Additions and Corrections

Experimental Determination of the Bond Density of Molecular Hydrogen in Momentum-Space by Binary (e,2e) Spectroscopy [J. Am. Chem. Soc. 1984, 106, 5859–5864]. K. T. LEUNG and C. E. BRION*

Page 5860, Table I: The order of column 1 should be reversed, thus it should read down as follows: $\langle p^{-2} \rangle$, $\langle p^{-1} \rangle$, $\langle p^0 \rangle$, $\langle p^1 \rangle$, $\langle p^2 \rangle$.

Cleavage of Ruthenium and Osmium Porphyrin Dimers: Formation of Organometallic Ruthenium Porphyrin Complexes and Highly Reduced Metalloprophyrin Species [J. Am. Chem. Soc. 1985, 107, 4570]. J. P. COLLMAN,* P. J. BROTHERS, L. MCELWEE-WHITE, E. ROSE, and L. J. WRIGHT

Page 4570: Dr. Eric Rose's name should have appeared as Eric Rose[‡] with the following footnote.

[†]Université P. et M. Curie, Laboratoire de Chimie Organique, Tour 45, UA 508, 4 Place Jussieu, 75230 Paris Cedex 05, France.

Page 4571: The acknowledgment should have included the following: Acknowledgment is also made to NATO Grant 85/300 for support of Dr. Rose's work.

Reactivity of Zero-Valent Metalloporphyrin Dianions toward Organic Electrophiles [J. Am. Chem. Soc. 1985, 107, 6110-6111]. J. P. COLLMAN,* P. J. BROTHERS, L. MCELWEE-WHITE, and E. ROSE

Page 6110: Dr. Eric Rose's name should have appeared as Eric Rose[‡] with the following footnote.

¹Université P. et M. Curie, Laboratoire de Chimie Organique, Tour 45, UA 408, 4 Place Jussieu, 75230 Paris Cedex 05, France.

Biomimetic Models for Cysteine Proteases. 3. Acylation of Imidazolium–Thiolate Zwitterions by p-Nitrophenylacetate as a Model for the Acylation Step and Demonstration of Intramolecular General-Base-Catalyzed Delivery of H_2O by Imidazole to Thiol Esters as a Model for the Deacylation Step [J. Am. Chem. Soc. 1985, 107, 7669–7679]. J. P. Street, K. I. Skorey, R. S. Brown,* and R. G. Ball

Page 7674, Table IV: pK_a values for **2b** and **3b** should be interchanged. This same change applies to Table I of the preliminary communication.

Page 7676, Table VI: k_1 for **2b** and **3b** should be interchanged, e.g., **2b**, $k_1 = 13.2 \text{ M}^{-1} \text{ min}^{-1}$; **3b**, $k_1 = 7.64 \text{ M}^{-1} \text{ min}^{-1}$. This same change applies to Table II of the preliminary communication.

X-ray and ¹H NMR Analyses of the Structure and Conformation of 2-Azacoformycin, a Potent Inhibitor of Adenosine Deaminase [J. Am. Chem. Soc. 1985, 107, 7635–7640]. GEORGE I. BIRNBAUM,* JEAN ROBERT BRISSON, STEVEN H. KRAWCZYK, OSCAR L. ACEVEDO, and LEROY B. TOWNSEND

Page 7635: The list of authors was incomplete with the inadvertent omission of Oscar L. Acevedo's name and should read as follows: George I. Birnbaum, *2a Jean Robert Brisson, 2a Steven H. Krawczyk, 2b Oscar L. Acevedo, 2b and Leroy B. Townsend. 2b

Deuterium Kinetic Isotopic Effects in the 1,4-Dimethylenecyclohexane Boat Cope Rearrangement [J. Am. Chem. Soc. 1986, 108, 468]. JOSEPH J. GAJEWSKI* and JOSE LEONARDO JIMENEZ

Page 469, Table III: The heading should be C not CD. Page 470, scheme below the first paragraph:

$$[K^{H}/K^{D}](1,2) = 1.16 (0.04) \rightarrow 1.16 \text{ not } 1/1.16$$

 $[K^{H}/K^{D}](1,3) = 1.04 \text{ not } 1/1.04$

Page 470, 2nd paragraph lines 15 and 16: TND(1) not TND(2).

Page 470, scheme below 2nd paragraph: interchange TND and TXD.

Page 471, Figure 1: interchange TND and AD.

These are typographical errors and do not affect the analyses derived from the data in Figures 3 and 4.

We thank Professor R. Magid for bringing these errors to our attention.

A New, Easily Accessible Reciprocal Chiral Stationary Phase for the Chromatographic Separation of Enantiomers [J. Am. Chem. Soc. 1986, 108, 352–354]. W. H. PIRKLE* and THOMAS C. POCHAPSKY

Page 354. The structure of the 19th entry in Table I was incorrectly drawn as

The proper structure is