RESEARCH HIGHLIGHTS

IN BRIEF

PATHOGEN GENETICS

Making male malaria mosquitoes

UK researchers have identified and characterized a master regulator of the sex determination process in the African malaria mosquito Anopheles gambiae. The team analysed the transcriptomes of male and female embryos and identified the gene Yob as a maleness-conferring Y chromosome-linked factor. Yob expression was observed from the onset of zygotic transcription throughout the life of males. The gene was found to encode a 56-amino-acid protein that controls the male-specific splicing of *doublesex* (*dsx*). Ectopic embryonic delivery of Yob mRNA was lethal for genetically female embryos, but had no effect on genetic males. By contrast, silencing of embryonic Yob expression resulted in male-specific lethality, suggesting a role of Yob in dosage compensation. Yob could be useful as a tool to produce male-only generations for transgenic approaches to control vector-borne diseases.

ORIGINAL ARTICLE Krzywinska, E. et al. A maleness gene in the malaria mosquito Anopheles gambiae. Science 353, 67–69 (2016)

TECHNIQUE

Genome-wide quantification of 5hmC in single cells

A new technique enables genome-wide detection and quantification of the epigenetic mark 5-hydroxymethylcytosine (5hmC) in single cells. Based on a method for bulk 5hmC sequencing, 5hmC marks are glucosylated using the T4 phage β -glucosyltransferase, then cut by the restriction enzyme AbaSI. Digested genomic DNA is ligated to double-stranded adapters containing a 2-nucleotide random 3' overhang, together with a cell-specific barcode, an Illumina 5' adapter and a T7 promoter. In vitro transcription is used to amplify the DNA fragments linearly in a strand-specific orientation, and the amplified RNA is fragmented and undergoes directional RNA library preparation. When applied to mouse embryonic stem cells, a median of 44,000 unique 5hmC sites per cell was detected. Substantial cell-to-cell variability existed between the number of 5hmC sites on the two strands of the same chromosome. Differences in age between the strands of a chromosome could explain the 5hmC strand bias, as confirmed by a new stochastic model. ORIGINAL ARTICLE Mooiiman, D. et al. Single-cell 5hmC sequencing reveals chromosome-

ORIGINAL ARTICLE Wooljman, D. et al. Single-cell shinc sequencing reveals chromosomewide cell-to-cell variability and enables lineage reconstruction. Nat. Biotechnol. <u>http://dx. doi.org/10.1038/nbt.3598</u> (2016)

CHROMATIN

Programmed R-loop formation

Using DNA-RNA immunoprecipitation followed by cDNA conversion coupled to high-throughput sequencing (DRIPc-seq), researchers have profiled the genome-wide prevalence and distribution of R loops in mouse and human cells. Mapping of R loops at near base-pair resolution and in a strand-specific manner showed that the co-transcriptional hybridization of nascent RNAs to template DNA is a conserved, prevalent and dynamic feature of mammalian chromatin that can impact gene expression. Epigenomic profiling revealed that R loops associate with specific epigenomic signatures: at promoters, R loops associate with an open, histone H3 lysine 4 (H3K4) hypermethylated and hyperacetylated state characteristic of strong CpG island promoters; at terminators, R loops associate with an enhancer- and insulator-like state; and R-loop formation seems to be a conserved hallmark of a broad class of transcription terminators.

ORIGINAL ARTICLE Sanz, L. A. et al. Prevalent, dynamic, and conserved R-loop structures associate with specific epigenomic signatures in mammals. *Mol. Cell* <u>http://dx.doi.org/10.1016/j.molcel.2016.05.032</u> (2016)