

sample cuvettes. Dimethyl sulfoxide had no effect on the binding spectra with addition up to 10% of the final suspension volume. The K_s value was determined from a Scatchard plot (absorbance difference vs absorbance difference/concentration = slope K_s) and is the average of three determinations ($\pm 10\%$).

Acknowledgment. This work was supported by grants from the Cancer Research Campaign (to the Biomolecular

Structure Unit) and from the Cancer Research Campaign/Medical Research Council (to the Section of Drug Development).

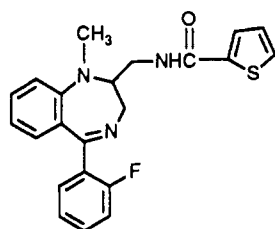
Supplementary Material Available: Tables listing hydrogen atom coordinates, anisotropic thermal parameters, and bond lengths and angles (19 pages). Ordering information is given on any current masthead page.

Additions and Corrections

1990, Volume 33

Mark G. Bock,* Robert M. DiPardo, Ben E. Evans, Kenneth E. Rittle, Willie L. Whitter, Daniel F. Veber, Roger M. Freidinger, Raymond S. L. Chang, T. B. Chen, and Victor J. Lotti: Cholecystokinin-A Receptor Ligands Based on the κ -Opioid Agonist Tifluadom.

Page 451. The structure for **2** is incorrect. The correct structure is



2, (*R,S*)-tifluadom
2a, (*R*)-(+)-tifluadom
2b, (*S*)-(-)-tifluadom

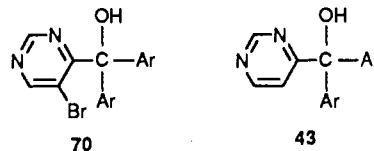
Robin D. Clark,* Jacob Berger, Pushkal Garg, Klaus K. Weinhardt, Michael Spedding,* Andrew T. Kilpatrick, Christine M. Brown, and Alison C. MacKinnon: Affinity of 2-(Tetrahydroisoquinolin-2-ylmethyl)- and 2-(Isoindolin-2-ylmethyl)imidazolines for α -Adrenoceptors. Differential Affinity of Imidazolines for the [^3H]Idazoxan-Labeled α_2 -Adrenoceptor vs the [^3H]Yohimbine-Labeled Site¹.

Page 597. Structure **6a,b** in Table I is incorrect. The correct structure is on page 596.

Page 597. The last sentence of the Chemistry section should read "Isoindolines **8a,b** ..." rather than **10a,b**.

C. David Jones,* Mark A. Winter, Kenneth S. Hirsch, Nancy Stamm, Harold M. Taylor, Howard E. Holden, James D. Davenport, Eriks V. Krumkalns, and Robert G. Suhr: Estrogen Synthetase Inhibitors. 2. Comparison of the in Vitro Aromatase Inhibitory Activity for a Variety of Nitrogen Heterocycles Substituted with Diarylmethane or Diarylmethanol Groups.

Page 419, Table IV, route 1. The structures for **70** and **43** are incorrect. The correct structures are



Gloria Cristalli, Palmarisa Franchetti, Mario Grifantini,* Giuseppe Nocentini, and Sauro Vittori: 3,7-Dideazapurine Nucleosides. Synthesis and Antitumor Activity of 1-Deazatubercidin and 2-Chloro-2'-deoxy-3,7-dideazaadenosine.

Page 1466. The ^1H NMR data for compound **18** (2'-deoxy- β -D-ribofuranosyl derivative) should read: δ 6.09 (s, 2, NH_2), 6.22 (pt, 1, $\text{C}_1\text{-H}$), 6.68 (d, 1, $J = 3$ Hz, $\text{C}_3\text{-H}$), 6.79 (d, 1, $J = 6$ Hz, $\text{C}_7\text{-H}$), 7.38 (d, 1, $J = 3$ Hz, $\text{C}_2\text{-H}$), 7.56 (d, 1, $J = 6$ Hz, $\text{C}_6\text{-H}$).