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Correction: Standardization of 16S rRNA gene sequencing using nanopore long read sequencing technology for clinical diagnosis of culture negative infections

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Affiliation “⁵Faculty of Biology, Medicine and Health, The University of Manchester” was omitted for author Ian Butler. This affiliation has now been added for author Ian Butler “Ian Butler^{1,2*} Olivia Turner¹, Kulsoom Mohammed¹, Mazeda Akhtar¹, Daniel Evans³, Jonathan Lambourne¹, Kathryn Harris¹, Denise M. O'Sullivan^{3,4} and Chrysi Sergaki⁵

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The corrected Acknowledgments section appears below.

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The original version of this article has been updated.

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