

CORRECTION

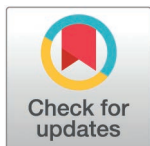
Correction: A pragmatic pipeline for drug resistance and lineage identification in *Mycobacterium tuberculosis* using whole genome sequencing

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Reference

1. Elton L, Aydin A, Stoker N, Rofael S, Wildner LM, Abdul JBPAA, et al. A pragmatic pipeline for drug resistance and lineage identification in *Mycobacterium tuberculosis* using whole genome sequencing. PLOS Glob Public Health. 2025;5(2):e0004099. <https://doi.org/10.1371/journal.pgph.0004099> PMID: [39928651](https://pubmed.ncbi.nlm.nih.gov/39928651/)



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