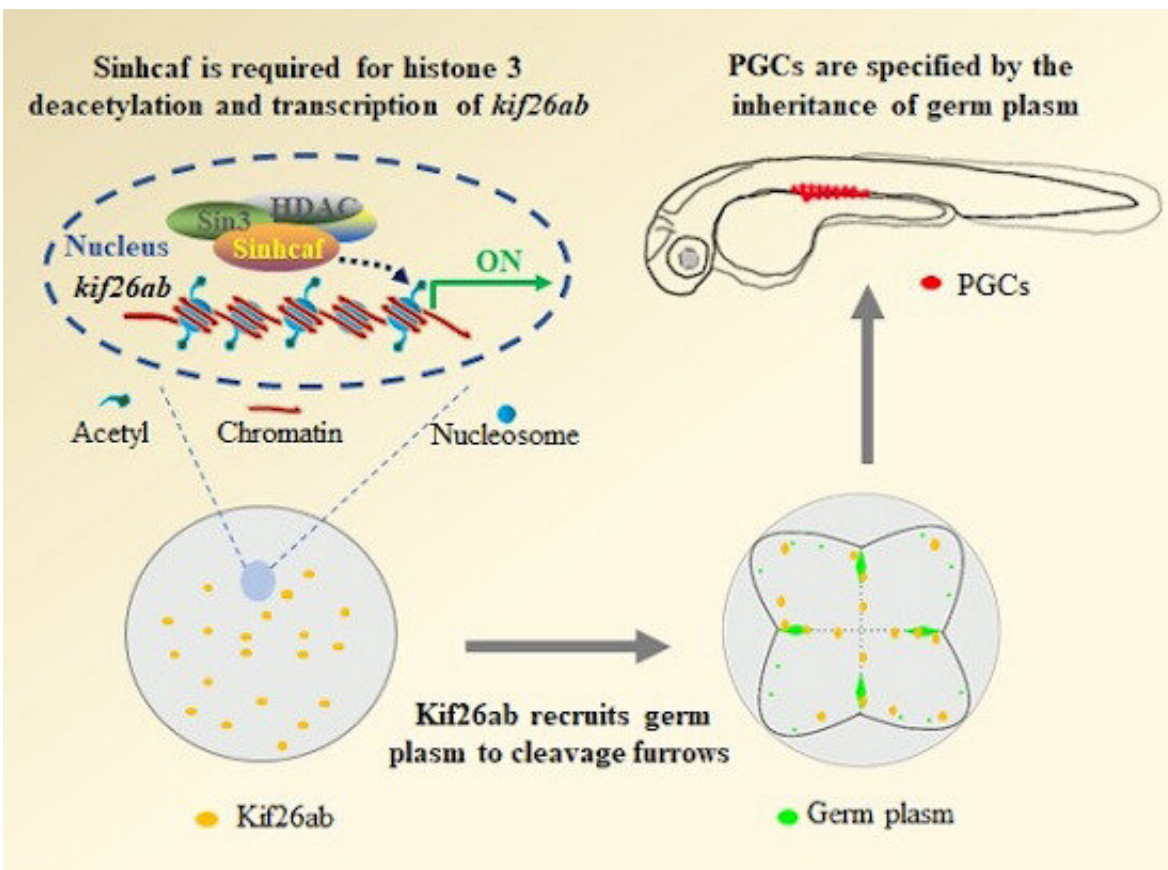


Sinhcaf-dependent histone deacetylation essential for primordial germ cell specification

May 10 2022, by Liu Jia



Graphical abstract. Credit: *EMBO reports* (2022). DOI: 10.15252/embr.202154387

Primordial germ cells (PGCs) are the first germ-cell population

established during development. The survival of species is dependent upon PGCs in sexually reproducing organisms because they are the founder cells for the germline.

In *Caenorhabditis elegans*, *Drosophila melanogaster*, *Xenopus laevis* and the zebrafish *Danio rerio*, PGCs are specified by inheritance of maternal germ plasm during early embryogenesis. Germ plasm is both sufficient and necessary for zebrafish PGC specification, and germ plasm accumulation at cleavage furrows requires coordination of microtubules, actin cytoskeletons and molecular motors. However, how this process is regulated remains poorly understood.

Recently, a research team led by Prof. Hu Wei from the Institute of hydrobiology (IHB) of the Chinese Academy of Sciences provides the first evidence that *Sinhcaf*-dependent histone deacetylation is essential for germ plasm aggregation and primordial germ cell specification. The study was published in *EMBO Reports*.

The researchers first generated *sinhcaf* knockout and overexpression zebrafish lines. They found that the maternal-zygotic *sinhcaf* mutants (MZ*sinhcaf*^{-/-}) exhibited germ plasm aggregation defects, decreased PGC abundance and male biased sex ratio. All of these effects were rescued by *sinhcaf* overexpression which led to excess PGCs and a female-biased sex ratio.

Using co-immunoprecipitation analysis, the researchers found that zebrafish *Sinhcaf* interacted with the core components of SIN3-HDAC complex. Western blot results indicated that loss of *Sinhcaf* resulted in an increased acetylation level of histone 3 in zebrafish full-grown stage (FG) follicles.

The researchers then conducted transcriptomic and real-time quantitative PCR analysis, and revealed that the level of *kif26ab* transcripts was

significantly decreased in *sinhcaf* mutant FG follicles and mature eggs. Chromatin immunoprecipitation results further demonstrated *Sinhcaf* can directly bind to the transcriptional regulatory region of *kif26ab*.

Loss of *sinhcaf* increased both acetyl-histone H3 (K9) and acetyl-histone H3 (K18) levels in the promoter region of *kif26ab*, said Prof. Hu, adding that *kif26ab* is transcriptionally activated by *Sinhcaf*, according to the luciferase-promoter assay.

The researchers demonstrated that suppressed expression of *kif26ab* caused defective germ plasm aggregation in cleavage furrows, and decreased PGCs number in genital ridge. Injection of *kif26ab* mRNA could partly rescue the decreased germ plasm components in cleavage furrows of *MZsinhcaf*^{-/-} embryos.

This study uncovers a critical role of *Sinhcaf* in germ plasm aggregation and subsequent PGC specification, which is mediated by regulating the histone acetylation status of *kif26ab* promoter to activate its transcription. The *sinhcaf*^{-/-} mutants offer an amendable *in vivo* model system to determine what controls PGC specification.

More information: Binbin Tao et al, *Sinhcaf*-dependent histone deacetylation is essential for primordial germ cell specification, *EMBO reports* (2022). [DOI: 10.15252/embr.202154387](https://doi.org/10.15252/embr.202154387)

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