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Molecular Biology and Translational Science

Modifications of Nuclear DNA and its Regulatory Proteins

edited by

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
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Contents

Contributors.....	xi
Preface.....	xiii
Introduction—Epiphanies in Epigenetics	1
Xiaodong Cheng and Robert M. Blumenthal	
I. What is an Epigenetic Code?	2
II. Inheritance.....	6
III. Modularity of Epigenetic Modifiers.....	7
References	14
Section I Evolution of DNA Methylation	23
Natural History of Eukaryotic DNA	
Methylation Systems	25
Lakshminarayan M. Iyer, Saraswathi Abhiman, and L. Aravind	
I. Introduction	27
II. DNA Methyltransferases	30
III. 5mC Demethylation and Potential DNA Demethylases.....	53
IV. Further Modifications of 5mC in Eukaryotic DNA.....	63
V. Domains Involved in Discrimination of Methylated Versus Nonmethylated Cytosines in DNA	69
VI. Domain Architectural Logic of Proteins Related to DNA Methylation	81
VII. Evolutionary Considerations	85
VIII. General Conclusions	88
References	90

Natural History of the Eukaryotic Chromatin Protein Methylation System	105
L. Aravind, Saraswathi Abhiman, and Lakshminarayan M. Iyer	
I. Introduction	107
II. The Categories of Protein Methylases and Their Role in Chromatin Protein Methylation	111
III. Enzymatic Mechanisms That Preempt or Reverse the Action of Protein Methylases in Chromatin	133
IV. Domains Involved in Discrimination of Methylated Peptides	147
V. Domain Architectures of Enzymes in the Chromatin Protein Methylation Network	154
VI. Evolutionary Considerations	159
VII. General Conclusions	162
References	163
 DNA Methylation in <i>Drosophila</i> —A Critical Evaluation	 177
Veiko Krauss and Gunter Reuter	
I. Evidence for 5-Methyl Cytosine in <i>Drosophila</i>	178
II. Evidence for DNA Methylation Activity of Dnmt2 Enzymes	180
III. Methylation of tRNA by Dnmt2	182
IV. Biological Function of Dnmt2 in <i>Drosophila</i>	184
V. Dinucleotide Abundances in “Dnmt2-only” Organisms: A Hint of DNA Methylation?	185
VI. Methyl-Binding Proteins in <i>Drosophila</i> —Evidence, Concepts, and Inconsistencies	187
References	188
 DNA Methylation in Zebrafish	 193
Mary G. Goll and Marnie E. Halpern	
I. Introduction	194
II. Methylation Profile of the Zebrafish Genome	195
III. The Zebrafish Methylation Machinery	197
IV. Link Between DNA and Histone Modification in Zebrafish	207
V. DNA Methylation Dynamics in Early Development	208

CONTENTS	vii
VI. Evidence for Active DNA Demethylation in Zebrafish	209
VII. Perspectives and Future Directions	211
VIII. Conclusions	213
References	214

Section II Mammalian DNA Methyltransferases – Structure and Function 219

Dnmt1: Structure and Function	221
--	------------

Željko M. Svedružić

I. Introduction	222
II. The Functional Domains of Dnmt1 and Dnmt1 Inhibitors	224
III. Structure and Function of Dnmt1 and Its Interaction with Other Molecules	236
IV. New Crystal Structures of Large C-Terminal Fragment of Mouse and Human Dnmt1	243
References	245

The DNMT3 Family of Mammalian <i>De Novo</i> DNA Methyltransferases	255
--	------------

Frédéric Chédin

I. Introduction	256
II. Discovery, Primary Structure, and Expression of Mammalian DNMT3A and DNMT3B	257
III. DNMT3A and DNMT3B Drive the Establishment of DNA Methylation Patterns <i>De Novo</i>	259
IV. The DNMT3L Protein, an Essential Accessory Factor for <i>De Novo</i> Methylation	269
V. Interactions Between the <i>De Novo</i> DNA Methylation Machinery and Chromatin	274
VI. Concluding Remarks and Outstanding Questions	277
References	277

Section III DNA Methylation and Demethylation. 287

Recruitment of Dnmt1: Roles of the SRA Protein Np95 (Uhrf1) and Other Factors. 289

Jafar Sharif and Haruhiko Koseki

I. Roles for DNA Methylation in Transcriptional Regulation and Development.....	290
II. Maintenance and <i>De Novo</i> DNA Methyltransferases: Two Modes of Action.....	291
III. Localization of Dnmt1 into Hemimethylated DNA: Search for Recruiter Molecule(s).....	292
IV. Np95 Is a Cell Cycle-Associated Protein That Localizes to the RF During S-Phase.....	293
V. Np95 Protein Domains: SRA and More.....	294
VI. Np95 Recruits Dnmt1 into Hemimethylated Sites During DNA Replication.....	296
VII. Np95 Is Required for Global DNA Methylation.....	298
VIII. Other Factors That Recruit/Interact with Dnmt1.....	299
IX. Concluding Remarks and Future Directions.....	305
References.....	307

Regulation of Expression and Activity of DNA (Cytosine-5) Methyltransferases in Mammalian Cells . . . 311

Shannon R. Morey Kinney and Sriharsa Pradhan

I. Transcriptional and Posttranscriptional Regulation of DNMTs.....	312
II. Regulation of DNMTs by Posttranslational Modifications.....	317
III. Altered Regulation of DNMTs During Disease.....	320
IV. Drug-Induced Reductions in DNMT Levels.....	326
V. Concluding Remarks and Future Directions.....	328
References.....	329

Mechanistic and Functional Links Between Histone Methylation and DNA Methylation 335

Taiping Chen

I. Introduction.....	335
II. An Evolutionarily Conserved Pathway Between H3K9 Methylation and DNA Methylation.....	337
III. A Role for LSD1 in Coordinating Histone and DNA Methylation?.....	339

CONTENTS	ix
IV. H3K4 Demethylation and Genomic Imprinting	341
V. Concluding Remarks	343
References	344

Section IV DNA Methylation and Silencing 349

RNA-Mediated Silencing Mechanisms in Mammalian Cells 351

Anton Wutz

I. Introduction.....	352
II. RNA Constitutes a Structural Component of the Mammalian Cell Nucleus	352
III. A Role for RNAs in Regulating Chromatin Modifications and Organization.....	353
IV. Repression of Ribosomal DNA Repeats by the NoRC Repressor Complex Is Regulated by IGS Noncoding RNAs	364
V. Pervasive Transcription: RNA at Centromeric and Telomeric Chromatin	366
VI. Aspects of the Evolution of Regulatory RNAs.....	367
VII. Concluding Remarks and Future Directions in RNA Research	369
References.....	370

Biological Functions of Methyl-CpG-Binding Proteins . . . 377

Pierre-Antoine Defossez and Irina Stancheva

I. Introduction	378
II. Families of Methyl-CpG-Binding Proteins.....	379
III. Biological Functions of Methyl-CpG-Binding Proteins.....	386
IV. Methyl-CpG-Binding Proteins and Human Disease.....	389
V. Concluding Remarks	391
References	392

Section V DNA Methylation in Medicine 399

Diseases Associated with Genomic Imprinting 401

Jon F. Wilkins and Francisco Úbeda

I. Overview of Genomic Imprinting.....	402
II. Disorders Associated with Particular Imprinted Genes and Regions	406

III. Psychiatric Disorders and Other Behavioral Effects	422
IV. The Cost of Imprinting.....	426
V. Conclusions	431
References.....	431
DNA Methylation Changes in Cancer	447
Minoru Toyota and Eiichiro Yamamoto	
I. Introduction	447
II. Mechanism Underlying DNA Methylation Changes in Cancer	448
III. The Roles of DNA Methylation in Cancer Signaling Pathways	449
IV. Genetic and Epigenetic Interaction in Cancer	451
V. Hypomethylation in Cancer.....	451
VI. Epigenetic Therapy	453
VII. Conclusions and Future Directions.....	454
References.....	454
Genome-Wide Distribution of DNA Methylation at Single-Nucleotide Resolution	459
Eleanor Wong and Chia-Lin Wei	
I. Impact of Single-Nucleotide-Based Detection on DNA Methylome Profiling	460
II. Overview of Molecular Approaches Used for Methylation Studies	461
III. DNA Methylation Patterns at Single-Nucleotide Resolution	470
IV. DNA Methylation, Histone Modifications, and Other Epigenetic Regulation	472
V. Detection of the 6th Base (5-Hydroxymethylcytosine) and Future Perspectives	473
VI. Concluding Remarks.....	473
References.....	474
Index	479

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Preface

It hardly seems possible that we are approaching two decades since the discovery that DNA methyltransferases act by everting the target base out of the double helix Klimasauskas et al., HhaI methyltransferase flips its target base out of the DNA helix. *Cell* 1994 76:357–69. Since 1994, a very substantial body of work has emerged from the attempts to understand both the molecular mechanisms and biological roles of methylation, not only of DNA but also of the associated histone proteins.

As is typical of important biological questions, there is no good stopping point at which we can say that the basic answers are known. Nevertheless, we feel the time is right for a summing up that allows establishment of context. For that reason, we are delighted that colleagues who study chromatin methylation from a variety of perspectives have agreed to contribute to this volume.

This book is divided into five sections, following an introduction by us that is intended to consolidate the most basic background material into one chapter. The next five paragraphs should have appropriate section numbers.

Section I explores the evolution of the DNA- and histone-methylating (and demethylating) enzymes and of the biological roles of this methylation. The former is presented in two chapters by Iyer et al. and Aravind et al., while the latter involves chapters on chromatin methylation in *Drosophila* (Krauss and Reuter) and zebrafish (Goll and Halpern).

Section II is focused on the major mammalian DNA methyltransferases, Dnmt1 (Svedruzic) and Dnmt3/3L (Chédin). One of the take-home messages is that the division of labor between *de novo* and maintenance methylation is much less strict than had been thought earlier.

Section III examines the processes and control of DNA methylation and demethylation. There are chapters on recruitment of Dnmt1 (Sharif and Koseki) and modulation of its stability (Morey-Kinney and Pradhan), along with the functional linkages between demethylation of DNA and histones (Chen).

Section IV covers the association between DNA methylation and silencing of gene expression. This can occur via an RNA-directed process (Wutz) or the involvement of proteins bearing a methyl-binding domain (Defossez and Stancheva).

Finally, Section V describes biomedical and biotechnological aspects of these epigenetic processes. There are chapters on chromatin methylation and cancer (Toyota and Yamamoto) and diseases associated with imprinting

(Wilkins and Úbeda). The section, and the volume, is brought to a close by a description of the methods to determine, and the implications of knowing, the distribution of DNA methylation at the single-nucleotide level (Wong and Wei).

We are happy that these chapters will be available online in addition to the print format, so it will be possible to update them periodically.

We would like to thank those who made this volume possible. First, of course, the authors of the various chapters, from whom we have learned so much. Second, Delsy Retchagar and the team at Elsevier, for laboring mightily to keep us on schedule. Third, our respective institutions and departments for encouragement and staff support (especially Suzanne Payne of the Department of Medical Microbiology and Immunology at the University of Toledo). Fourth, grant support that allows our ongoing research in this field—the U.S. National Institutes of Health (Grants GM-049245-17 and GM-068680-06 to X. C.) and the U.S. National Science Foundation (Grant MCB-0964728 to R. M. B.). Last, but by no means least, we thank our families and look forward to being able to spend a bit more time with them.

February, 2011

XIAODONG CHENG AND ROBERT M. BLUMENTHAL