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Edited by Lior Pachter and Bernd Sturmfels
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Algebraic Statistics for Computational Biology

“If you can’t stand algebra, keep out of evolutionary biology”

– John Maynard Smith
[Smith, 1998, page ix]

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Edited by
Lior Pachter and Bernd Sturmfels
University of California at Berkeley



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Preface

The title of this book reflects who we are: a computational biologist and an algebraist who share a common interest in statistics. Our collaboration sprang from the desire to find a mathematical language for discussing biological sequence analysis, with the initial impetus being provided by the introductory workshop on *Discrete and Computational Geometry* at the Mathematical Sciences Research Institute (MSRI) held at Berkeley in August 2003. At that workshop we began exploring the similarities between tropical matrix multiplication and the Viterbi algorithm for hidden Markov models. Our discussions ultimately led to two articles [Pachter and Sturmfels, 2004a,b] which are explained and further developed in various chapters of this book.

In the fall of 2003 we held a graduate seminar on *The Mathematics of Phylogenetic Trees*. About half of the authors of the second part of this book participated in that seminar. It was based on topics from the books [Felsenstein, 2003, Semple and Steel, 2003] but we also discussed other projects, such as Michael Joswig's polytope propagation on graphs (now Chapter 6). That seminar got us up to speed on research topics in phylogenetics, and led us to participate in the conference on *Phylogenetic Combinatorics* which was held in July 2004 in Uppsala, Sweden. In Uppsala we were introduced to David Bryant and his statistical models for split systems (now Chapter 17).

Another milestone was the workshop on *Computational Algebraic Statistics*, held at the American Institute for Mathematics (AIM) at Palo Alto in December 2003. That workshop was built on the algebraic statistics paradigm, which is that statistical models for discrete data can be regarded as solutions to systems of polynomial equations. Our current understanding of algebraic statistical models, maximum likelihood estimation and expectation maximization was shaped by the excellent discussions and lectures at AIM.

These developments led us to offer a mathematics graduate course titled *Algebraic Statistics for Computational Biology* in the fall of 2004. The course was attended mostly by mathematics students curious about computational biology, but also by computer scientists, statisticians, and bioengineering students interested in understanding the mathematical foundations of bioinformatics. Participants ranged from postdocs to first-year graduate students and even one undergraduate. The format consisted of lectures by us on basic principles

of algebraic statistics and computational biology, as well as student participation in the form of group projects and presentations. The class was divided into four sections, reflecting the four themes of algebra, statistics, computation and biology. Each group was assigned a handful of projects to pursue, with the goal of completing a written report by the end of the semester. In some cases the groups worked on the problems we suggested, but, more often than not, original ideas by group members led to independent research plans.

Halfway through the semester, it became clear that the groups were making fantastic progress, and that their written reports would contain many novel ideas and results. At that point, we thought about preparing a book. The first half of the book would be based on our own lectures, and the second half would consist of chapters based on the final term papers. A tight schedule was seen as essential for the success of such an undertaking, given that many participants would be leaving Berkeley and the momentum would be lost. It was decided that the book should be written by March 2005, or not at all.

We were fortunate to find a partner in Cambridge University Press, which agreed to work with us on our concept. We are especially grateful to our editor, David Tranah, for his strong encouragement, and his trust that our half-baked ideas could actually turn into a readable book. After all, we were proposing to write to a book with twenty-nine authors during a period of three months.

The project did become reality and the result is in your hands. It offers an accurate snapshot of what happened during our seminars at UC Berkeley in 2003 and 2004. Nothing more and nothing less. The choice of topics is certainly biased, and the presentation is undoubtedly very far from perfect. But we hope that it may serve as an invitation to biology for mathematicians, and as an invitation to algebra for biologists, statisticians and computer scientists. Following this preface, we have included a guide to the chapters and suggested entry points for readers with different backgrounds and interests. Additional information and supplementary material may be found on the book website at <http://bio.math.berkeley.edu/ascb/>

Many friends and colleagues provided helpful comments and inspiration during the project. We especially thank Elizabeth Allman, Ruchira Datta, Manolis Dermitzakis, Serkan Hoşten, Ross Lippert, John Rhodes and Amelia Taylor. Serkan Hoşten was also instrumental in developing and guiding research which is described in Chapters 15 and 18.

Most of all, we are grateful to our wonderful students and postdocs from whom we learned so much. Their enthusiasm and hard work have been truly amazing. You will enjoy meeting them in Part II.

Lior Pachter and Bernd Sturmfels
Berkeley, California, May 2005

Guide to the chapters

The introductory Chapters 1–4 can be studied as a unit or read in parts with specific topics in mind. Although there are some dependencies and shared examples, the individual chapters are largely independent of each other. Suggested introductory sequences of study for specific topics are:

- Algebraic statistics: 1.1, 1.2, 1.4, 1.5.
- Maximum likelihood estimation: 1.1, 1.2, 1.3, 3.3.
- Tropical geometry: 2.1, 3.4, 3.5.
- Gröbner bases: 3.1, 3.2, 2.5.
- Comparative genomics: 4.1, 4.2, 4.3, 4.4, 4.5, 2.5.
- Sequence alignment: 1.1, 1.2, 1.4, 2.1, 2.2, 2.3.
- Phylogenetics: 1.1, 1.2, 1.4, 2.4, 3.4, 3.5, 4.5.

Dependencies of the Part II chapters on Part I are summarized in the table below. This should help readers interested in reading a specific chapter or section to find the location of background material. Pointers are also provided to related chapters that may be of interest.

Chapter	Prerequisites	Further reading
5	1.4, 2.2, 2.3	6, 7, 8, 9
6	1.1, 1.2, 1.4, 2.3	5, 7, 8, 9
7	2.2, 2.3	8
8	1.1, 1.2, 1.4, 2.1, 2.2, 2.3	5, 7, 9
9	1.5, 2.2, 2.3, 4.4	5, 8
10	1.1, 1.2, 1.4	9, 11
11	1.1, 1.2, 1.3, 3.1, 3.2	12
12	1.3, 1.4	4.4, 11
13	1.1, 1.2, 1.4, 1.5	22
14	1.1, 1.2, 1.4, 1.5, 3.1	11, 16
15	1.4, 3.1, 3.2, 3.3, 4.5	16, 17, 18, 19, 20
16	1.4, 3.1, 3.2, 4.5	15, 19
17	1.1, 1.2, 1.4, 1.5, 2.4, 4.5	15, 18, 19
18	2.4, 4.5	20
19	2.4, 3.1, 4.5	15, 18
20	2.4, 4.5	17
21	1.4, 2.5, 4.5	17, 19
22	1.4, 4	7, 13, 21

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