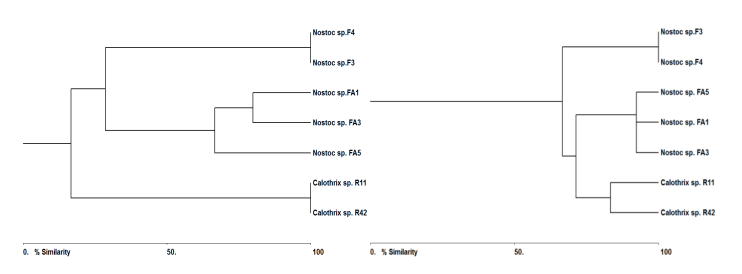
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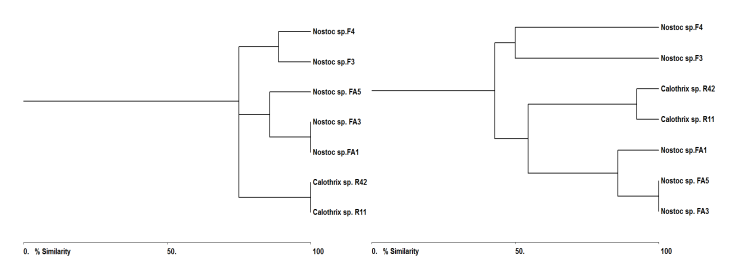
Genomic Fingerprinting Using Highly Repetitive Sequences to Differentiate Close Cyanobacterial Strains

Rezvan Shokraei^a, Hossein Fahimi^a, Saúl Blanco^b, Bahareh Nowruzi^c*

SUPPLEMENTARY DATA

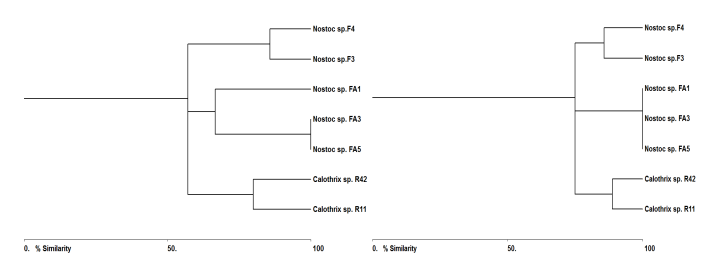


Supp. Figure 1. Composite dendrogram based on the ERIC1A (left) and ERIC1B (right) primers amplification profiles.

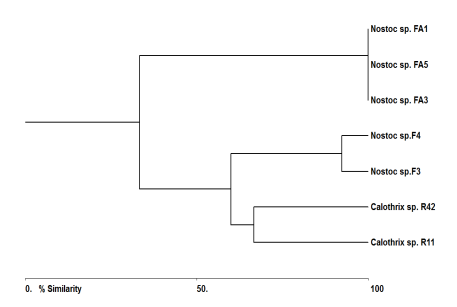


Supp. Figure 2. Composite dendrogram based on the HIP-CA (left) and HIP-AT (right) primers amplification profiles.

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 $Supp.\ Figure\ 3.\ Composite\ dendrogram\ based\ on\ the\ HIP-TG\ (left)\ and\ HIP-GC\ (right)\ primers\ amplification\ profiles.$



Supp. Figure 4. Composite dendrogram based on the STRR1a primers amplification profiles.