

Resource Summary Report

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18S and 16S rRNA genes amplicon generation for eukaryotic and prokaryotic metabarcoding

DOI:10.17504/protocols.io.qwhdxb6

Type: Protocol

Proper Citation

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dx.doi.org/10.17504/protocols.io.qwhdxb6

Protocol Information

URL: <https://dx.doi.org/10.17504/protocols.io.qwhdxb6>

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Group: Tara Oceans

Summary: This protocol describes the 18S and 16S rRNA genes amplicon generation for eukaryotic and prokaryotic metabarcoding for the Tara Oceans expedition and is part

of Viral to metazoan marine plankton nucleotide sequences from the Tara Oceans expedition. Figure 1: Overview of -omics analysis strategy applied on Tara Oceans samples.

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External URL: <https://www.nature.com/articles/sdata201793#methods>

Version: 1

Publication Date: 2020

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Record Creation Time: 20210329T220538+0000

Record Last Update: 20210329T221053+0000

Data and Source Information

Source: