

Additions and Corrections

Protonation of Nitro Groups. Diprotonation of β -Nitrostyrenes in Trifluoromethanesulfonic Acid [*J. Am. Chem. Soc.* **1986**, *108*, 3029–3032]. TOMOHIKO OHWADA, TOSHIHARU OHTA, and KOICHI SHUDO*

Page 3029: The intensity of the UV absorption of (*E*)- β -nitrostyrene should be $\log \epsilon_{\max} = 4.38$ instead of 4.86.

Page 3030: The digital resolutions in the observed NMR spectra were as follows: ± 0.4 Hz in ^1H NMR spectra (Table II) and ± 1.2 Hz in ^{13}C NMR spectra (Table III). The coupling constants in the tables should be read in the consideration of the digital resolutions.

Bimanes. 18. (Haloalkyl)-1,5-diazabicyclo[3.3.0]octadienediones (Halo-9,10-dioxabimanes): Sulfur "Extraction" by *syn*-(1-Bromoethyl,methyl)bimane. Glutathione "Sulfide" from the Tripeptide Thiol, Glutathione [*J. Am. Chem. Soc.* **1986**, *108*, 4532–4541]. ANNETTE E. RADKOWSKY, EDWARD M. KOSOWER,* DEBORAH EISENBERG, and ISRAEL GOLDBERG

1. Page 4536: The title for Scheme I should read as follows—Scheme I. Reactions of *syn*-(1-Bromoethyl,methyl)-bimane (**1**) with glutathione.^a

2. Page 4540: Left-hand column, ninth line from bottom, in the paragraph on *trans*- μ -(*S*)-*syn*-(CH(CH₃),CH₃)B (*trans*-2): Fluorescence (λ_{\max} , (ϕ_F) 3% CH₃CN/H₂O, pH 7.31) should be substituted for

UV (3% CH₃CN/H₂O, pH 7.31)

3. Page 4540: Right-hand column, second line from top: Fluorescence (λ_{\max} , (ϕ_F) 3% CH₃CN/H₂O, pH 7.31) should be substituted for

UV (3% CH₃CN/H₂O, pH 7.31)

4. Page 4541: Right-hand column, next to the last line in the Registry list, there is no compound **10** (isomer 3). For compound **10**, there are only isomer **1** and isomer **2**.

Synthesis and X-ray Crystal Structure of a Novel Bimetallic Bis(η^5 -dicarbollide) Aluminum Sandwich Complex [*J. Am. Chem. Soc.* **1986**, *108*, 5367]. WILLIAM S. REES, JR., DAVID M. SCHUBERT, CAROLYN B. KNOBLER, and M. FREDERICK HAWTHORNE*

Unfortunately, at the time that we recorded spectral data for compound **2**, we did not recognize that compound **1** was present as a contaminant. The correct spectral data for pure **2** are as follows: ^{11}B NMR (C₆H₆; referenced to external BF₃·OEt₂ at δ 0 with chemical shifts upfield of δ 0 reported as negative; ambient temperature) δ -31.7 (d, area 1), -21.6 (d, area 1), -16.8 (d, area 4), -12.3 (d, area 3), all resonances collapsed to singlets upon ^1H decoupling; ^1H NMR (C₆D₆, referenced to residual protons in C₆D₆ at δ 7.15; ambient temperature) δ 2.51 (singlet, carboranyl

C-H protons), 0.84 (t, CH₃), 0.17 (q, CH₂), 0.0–4.0 ppm (br, B-H protons); characteristic IR (cm⁻¹; NaCl, Nujol mull) 3023 (m, carboranyl C-H str), 2578 (s, B-H str), 2532 (s, B-H str), 1094 (m), 988 (m), 689 (m), 656 (m). We regret any confusion this mistake may have caused. A full paper describing this system is forthcoming.

Structure and Properties of HFe₃(CO)₉BH₂R and the Conjugate Bases [HFe₃(CO)₉BH₂R]⁻ (R = H and CH₃). Inorganometallic Analogues of Hydrocarbyltriiron Clusters [*J. Am. Chem. Soc.* **1986**, *108*, 3304]. JOSE VITES, CATHERINE E. HOUSECROFT, CHARLES EIGENBROT, MARGARET L. BUHL, GARY J. LONG, and THOMAS P. FEHLNER*

Page 3309: The ^{11}B NMR for III should read as follows: ^{11}B NMR (C₆D₆, 20 °C) δ 1.8 (br m, 230 Hz, fwhm), $\{^1\text{H}\}$ (br s, 150 Hz, fwhm).

Synthesis and Characterization of the First Stable Cyanocyclophosphazenes [*J. Am. Chem. Soc.* **1986**, *108*, 6089]. J. STEVEN RUTT, MASOOD PARVEZ, and HARRY R. ALLCOCK*

At the time of publication, we were unaware of an earlier paper by H. Roesky and M. Banek in *Z. Naturforsch.* **1979**, *34B*, 752, in which the cyclophosphazenes N₃P₃F₅CN and N₄P₄F₇CN were reported. The organic cosubstituents in the compounds we described, N₃P₃(OPh)₃CN and N₃P₃(NMe₂)₃(CN)₃, are needed for prolonged stabilization against moisture but apparently are not essential for stability under anhydrous conditions.

Double Nitrogen Inversion in Sesquibicyclo Hydrazines and Their Cation Radicals [*J. Am. Chem. Soc.* **1986**, *108*, 7926–7934]. STEPHEN F. NELSEN,* TIMOTHY B. FRIGO, YAESIL KIM, JAMES A. THOMPSON-COLON, and SILAS C. BLACKSTOCK

The name of S. C. Blackstock was regrettably omitted from the list of authors.

Synthesis of "Iso-EPSP" and Evaluation of Its Interaction with Chorismate Synthase [*J. Am. Chem. Soc.* **1986**, *108*, 8068–8071]. PAUL A. BARTLETT,* UDAY MAITRA, and PAUL M. CHOUINARD

The last line in the caption for Figure 1 should read: (□) no inhibitor, (■) 2.5 μM , (Δ) 5 μM , (\blacktriangle) 10 μM , and (○) 20 μM .

"Remote Control" of Flavin Reactivities by an Intramolecular Crown Ring Serving as a Metal-Binding Site [*J. Am. Chem. Soc.* **1987**, *109*, 923–924.]. SEIJI SHINKAI,* KEI KAMEOKA, KAORI UEDA, and OSAMU MANABE

Page 924, reference 19: $\Phi_f = 1.07E_T(30) - 0.017$ should read $\Phi_f = -0.017E_T(30) + 1.07$.