

CORRECTION **OPEN**

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Publisher Correction: A 39 kb structural variant causing Lynch syndrome detected by optical genome mapping and nanopore sequencing

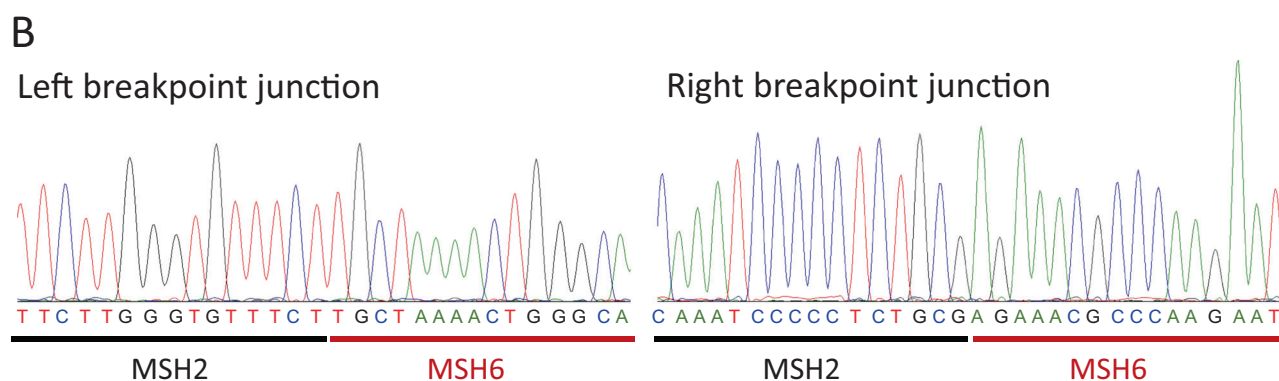
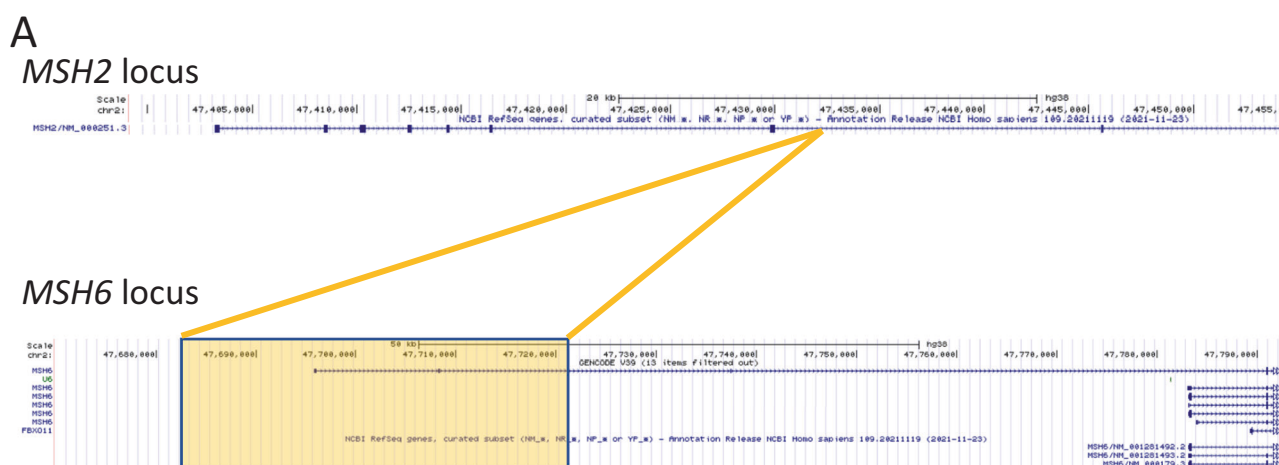
Pål Marius Bjørnstad , Ragnhild Aaløkken, June Åsheim, Arvind Y. M. Sundaram, Caroline N. Felde, G. Henriette Østby, Marianne Dalland , Wenche Sjursen, Christian Carrizosa, Magnus D. Vigeland, Hanne S. Sorte, Ying Sheng, Sarah L. Ariansen, Eli Marie Grindedal and Gregor D. Gilfillan

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Correction to: *European Journal of Human Genetics* <https://doi.org/10.1038/s41431-023-01494-7>, published online 29 November 2023

In this article, the wrong figure appeared as Fig. 3B due to typesetting mistake.; the Fig. 3 should have appeared as shown below.



The original article has been corrected.



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