The Complex Genome: From Structure to Function

Diwen (Steven) Gan May 17, 2023

Overview

In eukaryotes, the genome is extensively compacted inside the nucleus, hierarchically organized at a multiscale of structural units. One of the most critical roles of these delicately regulated structures is the control of transcriptional dynamics. Transcriptional regulations have been recognized as the primary regulatory mechanisms coordinating cellular activities, responses, cell type and fate determinations. No doubt, regulations on transcription are hugely achieved by changes in regulatory factors and structural configuration. Yet, with more than two hundred years of study on the genome, we are just starting to realize the complexity lies within ...

The course is designed for students to learn:

- 1) Basic knowledge of the genomic structure and its role in transcriptional regulation.
- 2) Cutting-edge techniques and bioinformatics used in the field.
- 3) Practical experience in bioinformatic analysis.
- 4) Frontier research principles and methodology.
- 5) Mysteries and dilemmas remained unsolved.

Schedule

Class meets online using Zoom/Lark on Tuesday and Saturday at 10:00 am - 12:00 pm (Beijing Time). Class on Saturday, Week 5 (Week $5 - 2^*$) is subject to change depending on the actual pacing.

Each class comprises 1 hour of lecture and $1 \sim 1.5$ hour of discussions or hands-on sessions, subject to change based on the progress of study.

Literature discussion is scheduled for each week. A research article and a related synopsis will be posted one or half week prior to the discussion. Students should read the article and complete the synopsis before the discussion. For each discussion, $1 \sim 2$ students will be selected as discussion leader to guide the discussion.

Week 2, 4 will have hands-on sessions, which requires Win/Mac setup, specific details to be determined.

Week 5 is scheduled for research proposal project. Recommended topics will be provided on week 3. However, students can pick other topics related to the course and discuss with the instructor no later than week 9.

Syllabus

Week 1 - 1

DNA, Chromatin, and Transcription

Introduction to the structure basis of DNA, nucleosomes, chromatins, and nucleus; Histone modifications and DNA methylations; Elements of genes and the transcription program; *Cis*- and *trans*-regulation.

Additional readings: Perspectives on transcription regulation
 Lis, J. T. A 50 year history of technologies that drove discovery in eukaryotic transcription regulation. Nat. Struct. Mol. Biol. 26, 777-782 (2019).

Week 1 - 2

Methods to Discovery

Quantification of gene expression (from RNA to proteins); Functional analysis of genes: fluorescent reporter assay, *in situ* hybridization, genetic perturbations (RNAi, CRISPR/Cas system, and protein degrons); Introduction to omics. *Principle of paper discussion*

- Discussion paper 1: Protein degrons
 Nishimura, K., Fukagawa, T., Takisawa, H., Kakimoto, T. & Kanemaki, M. An auxin-based degron system for the rapid depletion of proteins in nonplant cells. *Nat. Methods* 6, 917-922 (2009).
- Following paper: Advanced protein degrons
 Yesbolatova, A. et al. The auxin-inducible degron 2 technology provides sharp
 degradation control in yeast, mammalian cells, and mice. Nat. Commun. 11, 5701
 (2020).

Week 2 - 1

Sequence, Omics, and Bioinformatics I

Three generations of sequencing; Sequence alignment; Genomics and genome element (GWAS); Transcriptomics (RNA-seq & nascent RNA-seq). Paper 1 discussion; Hands-on session: Sequence alignment & database searching.

- Discussion paper 2: Loop extrusion model
 Fudenberg, G. et al. Formation of Chromosomal Domains by Loop Extrusion. Cell Rep. 15, 2038-2049 (2016).
- Additional readings: Review of high throughput sequencing
 Reuter, Jason A., Spacek, D. V. & Snyder, Michael P. High-Throughput Sequencing
 Technologies. Mol. Cell 58, 586-597 (2015).

Week 2 - 2

Sequence, Omics, and Bioinformatics II

Functional annotation (GO & KEGG); Epigenomics (ATAC-seq, ChIP-seq, BS-seq); Hi-C; Screening and lineage tracing. *Hands-on session: ChIP-seq analysis using Galaxy.*

Additional readings: Reviews of epigenetic technologies
 Rivera, Chloe M. & Ren, B. Mapping Human Epigenomes. Cell 155, 39-55 (2013).
 Mehrmohamadi, M., Sepehri, M. H., Nazer, N. & Norouzi, M. R. A Comparative
 Overview of Epigenomic Profiling Methods. Front. Cell Dev. Biol. 9 (2021).

Week 3 - 1

<u>Transcriptional Regulation I: Enhancers and Transcription factors</u>

Role of enhancers and TFs in transcription initiation, pause-and-release, etc.; Study higher-dimensional architecture of genomes; Model of enhancer-promoter interactions; Enhancer RNA. *Paper 2 discussion*

- Discussion paper 3: Cohesin and chromatin structure
 Rao, S. S. P. et al. Cohesin Loss Eliminates All Loop Domains. Cell 171, 305-320.e324 (2017).
- Parallel study: CTCF and chromatin structure
 Nora, E. P. et al. Targeted Degradation of CTCF Decouples Local Insulation of
 Chromosome Domains from Genomic Compartmentalization. Cell 169, 930-944.e922
 (2017).
- Additional readings: Reviews of epigenetic technologies
 Haberle, V. & Stark, A. Eukaryotic core promoters and the functional basis of transcription initiation. Nat. Rev. Mol. Cell Biol. 19, 621-637 (2018).
 Andersson, R. & Sandelin, A. Determinants of enhancer and promoter activities of regulatory elements. Nat. Rev. Genet. 21, 71-87 (2020).
 Li, W., Notani, D. & Rosenfeld, M. G. Enhancers as non-coding RNA transcription units: recent insights and future perspectives. Nat. Rev. Genet. 17, 207-223 (2016).

Week 3 - 2

Transcriptional Regulation II: Epigenetics

Epigenetic landscape of the genome; Epigenetic marks of gene activation and silencing; Regulation and dynamics of epigenetics and chromatin remodeling; Epigenetics in development. *Paper 3 discussion*

- Discussion paper 4: Phase separation at enhancer regions
 Boija, A. et al. Transcription Factors Activate Genes through the Phase-Separation
 Capacity of Their Activation Domains. Cell 175, 1842-1855.e1816 (2018).
- Additional readings: Reviews of epigenetic regulation & profile of histone marks

Klemm, S. L., Shipony, Z. & Greenleaf, W. J. Chromatin accessibility and the regulatory epigenome. *Nat. Rev. Genet.* **20**, 207-220 (2019). Barski, A. *et al.* High-Resolution Profiling of Histone Methylations in the Human Genome. *Cell* **129**, 823-837 (2007).

Week 4 - 1

<u>Transcriptional Regulation III: Condensates and Bursting?</u>

Lipid-lipid phase separation in cells; Novel (yet skeptical) theory of condensates/ clusters/hubs; Dynamics of condensate; Transcriptional bursting. Paper 4 discussion

- Discussion paper 5: Loop stacking model
 Hafner, A. et al. Loop stacking organizes genome folding from TADs to chromosomes.
 Mol. Cell 83, 1377-1392.e1376 (2023).
- Additional readings: Perspectives on condensate, loop, and hub model
 Lim, B. & Levine, M. S. Enhancer-promoter communication: hubs or loops? Curr. Opin.
 Genet. Dev. 67, 5-9 (2021).
 Furlong, E. E. M. & Levine, M. Developmental enhancers and chromosome topology.
 Science 361, 1341-1345 (2018).

Week 4 - 2

A Case of Study – Estrogen Receptor

Estrogen response pathway; Binding of estrogen receptor; Mechanism of estrogen-induced transcription; MegaTrans enhancers. *Hands-on session: Statistical Model of TF-DNA Interactions*

- Additional readings: MegaTrans of estrogen-regulated enhancers
 Liu, Z. et al. Enhancer Activation Requires Trans-Recruitment of a Mega Transcription
 Factor Complex. Cell 159, 358-373 (2014).
- Additional readings: Phase separation at estrogen-induced enhancers
 Nair, S. J. et al. Phase separation of ligand-activated enhancers licenses cooperative chromosomal enhancer assembly. Nat. Struct. Mol. Biol. 26, 193-203 (2019).

Week 5 - 1

A Case of Study - Enhancer-Promoter Contact and 3D Organization

Global analysis of chromatin folding; Advanced technologies (Micro-C & MERFISH); Shaping of enhancer-promoter contact; Discoveries against canonical theory; Problems of heterogenicity. Paper 5 discussion

Additional readings: Reviews of the 3D genome structure
 Zheng, H. & Xie, W. The role of 3D genome organization in development and cell
 differentiation. Nat. Rev. Mol. Cell Biol. 20, 535-550 (2019).

Hafner, A. & Boettiger, A. The spatial organization of transcriptional control. *Nat. Rev. Genet.* **24**, 53-68 (2023).

van Steensel, B. & Furlong, E. E. M. The role of transcription in shaping the spatial organization of the genome. *Nat. Rev. Mol. Cell Biol.* **20**, 327-337 (2019).

Week 5 - 2*

The 4D Genome and New Paradigm

Summary of the hierarchical 3D structure of chromatin; Reconstruction of the 3D genome; Reorganization of the genome during development; Structure-determined phenotype and diseases. *Research proposal discussion*

Additional readings: The 4D Genome
 Aboelnour, E. & Bonev, B. Decoding the organization, dynamics, and function of the 4D genome. Dev. Cell 56(11), 1562–1573 (2021).

 Dekker, J. et al. The 4D nucleome project. Nature. 549, 219–226 (2017).

Week 6

Final

Research proposal presentation

Q&A