_	\sim						
('HROMATIN	ARCHITECTURE	ARERRATIONS	IN PROSTATE	CANCER	ΔND	LEHKEMIA

by

James Hawley

A thesis submitted in conformity with the requirements for the degree of Doctor of Philosophy

Graduate Department of Medical Biophysics

University of Toronto

Chapter 1

Introduction

Cancer is one of the largest causes of death worldwide, ranking in the top ten most frequent causes in over 150 countries and most frequent in over 40 [1]. Disease treatment is complicated by the fact that cancers are a myriad of diseases with unique origins, symptoms, and treatment options, often related to the cell of origin [2]. However, numerous hallmarks of cancers have emerged over the last 50 years to provide understanding about what biological aberrations cause tumours to initiate, how they develop over time, and how they respond to the the repetition interventions [3–6] (Figure 1.1).

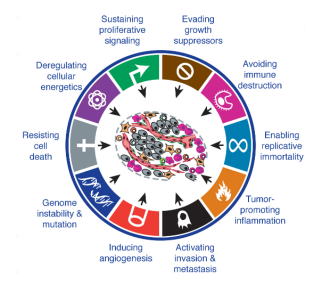


Figure 1.1: The hallmarks of cancer. Adapted from [REF 4].

Many of these hallmarks of cancer can be achieved through aberrations to the genome and the molecular machinery that enables cells to function normally [7]. For example, genome instability can be achieved by inhibiting DNA repair machinery, as is observed with abnormalities in *MLH1*

and MSH2 repair genes in colorectal cancers [8] or mutations to BRCA1, BRCA2, and ATM genes in prostate cancer (PCa) [9]. Similarly, replicative immortality can be achieved through telomere elongation by over-expression of the TERT gene [10]. Mutations to the TERT promoter, resulting in its over-expression, were first identified in melanomas [11, 12], but have since been further identified in bladder, thyroid, and brain cancers [10, 13, 14]. But while cancer has long been viewed as a disease of the genome [3, 7], there are many avenues cells can take to arrive these hallmarks resulting from aberrations of how genes are expressed inside the cell nucleus.

1.1 Normal chromatin architecture in mammalian cells

Genes, encoded as DNA, are expressed by being transcribed into messenger RNA (mRNA) and subsequently translated into proteins in the process known as the Central Dogma of molecular biology [15] (Figure 1.2a). The transcription of genes into mRNA requires RNA polymerase to bind at transcription start sites (TSSs) within DNA elements found at the beginning of genes, termed promoters [16]. Promoters are one example of a class of DNA elements, termed *cis*-regulatory elements (CREs) because of their roles in regulating the expression of genes on the same strand of DNA. The recruitment of RNA polymerase is aided by a special class of proteins, termed transcription factors (TFs), that can bind at DNA sequences either close to a gene's promoter, or far from it at other CREs such as enhancers and insulators [17–22] (Figure 1.2b). Together, the binding of TFs to the DNA at specific CREs is fundamental for to initiating transcription and expressing genes.

1.1.1 DNA elements and features regulating transcription

The ability of TFs to bind at specific CREs is dependent on multiple features of the DNA. Many TFs bind to DNA at specific sequences, termed motifs [18, 23]. The structural protein CCCTC-binding factor (CTCF) has a well-defined motif and binds to this sequence at thousands of locations across the human genome [24, 25]. Mutations to the sequence motif can alter CTCF's binding affinity for DNA, as is the case with many TFs [26–28]. Relying on more than just the genetic sequence, CTCF is also an example of a TF that is sensitive epigenetic features such as DNA methylation (DNAme), the addition of a methyl group to DNA nucleotides [29–33], as are DNA methyltransferases DNMT1, DNMT3A, and DNMT3B [34, 35]. TF binding to DNA can also be affected by the presence of other proteins at binding sites. TFs can bind in a combinatorial manner at the same location [18, 19, 23] or be blocked from binding altogether by the presence of nucleosomes, protein complexes that DNA winds around to make it compact in three-dimensional space [36, 37].

The collection of DNA, nucleosomes, DNA-bound transcription factors, and chemical modifications is defined as the chromatin, and the presence and density of nucleosomes, as well as DNA coiling, make certain segments of the chromatin more or less accessible for TF binding (euchromatin and heterochromatin, respectively). This can affect normal cellular behaviour such as cell-type-specific gene expression [38, 39] and DNA damage repair in inaccessible regions [40]. Thus, both genetic and epigenetic chromatin features affect how TFs can bind and regulate transcription.

In addition to TF binding, transcription regulation depends on the ability of CREs to localize together in three-dimensional space across large genomic distances [41–43] (Figure 1.2c). Localization of CREs tens to thousands of basepairs (bps) apart form focal interactions is aided by the formation of topologically associated domains (TADs), domains of chromatin whose boundaries are linked by structural proteins, including CTCF and cohesin [22, 44–46]. In addition to TADs which can range in size from $10^4 - 10^6$ bp, chromatin is also organized into active or inactive compartments (A and B compartments, respectively) that range in size from $10^5 - 10^6$ bp [22, 47–49]. These two modes of chromatin organization facilitate the proper localization of CREs and TFs at the right time. While TADs and compartments are largely conserved across cell types [25, 50, 51], focal chromatin interactions can differ up to 45 % between cell types, providing a further mechanism to change chromatin state [48]. Different chromatin states enable cells with the same DNA sequence to express genes differently [17, 19, 44, 52–54], and thus identifying the repertoire of CREs, chromatin interactions, TADs, and compartments are vital in determining the regulation of genes in various cell types.

1.1.2 Methods for identifying DNA elements and chromatin interactions

High throughput sequencing protocols have enabled the characterization of functional elements from across the genome and rely on a similar concept to do so. This concept is to take a molecular feature of interest, be it an RNA transcript or nucleosome position, associate it with a short fragment of DNA, sequence these DNA fragments, and map it to the reference genome to identify where the original molecules came from (Figure 1.3). RNA sequencing (RNA-seq) methods reverse transcribed RNA into DNA that map back to individual genes, with the abundance of fragments indicating how much the gene is expressed [55]. Protein binding sites and histone post-translational modifications can be identified by fragmenting DNA around antibodies that bind to these proteins with techniques like chromatin immunoprecipitation sequencing (ChIP-seq) and cleavage under targets and release using nuclease (CUT&RUN) [56–58]. Accessible and inaccessible chromatin can be assessed by the chromatin's propensity to be cut by enzymes like DNase I, Tn5 transposase, and micrococcal nucle-

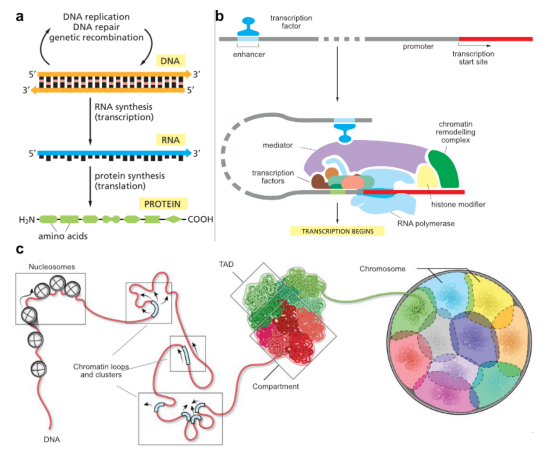


Figure 1.2: **The basics of gene expression inside the nucleus. a.** The central dogma of molecular biology. Adapted from [REF 15]. **b.** Schematic of the transcription machinery to initiate transcription. Adapted from [REF 15]. **c.** The scale of chromatin interactions across length scales. Adapted from [REF 46].

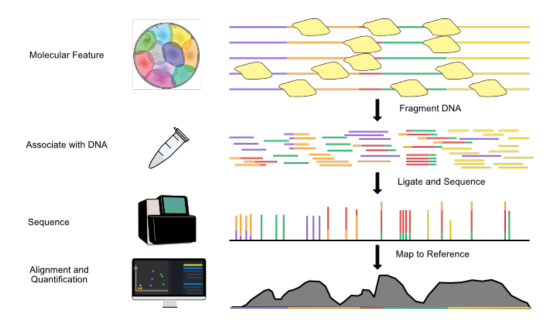


Figure 1.3: Characterizing functional DNA elements with high throughput sequencing.

ase in DNase I hypersensitive sequencing (DNase-seq), assay for transposase-accessible chromatin sequencing (ATAC-seq), and microccocal nuclease sequencing (MNase-seq) protocols, respectively [59–63]. DNAme can be measured with bisulfite-sequencing assays [64], and distal chromatin interactions can be identified with chromatin conformation capture (3C) and 3C-based methods such as Hi-C [25, 47, 48, 65, 66]. Yet while these measurements help in identifying candidate CREs and important regions of the genome, determining their function and which target genes they regulate is a further complicating problem.

Varying chromatin states across cell types means that multiple measurements across multiple cell types are necessary to understand the breadth of functions a single CRE may have. In 2007, the ENCODE Project aimed to catalogue all biochemically functional elements in the human genome to better understand all the ways genes are expressed and how they are regulated in different cell types [67, 68]. Using these genome-wide sequencing techniques across a variety of human cell lines and tissues, the ENCODE Project has since catalogued nearly 10⁶ candidate CREs, comprising nearly 8 % of the human genome [68]. Interpreting this data requires computational methods to correlate and interpret measurements across samples. Genome segmentation methods such as ChromHMM [69] and Segway [70, 71] classify genomic regions according to their predicted function which can be validated with in vitro or in vivo experiments. Many techniques for experimental validation, including clustered regularly interspaced short palindromic repeat (CRISPR)-Cas9, small interfering RNA (siRNA), and small hairpin RNA (shRNA), can interfere with candidate CREs by deleting them

from the genome, preventing TFs from binding to the chromatin, or preventing translation of mRNA transcripts into proteins [72, 73]. These same techniques can also be used to screen for candidate CREs themselves, through massively-parallel reporter assays (MRPAs) and CRISPR screens [73], necessitating their own suite of statistical and software tools for analyzing observations. Altogether, a collection of experimental and computational techniques enable the cataloguing and interpretation of thousands of CREs and chromatin interactions across many cell types. These catalogues facilitate understanding how genes are expressed within the complex chromatin architecture in normal cells and, importantly, how aberrations to this architecture can result in disease.

1.2 Aberrations to chromatin architecture in cancer

1.2.1 Genetic aberrations in cancer

Various types of mutations

single nucleotide variants (SNVs), copy number variants (CNVs), structural variants (SVs)

Coding and non-coding mutations

Impact of coding and non-coding mutations

1.2.2 Non-genetic aberrations in cancer

DNA methylation aberrations

Histone modifications

Chromatin accessibility and nucleosome positioning

Enhancer hijacking and chromatin organization aberrations

1.3 Interactions between chromatin architecture components in prostate cancer and leukemia

1.4 Dissertation structure

I begin with ?? by exploring the cis-regulatory landscape of PCa and delineating the CREs of the prostate oncogene FOXA1. I demonstrate the essentiality of FOXA1 for prostate tumours,

identify putative CREs based on integration of multiomic datasets in PCa cell lines, and assess the functional impact of recurrent PCa SNVs on *FOXA1* expression and TF binding.

With the *cis*-regulatory network of *FOXA1* established in PCa, I attempt to construct the *cis*-regulatory landscape genome-wide in PCa with 3C mapping in ??. Using Hi-C, I characterize the three-dimensional chromatin organization of PCa and investigate changes to this structure over oncogenesis, and explore the relationship between chromatin organization, SVs, and CRE hijacking.

In assessing the impact of SVs on chromatin organization, I uncovered a statistical problem stemming from the lack of recurrent SVs across PCa patients, leading to unbalanced experimental comparisons. To address this problem, I developed a statistical method for reducing error in gene expression fold-change estimates from unbalanced experimental designs in ?? and characterize the method.

Given the shared importance of mutations to TFs and epigenetic enzymes in prostate cancer and leukemias, in ?? I explore the epigenetic landscape of B-cell acute lymphoblastic leukemia (B-ALL) and its relapse after treatment. I characterize molecular changes to B-ALL tumours over the course of disease relapse and identify important changes to DNAme that indicate the reversion to a stem-like phenotype, often present in a subpopulation of cells at diagnosis.

Together, this dissertation investigates the multiple layers of the chromatin architecture that contribute to oncogenesis and cancer progression. I demonstrate that aberrations to the genome, epigenome, and three-dimensional organization of chromatin play important roles individually, and together, in the orchestration of the disease.

Glossary

3C chromatin conformation capture

AML acute myeloid leukemia

ANOVA Analysis of Variance

 \boldsymbol{AR} and rogen receptor

ATAC-seq assay for transposase-accessible chromatin sequencing

B-ALL B-cell acute lymphoblastic leukemia

bp basepair

cDNA complementary DNA

ChIP-seq chromatin immunoprecipitation sequencing

 ${f CLL}$ chronic lymphocytic leukemia

CMP common myeloid progenitor

CNV copy number variant

CPC-GENE Canadian Prostate Cancer Genome Network

 \mathbf{CpG} CG dinucleotide

crRNA CRISPR RNA

CRE cis-regulatory element

CRISPR clustered regularly interspaced short palindromic repeat

CTCF CCCTC-binding factor

GLOSSARY

CUT&RUN cleavage under targets and release using nuclease

DEPMAP Cancer Dependency Map

DHS DNase I hypersensitive sites

DMR differentially methylated region

DNAme DNA methylation

DNase-seq DNase I hypersensitive sequencing

dRI disease relapse-initiating

 $\mathbf{D}\mathbf{x}$ diagnosis

EarlyProB early progenitor B cell

 \mathbf{FDR} false discovery rate

FN false negative

FP false positive

FOX forkhead box

 ${f GLM}$ generalized linear model

 $\mathbf{GMP} \ \ \mathbf{granulocyte\text{-}macrophage} \ \mathbf{progenitor}$

GO gene ontology

 \mathbf{gRNA} guide RNA

HSC hematopoietic stem cell

HSPC hematopoietic stem and progenitor cell

IID independent and identically distributed

JS James-Stein

kbp kilobase

KO knockout

LDA limiting dilution assay

GLOSSARY 10

LMPP lymphoid-primed multi-potent progenitor

MeCapSeq DNA methylation capture sequencing

 \mathbf{MEP} megakaryocyte-erythrocyte progenitor

MNase-seq microccocal nuclease sequencing

MSE mean square error

mCRPC metastatic castration-resistant prostate cancer

MDS myelodisplastic syndrome

MLP monocyte-lymphoid progenitor

MPP multi-potent progenitor

MRPA massively-parallel reporter assay

NSG NOD scid gamma

OLS ordinary least squares

 \mathbf{mRNA} messenger RNA

PCa prostate cancer

PDX patient-derived xenograft

 $\mathbf{PreProB}$ pre-progenitor B cell

 \mathbf{ProB} progenitor B cell

Rel relapse

RNAi RNA interference

RNA-seq RNA sequencing

shRNA small hairpin RNA

siRNA small interfering RNA

 \mathbf{SNV} single nucleotide variant

SRA Sequence Read Archive

GLOSSARY 11

 ${f SNF}$ similarity network fusion

 ${f SV}$ structural variant

TAD topologically associated domain

TCGA The Cancer Genome Atlas

TSS transcription start site

 ${f TN}$ true negative

 ${f TP}$ true positive

 ${f TF}$ transcription factor

 ${f traceRNA}$ trans-activating CPRISR RNA

 ${f UTR}$ untranslated region

 \mathbf{WES} whole exome sequencing

 \mathbf{WGS} whole genome sequencing

 \mathbf{WT} wild-type

References

- Bray, F. et al. Global Cancer Statistics 2018: GLOBOCAN Estimates of Incidence and Mortality Worldwide for 36 Cancers in 185 Countries. en. CA: A Cancer Journal for Clinicians 68, 394–424. ISSN: 00079235 (Nov. 2018).
- Gilbertson, R. J. Mapping Cancer Origins. English. Cell 145, 25–29. ISSN: 0092-8674, 1097-4172 (Apr. 2011).
- 3. Hanahan, D. & Weinberg, R. A. The Hallmarks of Cancer. Cell 100, 57–70 (Jan. 2000).
- Hanahan, D. & Weinberg, R. A. A. Hallmarks of Cancer: The Next Generation. Cell 144, 646–674. ISSN: 1097-4172 (Electronic)\r0092-8674 (Linking) (Mar. 2011).
- Flavahan, W. A., Gaskell, E. & Bernstein, B. E. Epigenetic Plasticity and the Hallmarks of Cancer. Science 357, eaal2380—eaal2380 (July 2017).
- Pavlova, N. N. & Thompson, C. B. The Emerging Hallmarks of Cancer Metabolism. en. Cell Metabolism 23, 27–47. ISSN: 1550-4131 (Jan. 2016).
- Garraway, L. A. & Lander, E. S. Lessons from the Cancer Genome. English. Cell 153, 17–37.
 ISSN: 0092-8674, 1097-4172 (Mar. 2013).
- 8. Lengauer, C., Kinzler, K. W. & Vogelstein, B. Genetic Instabilities in Human Cancers. *Nature* **396**, 643–649 (Dec. 1998).
- Abeshouse, A. et al. The Molecular Taxonomy of Primary Prostate Cancer. en. Cell 163, 1011–1025. ISSN: 00928674 (Nov. 2015).
- Vinagre, J. et al. Frequency of TERT Promoter Mutations in Human Cancers. en. Nature Communications 4, 2185. ISSN: 2041-1723 (Oct. 2013).
- Huang, F. W. et al. Highly Recurrent TERT Promoter Mutations in Human Melanoma. en. Science 339, 957–959. ISSN: 0036-8075, 1095-9203 (Feb. 2013).

Horn, S. et al. TERT Promoter Mutations in Familial and Sporadic Melanoma. en. Science
 339, 959–961. ISSN: 0036-8075, 1095-9203 (Feb. 2013).

- Nagarajan, R. P. et al. Recurrent Epimutations Activate Gene Body Promoters in Primary Glioblastoma. Genome Research 24, 761–774 (May 2014).
- Stern, J. L., Theodorescu, D., Vogelstein, B., Papadopoulos, N. & Cech, T. R. Mutation of the TERT Promoter, Switch to Active Chromatin, and Monoallelic TERT Expression in Multiple Cancers. en. Genes & Development 29, 2219–2224. ISSN: 0890-9369, 1549-5477 (Nov. 2015).
- Alberts, B. Molecular Biology of the Cell Sixth edition. ISBN: 978-0-8153-4432-2 978-0-8153-4464-3 978-0-8153-4524-4 (Garland Science, Taylor and Francis Group, New York, NY, 2015).
- Goodrich, J. A. & Tjian, R. Unexpected Roles for Core Promoter Recognition Factors in Cell-Type-Specific Transcription and Gene Regulation. en. *Nature Reviews Genetics* 11, 549–558.
 ISSN: 1471-0056, 1471-0064 (Aug. 2010).
- Schoenfelder, S. & Fraser, P. Long-Range Enhancer-Promoter Contacts in Gene Expression Control. En. Nature Reviews Genetics, 1. ISSN: 1471-0064 (May 2019).
- 18. Spitz, F. & Furlong, E. E. M. Transcription Factors: From Enhancer Binding to Developmental Control. en. *Nature Reviews Genetics* **13**, 613–626. ISSN: 1471-0064 (Sept. 2012).
- Ong, C.-T. & Corces, V. G. Enhancer Function: New Insights into the Regulation of Tissue-Specific Gene Expression. en. *Nature Reviews Genetics* 12, 283–293. ISSN: 1471-0064 (Apr. 2011).
- Andersson, R. & Sandelin, A. Determinants of Enhancer and Promoter Activities of Regulatory Elements. en. Nature Reviews Genetics 21, 71–87. ISSN: 1471-0064 (Feb. 2020).
- Gaszner, M. & Felsenfeld, G. Insulators: Exploiting Transcriptional and Epigenetic Mechanisms. en. Nature Reviews Genetics 7, 703–713. ISSN: 1471-0064 (Sept. 2006).
- Oudelaar, A. M. & Higgs, D. R. The Relationship between Genome Structure and Function.
 en. Nature Reviews Genetics. ISSN: 1471-0056, 1471-0064 (Nov. 2020).
- Farnham, P. J. Insights from Genomic Profiling of Transcription Factors. en. Nature Reviews Genetics 10, 605–616. ISSN: 1471-0064 (Sept. 2009).
- 24. Kim, T. H. et al. Analysis of the Vertebrate Insulator Protein CTCF-Binding Sites in the Human Genome. en. Cell 128, 1231–1245. ISSN: 0092-8674 (Mar. 2007).
- Dixon, J. R. et al. Topological Domains in Mammalian Genomes Identified by Analysis of Chromatin Interactions. en. Nature 485, 376–380. ISSN: 1476-4687 (May 2012).

Kasowski, M. et al. Variation in Transcription Factor Binding Among Humans. en. Science
 328, 232–235. ISSN: 0036-8075, 1095-9203 (Apr. 2010).

- Maurano, M. T., Wang, H., Kutyavin, T. & Stamatoyannopoulos, J. A. Widespread Site-Dependent Buffering of Human Regulatory Polymorphism. *PLoS Genetics* 8. ISSN: 1553-7404 (Electronic)\n1553-7390 (Linking) (2012).
- Maurano, M. T. et al. Large-Scale Identification of Sequence Variants Influencing Human Transcription Factor Occupancy in Vivo. en. Nature Genetics 47, 1393–1401. ISSN: 1061-4036, 1546-1718 (Dec. 2015).
- Maurano, M. T. et al. Role of DNA Methylation in Modulating Transcription Factor Occupancy. en. Cell Reports 12, 1184–1195. ISSN: 22111247 (Aug. 2015).
- 30. Wang, H. et al. Widespread Plasticity in CTCF Occupancy Linked to DNA Methylation. Genome Research 22, 1680–1688. ISSN: 1549-5469 (Electronic)\n1088-9051 (Linking) (Sept. 2012).
- 31. Wiehle, L. et al. DNA (de)Methylation in Embryonic Stem Cells Controls CTCF-Dependent Chromatin Boundaries. en. Genome Research, gr.239707.118. ISSN: 1088-9051, 1549-5469 (Apr. 2019).
- 32. Xu, C. & Corces, V. G. Nascent DNA Methylome Mapping Reveals Inheritance of Hemimethylation at CTCF/Cohesin Sites. *Science* **359**, 1166–1170 (2018).
- 33. Viner, C. et al. Modeling Methyl-Sensitive Transcription Factor Motifs with an Expanded Epigenetic Alphabet. en. bioRxiv, 043794 (Mar. 2016).
- Goll, M. G. & Bestor, T. H. Eukaryotic Cytosine Methyltransferases. Annual Review of Biochemistry 74, 481–514. ISSN: 0066-4154 (June 2005).
- 35. Lister, R. et al. Human DNA Methylomes at Base Resolution Show Widespread Epigenomic Differences. Nature 462, 315–322. ISSN: 1476-4687 (Electronic)\n0028-0836 (Linking) (Nov. 2009).
- Henikoff, S. Nucleosome Destabilization in the Epigenetic Regulation of Gene Expression. en. Nature Reviews Genetics 9, 15–26. ISSN: 1471-0064 (Jan. 2008).
- 37. Jiang, C. & Pugh, B. F. Nucleosome Positioning and Gene Regulation: Advances through Genomics. en. *Nature Reviews Genetics* **10**, 161–172. ISSN: 1471-0064 (Mar. 2009).
- 38. Vierstra, J. et al. Global Reference Mapping of Human Transcription Factor Footprints. en. Nature 583, 729–736. ISSN: 1476-4687 (July 2020).

Cusanovich, D. A. et al. A Single-Cell Atlas of In Vivo Mammalian Chromatin Accessibility.
 en. Cell 174, 1309–1324.e18. ISSN: 0092-8674 (Aug. 2018).

- 40. Polak, P. et al. Cell-of-Origin Chromatin Organization Shapes the Mutational Landscape of Cancer. en. Nature 518, 360–364. ISSN: 1476-4687 (Feb. 2015).
- 41. Zhu, H., Wang, G. & Qian, J. Transcription Factors as Readers and Effectors of DNA Methylation. *Nature Reviews Genetics* **17**, 551–565 (Aug. 2016).
- Furey, T. S. ChIP-Seq and beyond: New and Improved Methodologies to Detect and Characterize Protein-DNA Interactions. en. Nature Reviews Genetics 13, 840-852. ISSN: 1471-0064 (Dec. 2012).
- 43. Carter, B. & Zhao, K. The Epigenetic Basis of Cellular Heterogeneity. en. *Nature Reviews Genetics* **22**, 235–250. ISSN: 1471-0064 (Apr. 2021).
- 44. Zhou, V. W., Goren, A. & Bernstein, B. E. Charting Histone Modifications and the Functional Organization of Mammalian Genomes. *Nature Reviews Genetics* **12**, 7–18 (Jan. 2011).
- 45. Dekker, J. & Mirny, L. The 3D Genome as Moderator of Chromosomal Communication. English. *Cell* **164**, 1110–1121. ISSN: 0092-8674, 1097-4172 (Mar. 2016).
- 46. Finn, E. H. & Misteli, T. Molecular Basis and Biological Function of Variability in Spatial Genome Organization. en. *Science* **365**, eaaw9498. ISSN: 0036-8075, 1095-9203 (Sept. 2019).
- Lieberman-Aiden, E. et al. Comprehensive Mapping of Long-Range Interactions Reveals Folding Principles of the Human Genome. en. Science 326, 289–293. ISSN: 0036-8075, 1095-9203 (Oct. 2009).
- 48. Rao, S. S. P. et al. A 3D Map of the Human Genome at Kilobase Resolution Reveals Principles of Chromatin Looping. English. Cell 159, 1665–1680. ISSN: 0092-8674, 1097-4172 (Dec. 2014).
- 49. Mirny, L. A., Imakaev, M. & Abdennur, N. Two Major Mechanisms of Chromosome Organization. en. *Current Opinion in Cell Biology. Cell Nucleus* **58**, 142–152. ISSN: 0955-0674 (June 2019).
- 50. Stergachis, A. B. *et al.* Conservation of Trans-Acting Circuitry during Mammalian Regulatory Evolution. *Nature* **515**, 365–370. ISSN: 1476-4687 (Electronic)\r0028-0836 (Linking) (2014).
- Berthelot, C., Villar, D., Horvath, J., Odom, D. T. & Flicek, P. Complexity and Conservation of Regulatory Landscapes Underlie Evolutionary Resilience of Mammalian Gene Expression. bioRxiv, 1–31 (2017).

52. Spurrell, C. H., Dickel, D. E. & Visel, A. The Ties That Bind: Mapping the Dynamic Enhancer-Promoter Interactome. English. *Cell* 167, 1163–1166. ISSN: 0092-8674, 1097-4172 (Nov. 2016).

- Buenrostro, J. D. et al. Single-Cell Chromatin Accessibility Reveals Principles of Regulatory Variation. en. Nature 523, 486–490. ISSN: 0028-0836, 1476-4687 (June 2015).
- Lee, T. I. & Young, R. A. Transcriptional Regulation and Its Misregulation in Disease. English.
 Cell 152, 1237–1251. ISSN: 0092-8674, 1097-4172 (Mar. 2013).
- 55. Conesa, A. et al. A Survey of Best Practices for RNA-Seq Data Analysis. Genome Biology 17, 13–13. ISSN: 1474-760X (Electronic)\r1474-7596 (Linking) (Dec. 2016).
- Robertson, G. et al. Genome-Wide Profiles of STAT1 DNA Association Using Chromatin Immunoprecipitation and Massively Parallel Sequencing. en. Nature Methods 4, 651–657. ISSN: 1548-7091, 1548-7105 (Aug. 2007).
- 57. Bailey, T. et al. Practical Guidelines for the Comprehensive Analysis of ChIP-Seq Data. en. PLOS Computational Biology 9, e1003326. ISSN: 1553-7358 (Nov. 2013).
- 58. Skene, P. J., Henikoff, J. G. & Henikoff, S. Targeted in Situ Genome-Wide Profiling with High Efficiency for Low Cell Numbers. en. *Nature Protocols* **13**, 1006–1019. ISSN: 1754-2189, 1750-2799 (May 2018).
- Boyle, A. P. et al. High-Resolution Mapping and Characterization of Open Chromatin across the Genome. English. Cell 132, 311–322. ISSN: 0092-8674, 1097-4172 (Jan. 2008).
- 60. Buenrostro, J. D., Giresi, P. G., Zaba, L. C., Chang, H. Y. & Greenleaf, W. J. Transposition of Native Chromatin for Fast and Sensitive Epigenomic Profiling of Open Chromatin, DNA-Binding Proteins and Nucleosome Position. *Nature Methods* 10, 1213–8 (Dec. 2013).
- Buenrostro, J., Wu, B., Chang, H. & Greenleaf, W. ATAC-Seq: A Method for Assaying Chromatin Accessibility Genome-Wide. Current Protocols in Molecular Biology 6, 356–372. ISSN: 6314442508 (2015).
- Corces, M. R. et al. An Improved ATAC-Seq Protocol Reduces Background and Enables Interrogation of Frozen Tissues. en. Nature Methods 14, 959–962. ISSN: 1548-7105 (Oct. 2017).
- Schones, D. E. et al. Dynamic Regulation of Nucleosome Positioning in the Human Genome.
 English. Cell 132, 887–898. ISSN: 0092-8674, 1097-4172 (Mar. 2008).
- 64. Laird, P. W. Principles and Challenges of Genome-Wide DNA Methylation Analysis. *Nature Reviews Genetics* **11**, 191–191. ISSN: 1471-0064 (Electronic)\r1471-0056 (Linking) (Mar. 2010).

 Dekker, J., Rippe, K., Dekker, M. & Kleckner, N. Capturing Chromosome Conformation. en. Science 295, 1306–1311. ISSN: 0036-8075, 1095-9203 (Feb. 2002).

- Nora, E. P. et al. Spatial Partitioning of the Regulatory Landscape of the X-Inactivation Centre.
 en. Nature 485, 381–385. ISSN: 0028-0836, 1476-4687 (May 2012).
- 67. Birney, E. et al. Identification and Analysis of Functional Elements in 1% of the Human Genome by the ENCODE Pilot Project. en. Nature 447, 799–816. ISSN: 1476-4687 (June 2007).
- 68. Moore, J. E. *et al.* Expanded Encyclopaedias of DNA Elements in the Human and Mouse Genomes. en. *Nature* **583**, 699–710. ISSN: 1476-4687 (July 2020).
- Ernst, J. & Kellis, M. ChromHMM: Automating Chromatin-State Discovery and Characterization. en. Nature Methods 9, 215–216. ISSN: 1548-7105 (Mar. 2012).
- Hoffman, M. M. et al. Unsupervised Pattern Discovery in Human Chromatin Structure through Genomic Segmentation. en. Nature Methods 9, 473–476. ISSN: 1548-7091, 1548-7105 (May 2012).
- Chan, R. C. W. et al. Segway 2.0: Gaussian Mixture Models and Minibatch Training. en. Bioinformatics 34 (ed Birol, I.) 669–671. ISSN: 1367-4803, 1460-2059 (Feb. 2018).
- 72. Zhou, S., Treloar, A. E. & Lupien, M. Emergence of the Noncoding Cancer Genome: A Target of Genetic and Epigenetic Alterations. *Cancer Discovery* **6**, 1215–1229 (Nov. 2016).
- 73. Gasperini, M., Tome, J. M. & Shendure, J. Towards a Comprehensive Catalogue of Validated and Target-Linked Human Enhancers. en. *Nature Reviews Genetics*, 1–19. ISSN: 1471-0064 (Jan. 2020).