














<https://doi.org/10.1038/s41467-020-20128-w>

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Author Correction: Retrospective evaluation of whole exome and genome mutation calls in 746 cancer samples

Matthew H. Bailey , William U. Meyerson, Lewis Jonathan Dursi , Liang-Bo Wang , Guanlan Dong , Wen-Wei Liang, Amila Weerasinghe , Shantao Li , Yize Li, Sean Kelso, MC3 Working Group*, PCAWG novel somatic mutation calling methods working group*, Gordon Saksena , Kyle Ellrott , Michael C. Wendl, David A. Wheeler , Gad Getz , Jared T. Simpson, Mark B. Gerstein , Li Ding & PCAWG Consortium*

Correction to: *Nature Communications*, <https://doi.org/10.1038/s41467-020-18151-y>, published online 21 September 2020.

The original version of this Article omitted from the author list the 9th author Yize Li, who is from the ‘The McDonnell Genome Institute at Washington University, St. Louis, MO 63108, USA and Department of Medicine, Division of Oncology, Washington University School of Medicine, St. Louis, MO 63108, USA’.

This has been corrected in both the PDF and HTML versions of the Article.

Published online: 30 November 2020



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