Corrections to A Medicinal Chemist’s Guide to Molecular Interactions [J. Med. Chem. 2010, 53, 5061. DOI: 10.1021/jm100112j], Caterina Bissantz, Bernd Kuhn, and Martin Stahl*

Page 5063. The first sentence of the second paragraph in the left column should be replaced by the following: “In an impressively comprehensive study on several series of thrombin inhibitors, the Hangauer and Klebe groups have analyzed the sources of cooperativity between a lipophilic interaction and a hydrogen bond”.

Page 5075. Figure 16 caption should read as follows: “Radial distribution of carbon atoms around phenyl rings: (a) CH₃ bound to O or N (CSD statistics); (b) CH₃ bound to sp³ C (CSD statistics); (c) CH₃ bound to O or N (PDB statistics). Darker gray corresponds to higher density; peaks above a numerical value of 90 are colored red”.

Page 5083. Reference 159 should be replaced by the following: Manley, P. W.; Cowan-Jacob, S. W.; Fendrich, G.; Mestan, J. Molecular interactions between the highly selective pan-Bcr-Abl inhibitor, AMN107, and the tyrosine kinase domain of Abl. Blood 2005, 106, 940a−941a. DOI: 10.1021/jm100950p

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