Hiroyuki Kagechika, Emiko Kawachi, Yuichi Hashimoto, Toshiyuki Himi, and Koichi Shudo*: Retinobenzoic Acids. 1. Structure-Activity Relationships of Aromatic Amides with Retinoidal Activity.

Page 2183. Table I. Footnote b should read "The ratio of $E\bar{D}_{50}$ (retinoic acid) to ED_{50} (a test compound), both values having been obtained in concurrent experiments. This is also the case in the other tables." The same is the case in the description of "relative activity" in the Experimental Section (page 2188, left column, lines 35-37).

Page 2186. The correct structure of Am586 (Chart II) should be as shown below.

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Richard A. Glennon: Brain 5-HT_{1A} Receptors. Edited by C. T. Dourish, S. Ahlenius, and P. H. Hutson..

Page 276. The first sentence of my review should read: "Brain 5-H T_{1A} Receptors is a compilation of 23 papers by an esteemed group of European authors who have made a significant contribution to this field of study."

Brigitte Jamart-Gregoire, Paul Caubere,* Marie Blanc, Jean-Pierre Gnassounou, and Charles Advenier*: A New Series of Tricyclic (Aryloximino)propanolamines Displaying Very High Selective β_2 -Blocking Properties.

Page 317. In Table III, second line of the last column, the value should read 343 and not 33.

James F. Kerwin, Jr,* Alex M. Nadzan, Hana Kopecka, Chun Wel Lin, Thomas Miller, David Witte, and Stanley Burt: Hybrid Cholecystokinin (CCK) Antagonists: New Implications in the Design and Modification of CCK Antagonists.

Page 740. Reference 5 should be amended to read: (5) Preliminary reports of our work were presented at the Cold Spring Harbor Workshop on "CCK Antagonists", Nadzan, A. M.; Kerwin, J. F., Jr.; Kopecka, H.; Lin, C. W.; Miller, T.; Witte, D.; Burt, S. Cholecystokinin Antagonists, Neurology and Neurobiology, Vol. 47; Wang, R. Y., Schoenfeld, R., Eds.; Alan R. Liss, Inc.: New York, 1988; pp 93-103, and at the 21st National Medicinal Chemistry Symposium, June 19–23, 1988, Minneapolis, MN. Since these presentations, reports addressing this point have appeared: Makovec, F.; Chiste, R.; Rovati, L. C.; Setnikar, I. Gastroenterology 1988, 94(5), A279. Freidinger, R. M. European Patent Application 87305088.4. Makovec, F.; Chiste, R.; Peris, W.; Rovati, L. European Patent Application 87830442.7.

Federico Maria Arcamone,* Fabio Animati, Brunella Barbieri, Emanuela Configliacchi, Roberto d'Alessio. Cristina Geroni, Fernando Carlo Giuliani, Ettore Lazzari, Milena Menozzi, Nicola Mongelli,* Sergio Penco, and Maria Antonietta Verini: Synthesis, DNA-Binding Properties, and Antitumor Activity of Novel Distamycin Derivatives.

Page 774. The structure for 15 is as shown below:

James Inglese, Richard A. Blatchly, and Stephen J. Benkovic*: A Multisubstrate Adduct Inhibitor of a Purine Biosynthetic Enzyme with a Picomolar Dissociation Constant.

Page 937. (1) Replace ref 3 by: Jones, T. R.; Calvert, A. H.; Jackman, A. L.; Eakin, M. A.; Smithers, M. J.; Betteridge, R. F.; Newell, D. R.; Hayter, A. J.; Stocker, A.; Harland, S. J.; Davies, L. C.; Harrap, K. R. J. Med. Chem. 1985, 28, 1468-1476. (2) In sentence 4, replace 5,10-dideazatetrahydroaminopterin by 5,10-dideazatetrahydrofolate. (3) Reference 5 is not applicable.

Munefumi Kanao,* Yoshifumi Watanabe, Youchi Kimura, Junji Saegusa, Kenjiro Yamamoto, Hideyuki Kanno, Naoaki Kanaya, Hideo Kubo, Shin-ichiro Ashida, and Fumiyoshi Ishikawa: Thromboxane A₂ Synthetase Inhibitors. 2. Syntheses and Activities of Tetrahydronaphthalene and Indan Derivatives.

Page 1330. Table I, line 1, the compound 47 should read 47a.

Page 1333. Column 2, line 42, 243-251 °C, the melting point of the compound 47a, should be 270-276 °C.

Christopher F. Bigge, James T. Drummond, Graham Johnson,* Thomas Malone, Albert W. Probert, Jr., Frank W. Marcoux, Linda L. Coughenour, and Laura J. Brahce: Exploration of Phenyl-Spaced 2-Amino-(5-9)-phosphonoalkanoic Acids as Competitive N-Methyl-Daspartic Acid Antagonists.

Page 1580. The biological results of compounds 10 (the meta,1,1 derivative) and 15 (the para,1,1 derivative) have been inverted. Compound 10 has an IC₅₀ value of 3.3 μ M in the receptor binding assay (Table I), and further, the results listed under compound 15 in Table II are actually those of compound 10. All discussion under the heading of Pharmacology and Discussion in the text that refers to compound 15 actually is a discussion of the activity of the meta derivative 10.