Taking a Measured Approach to Toxicity Data in Phase I Oncology Clinical Trials

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The standard categorical system for assessing attribution of toxicity to study drug(s) in phase I trials is cumbersome and uninformative. Although a binary system ("related" vs. "unrelated") would be sufficient to define maximum tolerated dose (MTD), a probability estimation would better support dose selection for randomized dose-ranging phase II trials.

In this issue of Clinical Cancer Research, Eaton and colleagues (1) analyzed data from 11,909 toxicities on 38 phase I trials sponsored by the NCI and reported that the rate of drug-related toxicity increased with dose, whereas the rate of unrelated toxicity did not. They found that these relationships to dose were similar when "unrelated" and "unlikely" related toxicities were considered separately or grouped together as "unrelated," and when "possibly," "probably," and "definitely" related toxicities were considered separately or grouped together as "related." They propose a simplified binary system of "related" versus "unrelated" when assessing the attribution of adverse events (AE) to study drug(s) in phase I trials.

The Common Toxicity Criteria (now the Common Terminology Criteria for Adverse Events) was first developed in 1983 by the Cancer Treatment and Evaluation Program (CTEP) of NCI to standardize the language of AEs reported on NCI-sponsored trials. With respect to attribution, the International Conference on Harmonisation (ICH) in 1994 stated in its E2A guideline: "the expression 'reasonable causal relationship' is meant to convey in general that there are facts (evidence) or arguments to suggest a causal relationship" between the AE and the investigational agent(s)/intervention (2). Citing the ICH E2A guideline, CTEP requires that AEs be reported with one of five attributions to study drug: "definite," "probable," "possible," "unlikely," and "unrelated," where these mean the AE "is clearly related," "is likely related," "may be related," "is doubtfully related," or "is clearly not related" to the intervention, respectively (3).

The distinctions between adjacent categories (e.g., "possibly" vs. "unlikely" related) are inherently ambiguous and subjective, although studies to formally assess interrater variability are lacking. Misattribution is common, with nearly 50% of AEs in the placebo arms of two randomized phase III trials attributed to study drug (4). Disease-related symptoms and drug toxicities can look similar, especially for AEs that are common, not previously known to be associated with the drug (or its class), and not causally related to its mechanism of action. Fatigue is an AE that can easily be misattributed to study drug, whereas the acneiform rash associated with anti-EGFR therapy would almost never be misattributed.

The study by Eaton and colleagues has sound methodology and should inform the conduct of traditional first-in-human studies that aim to define MTD. Most such studies are being conducted by industry sponsors (rather than NCI) and are already using a binary attribution system. However, several limitations of the study minimize the generalizability of the findings to modern oncology drug development. First, their analysis was restricted to monotherapy trials, whereas many current phase I trials are combination studies with the potential for pharmacokinetic and/or pharmacodynamic interactions between the drugs, and the issue of attribution to the investigational drug versus the standard drug (5). Second, they only considered toxicities recorded during the first cycle of therapy as defined in each protocol. Long-term and often less severe toxicities are important in assessing the tolerability of oral targeted therapies (6), and we are now in the era of investigational immunotherapy where immune-related AEs may be delayed or insidious (7).

An underemphasized finding of this study is the high rate of misattribution of toxicities as being related to study drug when they are almost certainly not related. The y-intercepts on Fig. 1A, from Eaton and colleagues' article, demonstrate that approximately 60% of patients have grade $\geq 1$ toxicities and approximately 12% of patients have grade $\geq 3$ toxicities attributed to study drug at doses that are approaching zero as a percentage of the MTD. There are major differences between "possibly," "probably," and "definitely" related attributions in this regard, as Fig. 2C–E, from Eaton and colleagues' article, show that approximately 60%, 20%, and 10% of patients have grade $\geq 1$ toxicities, respectively, that are attributed to study drug at these lowest doses. The authors hypothesize that toxicities observed at the lowest doses may be idiosyncratic (i.e., dose independent), but it is not plausible for this to account for toxicities in approximately 60% of patients across a variety of drugs and therapeutic classes. Instead, it is much more plausible that these toxicities were documented by investigators as being related to study drug when they were actually due to disease or other factors, as in the placebo arms of the study by Hillman and colleagues (4).

A binary classification of "related" versus "unrelated" with respect to toxicity attribution is analogous to a classification of...
“responders” versus “nonresponders” with respect to efficacy. Just as response rates do not tell the whole story regarding efficacy and are supplemented by waterfall plots and spider plots that depict change in tumor size as a continuous variable (8), a “related” versus “unrelated” classification would not tell the whole story regarding toxicity. A “possibly” related toxicity is like “stable disease” in that it can be due to either drug or disease. In modern oncology drug development, the primary objective of phase I trials should be to select doses for randomized dose-ranging phase II trials, rather than to identify the MTD (9). The optimal selection of doses for a dose-ranging phase II trial requires a sophisticated understanding of the relationship between dose toxicity and dose efficacy. The understanding of the relationship between dose and toxicity would be enhanced by assigning more weight to toxicities that are more likely to be related to study drug(s), as this would minimize the impact of misattribution discussed above. Weighting by attribution would also support quantitative comparisons between randomized arms in a phase II trial with respect to toxicity.

Figure 1 illustrates a proposed system for assessing attribution of AEs to study drugs using probability estimation. Investigators would electronically click on an arrow and move it along a probability distribution that ranges between 0 and 1. The investigator would move the arrow to provide a quantitative (and visual) assessment of the probability that an AE is related to study drug(s) at a point in time. In the hypothetical example to the left, the probability decreases between day 8 and day 15 because the toxicity did not resolve after the study drug was discontinued.

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No potential conflicts of interest were disclosed.

**Authors’ Contributions**
Conception and design: M.R. Sharma, M.J. Ratain
Development of methodology: M.R. Sharma
Acquisition of data (provided animals, acquired and managed patients, provided facilities, etc.): M.R. Sharma
Analysis and interpretation of data (e.g., statistical analysis, biostatistics, computational analysis): M.R. Sharma
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**References**


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