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## Axenic Diatoms cultures protocol V.3

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## **ABSTRACT**

Axenic cultures protocol

Axenicity of cultures is obtained by multi-antibiotic treatment: by adding antibiotics directly to the medium in which the strain will be inoculated. After a variable amount of days we refresh the coltures with new medium.

## antibiotics concentration

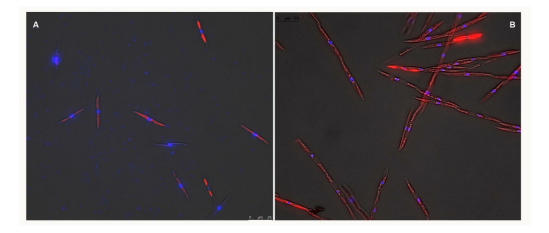
- The antibiotics used are streptomycin, penicillin and ampicillin; We used the following final concentration:
  - Streptomycin 0,5 mg/ml
  - Penicillin 0,05 mg/ml
  - Ampicillin 0,03 mg/ml
- 2 Add antibiotics in F/2 medium. Start from 20/50.000 cells in 25 ml of F/2 medium with antibiotics. Refresh 3 times the cultures every 3 days with new medium with antibiotics.
- 3 Transfer the cultures into a variable higher volume flask containing F/2 medium and antibiotics. Arrive progressively at the final volume.

## contamination test

4 Assess axenicity of the strains by fluorescence microscopy using DAPI staining (4',6-diamidino-2-phenylindole-a DNA stain).

(DAPI stains DNA, if bacteria are present in the culture bacterial nucleoids can be visualised as fluorescent spots).

To fix cells use neutralized formaldehyde 1.6% DAPI final concentration 1:1000



Pseudo-nitzschia multistriata fluorescence microscopy images using DAPI staining.

- A. Normal culture with bacteria
- B. Axenic culture without bacteria.

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