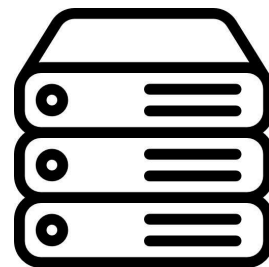


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Bioinformatic procedure for shotgun metagenomic analysis of human saliva microbiota

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We use this protocol and it's working

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Disclaimer

NA

Abstract

This protocol describes the bioinformatical procedure for the study of human saliva microbiota.

Attachments



Bioinformatic_proced...

167KB

Guidelines

NA

Materials

NA

Safety warnings

 NA

Ethics statement

NA

Before start

NA



General description

- 1 *Read Quality check*
- 2 *Mapping and microbial gene count procedure*
- 3 *Metagenomic Pangenome Species MSP profiles*

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