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Symposium-in-Print

POLYMER SUPPORTED CATALYSTS AND REAGENTS IN SYNTHETIC ORGANIC CHEMISTRY

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3635 Characterization of Mono- and Diaminopyrimidine Derivatives as Novel,
Nonpeptide Gonadotropin Releasing Hormone (GnRH) Receptor
Antagonists

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3641 Unique Hole-Trapping Property of the Degenerate Base, 2-Amino-7-dea-
zaadenine

3645 Erratum

I Instructions to Contributors