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Errata

- Bellamy F.D. and Ou K., 25,839-842(1984): corrected procedure for reduction of p-nitrobenzoic acid, 1362
- Berkowitz W.F., Perumattam J., and Amarasekara A., 26, 3665(1985): acknowledgement of similar work published in Chem. Letters, 4562
- Brinkman K. and Helquist P., 26, 2845(1985): formula, last product in Table on p.2846 corrected and reference 8a replaced, 4154
- Danikiewicz W. and Makosza M., 26, 3599(1985): scheme 1 which was omitted from p 3599, 6523
- Davies J.W., Malpas J.R. and Moss R.E., 26, 4533(1985): last word of second paragraph on p.4533 diazabicyclo; on p.4535 energy values corrected; revised Table 1 produced, 5226
- Deshmukh M., Dunach E. and Kagan H.B., 25, 3467(1984): Line 15 after Abstract - correction, 402
- Gamboni R., Mohr P., Waespe-Sarcevic N. and Tamm C., 26,203(1985): structure 1d is corrected, 3058
- Guo-qiang L., Hai-jian X., Bi-chi W., Guong-zhong G. and Wei-shan Z., 26, 1233(1985): reaction conditions for Schemes 1 and 2 corrected and arrangement of comps in Table 1 changed, 1690
- Jackson A.H., Rao K.R.N., Ooi N.S. and Adelakun E., 25, 6049(1984): p.6049 three corrections, 3522
- Kondo N., Imai K., Isobe M., Goto T., Murasugi A., Wada C., Nakagawa C. and Harashi Y., 25, 3869(1984): corrected structures for formulae 1 and 2, 690
- Mak C-P. and Filiri H., 26, 1433(1985): on p 1434 entry 8 corrected, 4010
- Newcomb M., Williams W.G. and Crumpacker E.L., 26, 1183(1985): yield of isolated bromide 1 corrected, 2727
- Otera J., Yoshinaga Y., Hirakawa K. and Nakata T., 26,3219(1985): entries 6 and 7 in Table 1 should be deleted, 3746
- Overman L.E. and Burk R.M.,

- line from bottom of p.5737, 4154
- Shirahama H., Arora G.S., Osawa E. and Matsumoto T., 24,2869-2872(1983): Formulas 10 and 11 on p.2871 corrected, 2
- Sunitha K., Balasubramanian K.K. and Rajagoplan K., 26, 4393: corrections in References and Notes, 5490
- Swindell C.S., de Solms S.J., 25,3801(1984): data listed for conformation B of structure 3b corrected, 1474
- Uma R., Swaminathan S. and Rajagopalan K.N., 25,5825-8 (1984): Sentence on lines 20-24, page 5827 rewritten, 1114
- Wakamiya T., Nakamoto H. and Shiba T., 25,4411(1984): chemical shift of ¹H NMR corrected, 2138

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