ERRATUM

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Erratum to: 'Reference-free inference of tumor phylogenies from single-cell sequencing data'

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Unfortunately, the original version of this article [1] contained an error. Figures 2, 4 and 5 were incorrect and the captions for Figs. 4 and 5 were incorrect. Below are the correct figures and captions:

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Reference

1. Schwartz R, Subramanian A. Reference-free inference of tumor phylogenies from single-cell sequencing data'. BMC Genomics. 2015;16 Suppl 11:S7.

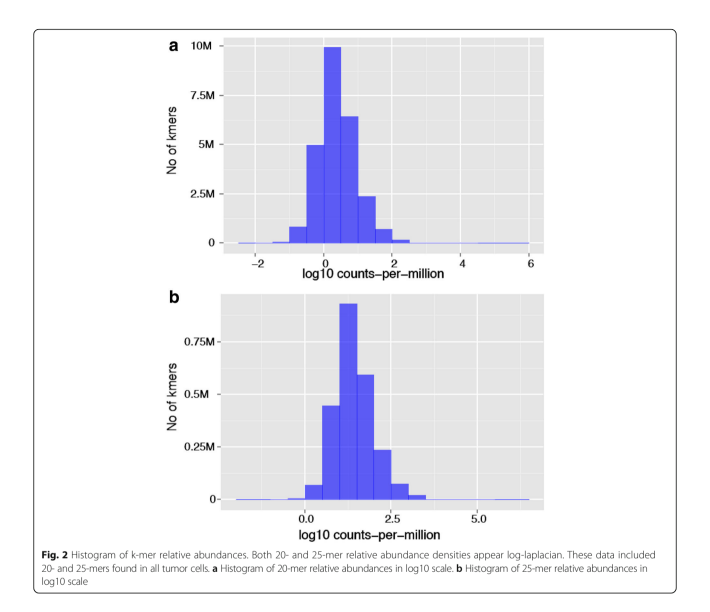
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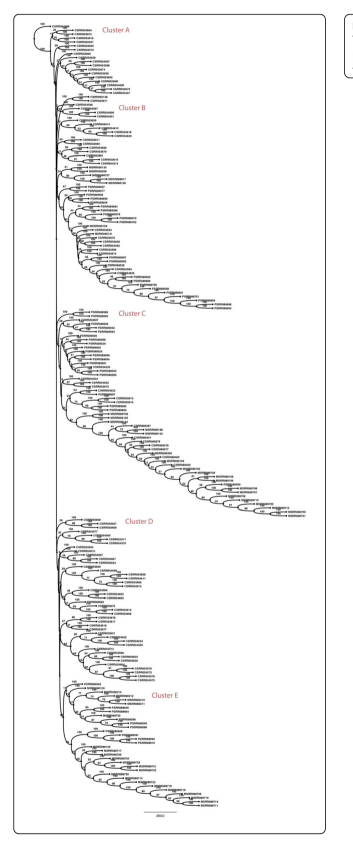


Fig. 4 20-mer bootstrap consensus neighbor-joining tree built from T10 primary breast tumor cells (prefix C), T16 primary (prefix P) and metastatic data (prefix M). Distinct groupings of cells are labeled as clusters

