CORRECTION TO “THREE-DIMENSIONAL QUANTITATIVE STRUCTURE-ACTIVITY RELATIONSHIP ANALYSIS OF HUMAN CYP51 INHIBITORS”

In Table 1 on page 496 of the article above [Ekins S, Mankowski DC, Hoover DJ, Lawton MP, Treadway JL, and Harwood HJ Jr (2007) Drug Metab Dispos 35:493–500], the chemical structure for analog L was inadvertently dropped and replaced with a duplicate of the structure for benzimidazole. The correct structure for analog L appears below.

The online version of this article will be corrected in departure from the print version.

The printer regrets this error and apologizes for any confusion or inconvenience it may have caused.