperfect nature of existing measures of quality of life, including those we selected, and the vigorous research activity that is generating new instruments [e.g., the developmental work on a core instrument by the European Oncology Group on Research and Treatment of Cancer (3)]. Consequently, we have elected annual re-evaluation of the questionnaires constituting our Quality of Life Questionnaire. These reviews will allow us to judge the appropriateness of the measures for different patient groups. In this vein, we are exploring the possibility of translating our Quality of Life Questionnaire into Spanish so we do not eliminate from our database patients who speak Spanish but not English. Communication with other investigators involved in measuring quality of life in clinical trials or multi-institution research settings can facilitate this process of continual examination of the appropriateness of selected questionnaires.

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