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Preface

This issue of Transactions on Computational Systems Biology contains a fully-refereed selection of papers from the Fourth International Conference on Computational Methods in Systems Biology, held in Edinburgh, Scotland, April 3–5, 2005. I would like to thank both the referees for all their hard work and also the CMSB 2005 programme committee for their help in choosing which papers to invite for submission.

June 2006

Gordon Plotkin
Program Chair
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