






<https://doi.org/10.1038/s41467-022-29652-3>

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Author Correction: Assortative mating biases marker-based heritability estimators

Richard Border , Sean O'Rourke, Teresa de Candia, Michael E. Goddard, Peter M. Visscher , Loic Yengo, Matt Jones & Matthew C. Keller 

Correction to: *Nature Communications* <https://doi.org/10.1038/s41467-022-28294-9>, published online 03 February 2022.

The original version of this Article contained an error in the third sentence of the results subsection “AM-induced bias persists when not all causal variants are measured” which incorrectly read “in simulated data that randomly dropped $\pi = 0, 50, \text{ or } 75\%$ of all $p = 10^6$ variants, including both causal ($m = 10^4$) and non-causal ($p - m = 990,000$) SNPs (Fig. 4)”. The correct version refers to “Fig. 3b” instead of “Fig. 4”. This has been corrected in both the PDF and HTML versions of the Article.

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