

How the 'master regulators' of cells make DNA accessible for gene expression

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Cbf1 pioneer factor/nucleosome complex

New research by researchers at Penn State and Ohio State helps explain how proteins called pioneer factors are able to open up the typically dense genetic material within our cells, packaged into complexes called nucleosomes. This image depicts the molecular structure of the yeast Cbf1 pioneer factor interacting with both DNA and histone components of the nucleosome. Credit: Song Tan lab, Penn State



New insight into proteins called "pioneer factors" helps to explain their unusual ability to open up the typically dense genetic material within our cells. This behavior makes the genetic material accessible for proteins involved in important cellular processes, such as DNA replication and repair, gene expression, and the creation of proteins.

The study, by a team of researchers from Penn State and the Ohio State University, used a unique combination of structural biology, biophysics, and <u>cell biology</u> to understand how these "master regulators" of the genome interact with nucleosomes—the basic unit of the genome in all <u>eukaryotic cells</u>, which range from yeast to humans.

DNA inside the nucleus of cells is generally wrapped around proteins called histones and packed into dense complexes called nucleosomes. The nucleosomes, which resemble beads on a string of DNA, are further packed together forming chromatin, which then make up chromosomes.

"Nucleosomes are a barrier for a lot of proteins to associate with chromatin, but pioneer factors have a special property that they can invade into the nucleosome and generate open regions that can be accessed by other factors," said Lu Bai, associate professor of biochemistry and <u>molecular biology</u> and of physics at Penn State and one of the leaders of the research team.

"Because of this, they are sometimes thought of as 'master regulators' of genes. In this study, we used a combination of approaches to better understand this how pioneer factors can invade the nucleosome."

The study appears in the journal Molecular Cell.

Pioneer factors are a type of transcription factor—proteins critical to the process of transcription, where DNA is copied into the RNA blueprints for the creation of proteins. While many <u>transcription factors</u> can bind to



the nucleosome, most fall off very quickly. By contrast, pioneer factors have what is called a "dissociation compensation mechanism" that allows them to remain stably bound to the nucleosome for an extended period of time.

The research team compared a pioneer factor called Cbf1 and a nonpioneer transcription factor called Pho4 in budding yeast. The two proteins have similar overall structures and the ability to recognize the same DNA sequences but they behave differently around the nucleosome. Using a sensitive imaging technique called <u>cryo-electron</u> <u>microscopy</u>, the researchers identified a structure on the pioneer factor that they believe contributes to its ability to invade the nucleosome.

"Based on our cryoelectron microscopy structure, Cbf1 interacts not only with the DNA part of the nucleosome, but also with histones within the nucleosome through its helix-loop-helix region," said Song Tan, Verne M. Willaman Professor of Molecular Biology at Penn State and one of the leaders of the research team. "We suspect this interaction helps prevent Cbf1 from dissociating as quickly as non-pioneer factors."

To confirm the role of the helix-loop-helix region, the research team made "chimeras" of each of the proteins, removing a helix-loop-helix from the pioneer factor and adding one to the non-pioneer transcription factor. Then they measure how quickly these chimeras dissociated from the nucleosome.

"Removing the helix-loop-helix from the pioneer factor caused quicker dissociation, which resulted in a reduced dissociation compensation phenomenon and a dramatically less effective pioneer factor," said Michael Poirier, professor and chair of physics at Ohio State and one of the leaders of the research team. "Amazingly, adding the helix-loophelix region conferred pioneer factor properties to the non-pioneer factor."



To further confirm that their results, the researchers explored how these chimeras functioned within live yeast cells. They found that the pioneer factor and non-pioneer chimera modified to behave like a pioneer factor both facilitated invasion and opening up of the nucleosome. These results collectively help explain how pioneer factors like Cbf1 can access the nucleosome and increase accessibility of DNA for other factors.

"This study would not have been possible without combining expertise from our three separate groups," said Tan

"My group focused on structural biology, Michael Poirier's group at Ohio State focused on single molecule biophysics, and Lu Bai's group at Penn State focused on in vivo cellular biology. This has been a wonderfully synergistic collaboration and has produced ideas and results that would not have been generated with one or even two of our groups on their own. We are continuing this collaboration to explore interactions of other pioneer factors and the <u>nucleosome</u>."

More information: Benjamin T. Donovan et al, Basic helix-loop-helix pioneer factors interact with the histone octamer to invade nucleosomes and generate nucleosome-depleted regions, *Molecular Cell* (2023). DOI: 10.1016/j.molcel.2023.03.006

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