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EDITED AND REVIEWED BY Meizhu Bai, Yale University, United States

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RECEIVED 26 February 2024 ACCEPTED 28 February 2024 PUBLISHED 11 March 2024

CITATION

Kim S, Yuan JB, Woods WS, Newton DA, Perez-Pinera P and Song JS (2024), Corrigendum: Chromatin structure and context-dependent sequence features control prime editing efficiency. *Front. Genet.* 15:1391923. doi: 10.3389/fgene.2024.1391923

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Corrigendum: Chromatin structure and context-dependent sequence features control prime editing efficiency

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KEYWORDS

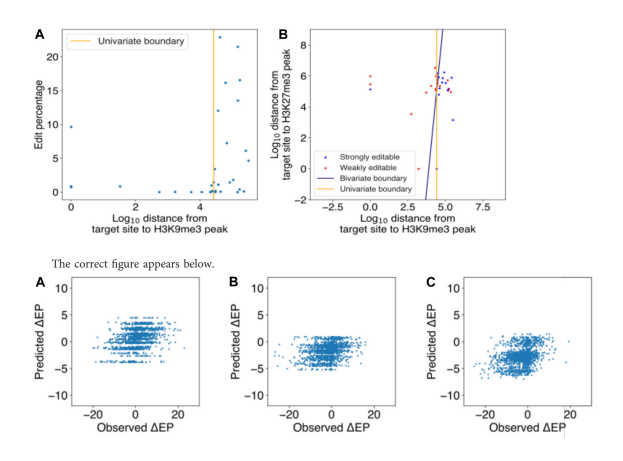
prime editing, CRISPR-Cas9, heterochromatin, nucleosome positioning, DNA-RNA hybridization, nucleotide preference, machine learning, neural network interpretation

A Corrigendum on

Chromatin structure and context-dependent sequence features control prime editing efficiency

by Kim S, Yuan JB, Woods WS, Newton DA, Perez-Pinera P and Song JS (2023). Front. Genet. 14: 1222112. doi: 10.3389/fgene.2023.1222112

In the published article, there was an error in **Supplementary Figure S3**. Even though the figure caption for Supplementary Figure S3 was correct, Figure S3 was inadvertently duplicated from Figure S2 and appeared as:



The authors apologize for this error and state that this does not change the scientific conclusions of the article in any way. The original article has been updated.

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