## ADDITIONS AND CORRECTIONS

## 2000, Volume 104A

**István Z. Kiss,\* Géza Mándi, Mihály T. Beck\*:** Artificial Neural Network Approach to Predict the Solubility of  $C_{60}$  in Various Solvents

Page 8087. The first sentence of the third paragraph in the Discussion section should cite ref 4 instead of ref 14.

10.1021/jp003302w Published on Web 10/28/2000

## 2000, Volume 104A

**Jon Applequist:** Electronic Normal Modes and Polarization Waves in Translational Polymer Helices. Application to Fully Extended Poly[(R)- $\beta$ -aminobutyric acid] Chains

Page 7135. Equation 16 should read

$$\tau_{m1}^{(n)} = c_1 \cos(mk_n a + \delta) \tag{16}$$

10.1021/jp003423w Published on Web 10/21/2000

## 2000, Volume 104A

N. Bellec, K. Boubekeur, R. Carlier, P. Hapiot\*, D. Lorcy, and A. Tallec: Controlling the Conformation Changes Associated to Electron Transfer Steps through Chemical Substitution: Intriguing Redox Behavior of Substituted Vinylogous TTF

Page 9750. The paper entitled Controlling the Conformation Changes Associated to Electron Transfer Steps through Chemical Substitution: Intriguing Redox Behavior of Substituted Vinylogous TTF includes supporting information. This information is described as drawings of the optimized geometries at the B3LYP/6-13G\* level with the corresponding cartesian coordinates followed by the total atomic charges or total spin densities for open-shell calculations in the case of radicals, the values of the energies in hartrees, and S\*\*2 showing the spin contaminations for the neutral, cation radicals, and dications of the investigated TTFs. This material is available free of charge via the Internet at http://pubs.acs.org.

10.1021/jp003725y Published on Web 10/25/2000