## Additions and Corrections

The Secondary  $\beta$ -Deuterium Isotope Effect and the Cage Effect in the Thermal Decomposition of Azobis- $\alpha$ -phenylethane [J. Am. Chem. Soc., 88, 3775 (1966)]. By STANLEY SELTZER\* and EDWIN J. HAMILTON, JR., Chemistry Department, Brookhaven National Laboratory, Upton, New York 11973.

The abscissa of Figure 2 is incorrect and should read 2[IV]/[III + IV + V]. The caption should read: "A plot of the ratio of rate constants vs. twice the ratio of cross-product to the sum of all products for equal concentrations of starting material."

Line 2, left column of p 3780, should read: "2[IV]/[III + IV + V] = 0.9967."

Ten lines from the bottom of the right column of p 3779 should read: "The variation of the ratio of cross-product to total product as a function of the ratio of first-order rate constants, in the absence of a cage effect, is shown in Figure 2."

Mass Spectral Rearrangements of Siliconium Ions. Migration of Electronegative Groups from Carbon to Silicon [J. Am. Chem. Soc., 91, 6544 (1969)]. By WILLIAM P. WEBER,\* RAYMOND A. FELIX, and ALVIN K. WILLARD, Department of Chemistry, University of Southern California, Los Angeles, California 90007.

On reexamination of the mass spectra of methyl 3-trimethylsilylpropionate under high resolution conditions, an error in the composition of the m/e 105 ion became apparent. It is not  $C_3H_9SiO_2$  as indicated in eq 3 and Figure 1, but rather  $C_4H_{13}SiO$ . Peak matching high resolution data: calcd for  $C_4H_{13}SiO$ , 105.0732; found, 105.0733. A metastable peak at m/e 68.9 =  $(105)^2/160$  indicates that this ion may be formed directly from the parent ion by loss of  $C_3H_3O$ .

Stable Carbocations. CXXXVII. Cycloheptenyl, Cyclooctenyl, and Cyclononenyl Cations [J. Am. Chem. Soc., 94, 6434 (1972)]. By GEORGE A. OLAH\* and GAO LIANG, Department of Chemistry, Case Western Reserve University, Cleveland, Ohio 44106.

In Table II, cmr shifts for the first and third ions were interchanged and should be:

Ambident Behavior of Nitrite Ion. Reactions of 2,4-Dinitro-halobenzenes and 1,2,4-Trinitrobenzene with Sodium Nitrite [J. Am. Chem. Soc., 95, 2133 (1973)]. By DAVID H. ROSENBLATT,\* WILLIAM H. DENNIS, JR., and RICHARD D. GOODIN, Physical Research Division, Chemical Laboratory, Edgewood Arsenal, Edgewood Arsenal, Maryland 21010.

An error has been found in the calculation of rate constant  $k_1$  from  $\Delta OD/\Delta t$  for the compounds 2,4-dinitrochlorobenzene (DNClB) and 2,4-dinitrobromobenzene (DNBrB) for their reactions with nitrite by O-attack. In

Table I the values for  $k_1$  should read, for DNC1B and DNBrB,  $2.5 \times 10^{-5}~M^{-1}~{\rm min^{-1}}$  and  $2.2 \times 10^{-5}~M^{-1}~{\rm min^{-1}}$ , respectively. Therefore the N:O ratio of nitrite attack on DNC1B is 70:1 and not 18,000:1, as implied by the previously stated rate constants. This correction has no effect upon the other rate constants nor upon the conclusions stated.

Structure of the Peptide Antibiotic Amphomycin [J. Am. Chem. Soc., 95, 2352 (1973)]. By MIKLOS BODANSZKY,\* GERALD F. SIGLER, and AGNES BODANSZKY, Department of Chemistry, Case Western Reserve University, Cleveland, Ohio 44106.

The + sign preceding the name of 3-isododecenoic acid should be deleted.

Stereochemistry of  $\alpha$ -Halo Sulfoxides. II. Interdependent Stereochemistry at Sulfur and  $\alpha$ -Carbon in the  $\alpha$ -Halogenation of Sulfoxides [J. Am. Chem. Soc., 95, 7431 (1973)]. By PAOLO CALZAVARA, MAURO CINQUINI, STEFANO COLONNA, ROBERTO FORNASIER, and FERNANDO MONTANARI,\* Centro C.N.R. e Istituto di Chimica Industriale dell' Università, Milan 20133, Italy.

On page 7432, in SchemeI, the notations of absolute configuration (S)-(-)-5b,6b and (S)-(-)-5a,6a should be replaced by (R)-(-)-5b,6b and (R)-(-)-5a,6a, respectively.

Photochemical Addition of Benzene to Cyclopentene [J. Am. Chem. Soc., 95, 8250 (1973)]. By V. Y. MERRITT, J. CORNELISSE, and R. SRINIVASAN,\* IBM Thomas J. Watson Research Center, Yorktown Heights, New York 10598.

The correct structure of VIIIa is

$$H_{C}$$
 $H_{C}$ 
 $H_{D}$ 
 $H_{E}$ 
 $H_{E}$ 
 $H_{F}$ 

In VIIIc, positions A-E and G are deuterated.

<sup>13</sup>C Nuclear Magnetic Resonance Studies of Organometallic Compounds. I. trans-Methylplatinum(II) Derivatives [J. Am. Chem. Soc., 95, 8574 (1973)]. By M. H. CHISHOLM, H. C. CLARK,\* L. E. MANZER, J. B. STOTHERS, and J. E. H. WARD, Department of Chemistry, The University of Western Ontario, London, Ontario, N6A 3K7, Canada.

Table II, item 14 under other <sup>13</sup>C nmr data:  $\delta CH_3$  is 49.0 not 9.2 and  ${}^2J_{PtCC}$  was not observed and should be denoted  ${}^2J_{PTCC}^d$ .

Table V, item 8: the left-hand  $|{}^3J_{\text{PtCNC}}|$  value should be  $66(\pm 1)$ , not  $56(\pm 1)$ ; item 15:  $\delta C^1$  should be 52.0, not 41.6, while  $\delta C^2$  should be 41.6, not 52.0.

Transition Metal Hydroborate Complexes. VI. The Solid-State Structure of  $\mu$ -Bis(cyanotrihydroborato)-tetrakis(tri-