

CORRECTION

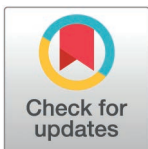
Correction: Metagenomic characterization of bacterial abundance and diversity in potato cyst nematode suppressive and conducive potato rhizosphere

John Kamathi Kiige, Agnes Mumo Kavoo, Mwashasha Rashid Mwajita, Derleen Mogire, Stephen Ogada, Tofick Barasa Wekesa, Leonard Muriithi Kiirika

The reference list is incorrectly numbered. The correct list is:

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