

# **Epigenetic regulation of gene expression**

PRC2 methylates histones and DNA	
TET1,2,3 and TDG demethylate DNA	
Negative epigenetic regulation of rRNA expression	
Formation of WDR5-containing histone-modifying complexes	
nucleoplasm	
stosol fread	

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This is just an excerpt of a full-length report for this pathway. To access the complete report, please download it at the <u>Reactome Textbook</u>.

04/05/2024

### Introduction

Reactome is open-source, open access, manually curated and peer-reviewed pathway database. Pathway annotations are authored by expert biologists, in collaboration with Reactome editorial staff and cross-referenced to many bioinformatics databases. A system of evidence tracking ensures that all assertions are backed up by the primary literature. Reactome is used by clinicians, geneticists, genomics researchers, and molecular biologists to interpret the results of high-throughput experimental studies, by bioinformaticians seeking to develop novel algorithms for mining knowledge from genomic studies, and by systems biologists building predictive models of normal and disease variant pathways.

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#### Literature references

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- Sidiropoulos, K., Viteri, G., Sevilla, C., Jupe, S., Webber, M., Orlic-Milacic, M. et al. (2017). Reactome enhanced pathway visualization. *Bioinformatics*, 33, 3461-3467. A
- Fabregat, A., Jupe, S., Matthews, L., Sidiropoulos, K., Gillespie, M., Garapati, P. et al. (2018). The Reactome Pathway Knowledgebase. *Nucleic Acids Res, 46*, D649-D655. ↗
- Fabregat, A., Korninger, F., Viteri, G., Sidiropoulos, K., Marin-Garcia, P., Ping, P. et al. (2018). Reactome graph database: Efficient access to complex pathway data. *PLoS computational biology*, *14*, e1005968. *オ*

This document contains 5 pathways (see Table of Contents)

#### **Epigenetic regulation of gene expression ↗**

Stable identifier: R-DME-212165

Compartments: nucleoplasm

Inferred from: Epigenetic regulation of gene expression (Homo sapiens)

PRC2 methylates histones and DNA	
TET1,2,3 and TDG demethylate DNA	
Negative epigenetic regulation of rRNA expression	
Formation of W DR5-containing histone-modifying complexes	
sol	

This event has been computationally inferred from an event that has been demonstrated in another species.

The inference is based on the homology mapping from PANTHER. Briefly, reactions for which all involved PhysicalEntities (in input, output and catalyst) have a mapped orthologue/paralogue (for complexes at least 75% of components must have a mapping) are inferred to the other species. High level events are also inferred for these events to allow for easier navigation.

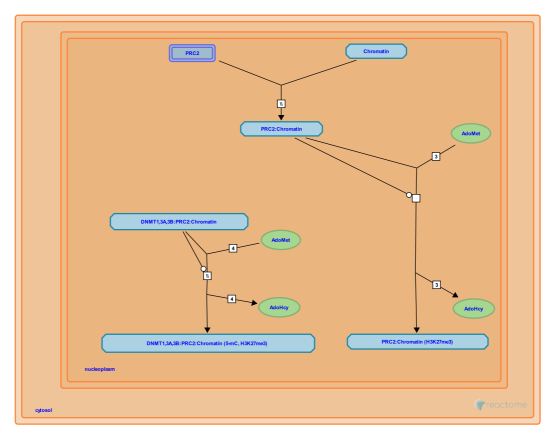
#### PRC2 methylates histones and DNA 7

Location: Epigenetic regulation of gene expression

Stable identifier: R-DME-212300

Compartments: nucleoplasm

Inferred from: PRC2 methylates histones and DNA (Homo sapiens)



This event has been computationally inferred from an event that has been demonstrated in another species.

The inference is based on the homology mapping from PANTHER. Briefly, reactions for which all involved PhysicalEntities (in input, output and catalyst) have a mapped orthologue/paralogue (for complexes at least 75% of components must have a mapping) are inferred to the other species. High level events are also inferred for these events to allow for easier navigation.

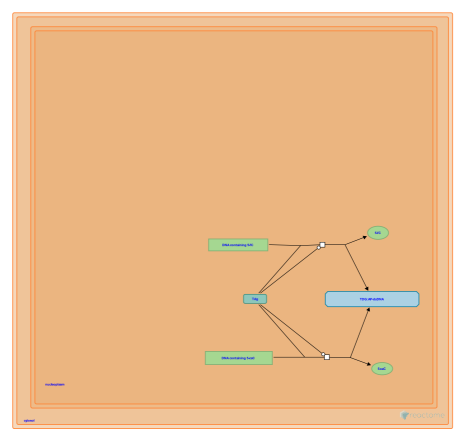
#### TET1,2,3 and TDG demethylate DNA 7

Location: Epigenetic regulation of gene expression

Stable identifier: R-DME-5221030

Compartments: nucleoplasm

Inferred from: TET1,2,3 and TDG demethylate DNA (Homo sapiens)



This event has been computationally inferred from an event that has been demonstrated in another species.

The inference is based on the homology mapping from PANTHER. Briefly, reactions for which all involved PhysicalEntities (in input, output and catalyst) have a mapped orthologue/paralogue (for complexes at least 75% of components must have a mapping) are inferred to the other species. High level events are also inferred for these events to allow for easier navigation.

#### Negative epigenetic regulation of rRNA expression 7

Location: Epigenetic regulation of gene expression

Stable identifier: R-DME-5250941

Compartments: nucleoplasm

Inferred from: Negative epigenetic regulation of rRNA expression (Homo sapiens)



This event has been computationally inferred from an event that has been demonstrated in another species.

The inference is based on the homology mapping from PANTHER. Briefly, reactions for which all involved PhysicalEntities (in input, output and catalyst) have a mapped orthologue/paralogue (for complexes at least 75% of components must have a mapping) are inferred to the other species. High level events are also inferred for these events to allow for easier navigation.

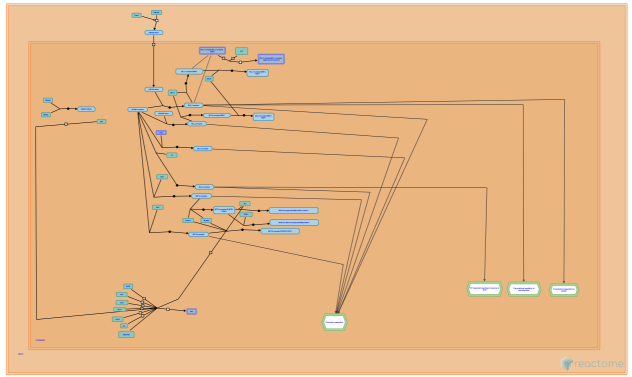
#### Formation of WDR5-containing histone-modifying complexes 7

Location: Epigenetic regulation of gene expression

Stable identifier: R-DME-9772755

Compartments: nucleoplasm

Inferred from: Formation of WDR5-containing histone-modifying complexes (Homo sapiens)



This event has been computationally inferred from an event that has been demonstrated in another species.

The inference is based on the homology mapping from PANTHER. Briefly, reactions for which all involved PhysicalEntities (in input, output and catalyst) have a mapped orthologue/paralogue (for complexes at least 75% of components must have a mapping) are inferred to the other species. High level events are also inferred for these events to allow for easier navigation.

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