The ISMB conference series provides a general forum for disseminating the latest developments in bioinformatics. ISMB is a multidisciplinary conference that brings together scientists from computer science, mathematics, molecular biology and statistics. Its scope includes the development and application of advanced computational methods for biological problems. Relevant computational techniques include, but are not limited to: machine learning, pattern recognition, knowledge representation, databases, combinatorics, stochastic modeling, string and graph algorithms, linguistic methods, robotics, constraint satisfaction, and parallel computation. Biological areas of interest include biomolecular structure, genomics, biomolecular sequence analysis, evolution and phylogenetics, biomolecular interactions, metabolic pathways, regulatory networks, developmental control, and molecular biology generally. Emphasis is placed on the validation of methods using real data sets, on practical applications in the biological sciences, and on development of novel computational techniques.

Previous ISMB conferences were held at:
- 1993: National Library of Medicine, Bethesda, MD, USA
- 1994: Stanford University, Stanford, CA, USA
- 1995: Cambridge University, Cambridge, UK
- 1996: Washington University, St. Louis, MO, USA
- 1997: Halkidiki, Greece
- 1998: Montreal, Quebec, Canada

Meanwhile ISMB has established itself as one of the leading refereed international conference series in bioinformatics.

ISMB '99 has received 91 submissions of contributed papers out of which a program committee consisting of 47 distinguished researchers in the field of bioinformatics has selected 34 papers to be presented at the conference and included in these proceedings. In the rigorous selection process each paper was reviewed by at least three members of the program committee and, after all the referee's reports had been completed, an extensive web-based discussion of all papers has taken place. The accepted papers have authors from 12 countries and 4 continents.

The contributed papers presented in these proceedings are only one part of the strong scientific program of the ISMB conferences. Another very important part of the conference is its set of invited talks. ISMB '99 expanded the number of invited talks from previous ISMB conferences from three to eight. Four invited talks were arranged to inject new problems and ideas from molecular biology and emerging experimental technologies into the bioinformatics community. These talks were given by Manfred Eigen, from the Max-Planck Institute for Biophysical Chemistry in Göttingen, Germany, Rob Lipshutz from Affymetrix Inc., Santa Clara, California, Matthias Mann from Odense University, Denmark and Anthony Kerlavage from Celera Genomics Corporation in Rockville, Maryland. Four other talks concentrated on bioinformatics issues. These talks were given by Amos Bairoch from the University of Geneva, Switzerland, Richard Karp from the University of Washington in Seattle, Washington, Eugene Koonin from the National Center for Biotechnology Information (NCBI), Bethesda, Maryland, and Michael Sternberg from the Imperial Cancer Research Fund (ICRF) London, UK. Short abstracts of all invited talks are contained in these proceedings.

Ten tutorials were given on the first day of the conference (August 6, 1999):
- Pierre Baldi: Probabilistic Graphical Models
- Douglas Brutlag: Bioinformatics and Molecular Biology
- Kevin Karplus, Melissa Cline, Christian Barrett: Getting the Most out of Hidden Markov Models
- Arthur Lesk: Sequence-Structure Relationships and Evolutionary Structure Changes in Proteins
- Rob Miller, Alan Christoffels, Winston Hide: EST Clustering
- Martin Reese, Suzanna Lewis: The Challenge of Annotating a Complete Eukaryotic Genome: A Case Study in Drosophila melanogaster
- David States, Brian Dunford Shore: PERL Abstractions for Databases and Distributed Computing
- Zoltan Szallasi: Genetic Network Analysis—From the Lab Bench to Computers and Back
- Tandy Warnow, Junhyong Kim: Computational and Statistical Challenges Involved in Reconstructing Evolutionary Trees
- Thomas Werner: The Biology and Bioinformatics of Regulatory Regions in Genomes

Furthermore, the conference included a lively poster session, an industrial exhibition of bioinformatics software, and a job fair. BIOPERL '99 took place in Heidelberg on August 5, 1999 directly before ISMB '99. Information on this workshop can be found on its web site (http://bio.perl.org/bioperl-99/). ISMB '99 directly preceded a total solar eclipse that happened just south of Heidelberg on August
The planning and organization of a conference such as ISMB ’99 with the commitment to publishing a high quality refereed proceedings in a short time frame relies extensively on the enthusiasm and commitment of the program committee. The organizing committee wishes to formally express their gratitude to all members of the program committee.

In conclusion the organizing committee would like to thank Ms. Ingeborg Fatscher in Heidelberg for her exceptional engagement in all on-location aspects of the conference preparation. Thanks go also to Hannelore Napieraj at GMD, for secretarial help, Ingrid Filter and Theo Mevissen at GMD for help with the web presentation of the conference, Ulrich Müller, Katharina Rusch, and Karina Schygulla at LION bioscience AG for graphics design, and Mike Hamilton at AAAI Press for once again producing a proceedings of excellent quality.

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Peer Bork
Douglas Brutlag
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Hans-Werner Mewes
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ISMB ’99 Organizing Committee