Introduction to Protein–DNA Interactions

Structure, Thermodynamics, and Bioinformatics
Also from Cold Spring Harbor Laboratory Press

Other Titles of Interest

Bioinformatics: Sequence and Genome Analysis, Second Edition
Genes & Signals
A Genetic Switch, Third Edition: Phage Lambda Revisited
To my parents, Milo and Claryce, who gave me the love of learning and the encouragement to follow wherever that led.

To my wife, Susan Dutcher, and my children, Ben and Adrienne, who have enriched my life immeasurably.
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The biological importance of protein–DNA interactions has been recognized since the early 1960s, starting with the discovery by Jacob and Monod of the lac operon and its regulation in *Escherichia coli*. In the intervening 50 years, studies of protein–DNA interactions have made significant contributions to most areas of molecular, cellular, and developmental biology. A wide range of approaches has been applied in those studies, but they can be broadly classified into the three types that are the focus of this book: structural, thermodynamic, and bioinformatic. The earliest studies used biochemical and biophysical methods to analyze the thermodynamic and kinetic aspects of protein–DNA interactions. The first binding site sequences were determined in the early 1970s, which led to hypotheses about recognition mechanisms and the information required for regulatory systems to function. Technological advances in the late 1970s and the early 1980s, including the ability to sequence and synthesize DNA and to clone, express, and purify large quantities of proteins, facilitated many new types of studies. The earliest bioinformatics approaches were developed in the late 1970s, as soon as there were enough sequences for statistical analyses to be worthwhile. Shortly after that, as it became much easier to synthesize and purify sufficient quantities of specific proteins and DNA sequences of interest, structural studies rapidly increased. Further technological advances in the last two decades have continued to accelerate the pace of discovery. Most important have been further efficiencies in DNA sequencing that have resulted not only in whole-genome sequences for many species but also whole-genome and mRNA sequences from individuals as well as a variety of other sequence-based data sets. Our understanding of protein–DNA interactions and their roles in a wide range of biological processes has grown enormously, but there is still much we do not know and the field continues to be ripe for further discovery.

The primary goal of this book is to provide an introduction to protein–DNA interactions that bridges the three classes of approaches. Experts in any of the fields are not
likely to learn anything new within their field; in fact, they will undoubtedly find examples of details being glossed over in favor of a simplified presentation. But experts in one area tend to have more cursory knowledge of the other fields and thus may learn from other sections of the book. Those who are new to the study of protein–DNA interactions or those outside the field with a casual interest in the topic may gain new insights throughout the book. If so, the book has succeeded even beyond the fact that I learned something in the process of writing every chapter.

The regulation of gene expression has fascinated me since my graduate school days. I have ventured into other topics, mostly related to how computer programs can help to uncover biological knowledge, but the majority of my efforts have been focused on understanding how networks of transcription factors regulate gene expression and control cell fates and phenotypes. I have been extremely fortunate to have been associated throughout my career with teachers and students, colleagues and collaborators, and most of all friends who have taught and encouraged me and made my whole adventure enjoyable. The list of those who made significant contributions to my research, many of whom I have never met but have benefited from immensely through reading their papers, is too long to include in this preface. But a few have had such a large influence that I must thank them here. Larry Gold, my graduate and postdoc advisor, kept research always fun and gave me the freedom and encouragement to follow an unconventional path. Tom Schneider, a fellow student in Larry’s lab, and Andrej Ehrenfeucht, a mentor in all things computational, were there from the beginning and opened my eyes to new horizons that I would have missed without them. I have had many great collaborators over the years but special thanks go to John Heumann, Alan Lapedes, and Charles “Chip” Lawrence, each of whom has filled gaps in my knowledge and provided numerous insights into my own work that were initially invisible to me. I have also had many great students and postdocs who made progress possible and who taught me at least as much as I taught them.

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