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Conjugation of Thalassiosira pseudonana Version 2

Jernej Turnsek

Abstract

This protocol was used to express a gene encoding a silaffin precursor TpSil3p-APEX2 fusion protein in *T. pseudonana* strain <u>CCMP1335</u> via conjugation largely following Karas et al. (2015). In addition to the protocol I am listing some of my observations working with this diatom species. Please check the 'Before start' and 'Guidelines' sections.

1. B. J. Karas *et al.*, Designer diatom episomes delivered by bacterial conjugation. *Nat. Commun.* **6**, 6925 (2015).

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Guidelines

[1] 'Crashing phenomenon' in T. pseudonana cultures

In my experience *T. pseudonana* CCMP1335 has to be subcultured frequently - every ~2 weeks - to avoid 'crashing'. I usually dilute my cultures 1:10 in fresh L1 medium. By 'crashing' I mean sudden take over of what I believe is a naturally associated bacterium in the culture. This usually happens in the stationary phase over the course of 1 or 2 days during which brown diatom culture becomes milky due to bacteria. It could be a standard contamination issue, but I've pretty consistently observed this phenomenon even after ordering completely fresh stocks from the NCMA. I analyzed the bacterium via 16S sequencing and some simple growth experiments and confirmed it was a *Pseudomonas* species. You can read more about how I've done this here.

Interestingly, this T. pseudonana-Pseudomonas interplay was already observed in the late 1970's.

[2] Episomes for *T. pseudonana* conjugation available upon request

All episomes used in my conjugation experiments including their maps are available upon request. Send an email to turnsek@fas.harvard.edu, turnsek@ucsd.edu or jturnsek@jcvi.org and we'll go from

there/I can provide more information. Here is a list of available episomes:

pTpPuc3 - empty nourseothricin resistance cassette-containing episome for *T. pseudonana* conjugation

eGFP - 2 different enhanced GFP constructs

<u>DreidelTeal</u> - blue chromogenic protein

TpSil3p-eGFP - frustule-associated silaffin precursor protein TpSil3p tagged with eGFP

TpSil3p-APEX2-FLAG - frustule-associated silaffin precursor protein TpSil3p tagged with APEX2 and FLAG

TpSil3p-APEX2-6xHis - frustule-associated silaffin precursor protein TpSil3p tagged with APEX2 and 6xHis

NOTE:

All the genes above were cloned into pTpPuc3 backbone via Gibson Assembly starting 36 nucleotides downstream of the HIS3 gene. They are flanked by a constitutive fcp promoter and an associated terminator. Here are their sequences in FASTA format:

>Tp_fcp_promoter

GCGCTTTTTCCGAGAACTCCCCATAAGTCAACGGCTCCAATCAAGAATGTATCCGACAACGGCGAGCATAGCAACGGCCGTCTTTGGAGTAGAATCATCATGTTGTGGATGAATACACAGA

TGAATGACATTAAAAGCATGAACATGTTAGAGAGTAGGAGGTAGAGATTGATATGGTAGCATTGCGATGTTTGT
TTTTGGTCAGCATATGATGAGTGGATACCAATATGATGAAAGTTGAATCT

TGGAGGTAGAAACTCATCCCAGTTGAGAAGAAGTGAAGGCAGTGGCGGTGGCGAAAGCAGAGGCAACGAGGA CAGACTTCCTGTGGGTTGATGCAACGAATATTTCCAGAAGGAGAAG

CGATCATTCACCTCCACACTGCAACACACGGTACTTCTTCCGCGGCAGGTCTCTGTCGCCATTCTCTTGTCCTG
TTGTTGGCTGTGAGACGACGACAACGACAACGTTTCACAAAAGGG

AGTTCCTTTAACGAGATATGTTTTTTATAAAGAGTCCCAATAGAAAGACAAATTGATTCCTCCGTGCAAACGCGC AAATAAACACCACGTCCATTATATCCATATCTTTCAGAGTATCCAACAAGT GTTGAAGGACAGGTAGTTGAAGTAACGTATCTTCCCCCCTCGACTGGATCCATCAACAAGGCGAACAAATCCATTCAACCTCTCATAAATTATCTGATTTACCAAACC

>Tp_fcp_terminator

TGTCCTCTGTCAGGAGTAGATAGTAGATGTTCTTTTTAAACTAAAATGCTAACTGTTCCGAATTCCTCATCGCAGC TAATCCGTACATCAAAAGACAAAATGCTAGGTATGTGTACTACATCTCC

TGTTGCTAGATAAGACATATGATAGGAAACACCATCAATAGTCATTGTAGCTTTACTTATACTACGCATTTGCACTTTCCCCTGAGTGGCAGAGGCGCATTGAGAAAATCGATCTCAACATAG

TTTATGTAGCATCCCTAGATCCATTACTTTAAGTCTCCTTCGTCTTTGGTGTAGGCATGTTGGACACAACGAGG
TAAAACACAACAACAACAATGTGTCCAGCAAAGTAGTAGCTGCTCCAGT

TCT

Before start

Preparation of donor bacteria

- 1. Any *E. coli* strain will probably work just fine, but I stuck with the one from the original publication: <u>EPI300</u> (Epicentre, catalog # EC300110).
- 2. Electroporate pTA-Mob into EP300 cells and select on LB agar plates w/ 10 or 20 μ g/mL gentamycin.
- 3. Make pTA-Mob-containing strain chemically competent.
- 4. Tranform your engineered pTpPuc3 cargo plasmid into pTA-Mob-containing cells.
- 5. Select double transformants on LB agar plates with 10 or 20 μ g/mL gentamycin and 50 μ g/mL kanamycin.
- 6. Store plates at 4 °C.

Preparation of L1 medium and L1 1% agar plates

I was using <u>L1 Medium Kit</u> and <u>Gulf of Maine Seawater</u>, both from NCMA, to make my liquid medium and plates.

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L1 medium

I would usually make batches of 1L by combining seawater, N, P and Si sources from the kit followed by autoclaving, cooling down to ~ 50 $^{\circ}$ C and adding vitamins and antibiotics. The resulting medium had pH ~ 8.5 . Before starting a diatom culture I would adjust the pH with HCl to ~ 8.0 .

NOTE:

The resulting medium contain some precipitates. You can use a combination of microwaving and filter sterilization instead which should prevent precipitation.

1/2xL1 1% agar plates w/ 5% LB

I would always prepare 0.5L and make \sim 12 plates 40 mL each. For 0.5L combine and mix seawater, N, P and Si sources from the kit, add agar and 5% LB. Autoclave, cool down to \sim 50 $^{\circ}$ C and add vitamins.

NOTES:

- 5% LB is v/v not w/v. I had success preparing plates w/v, but v/v is what you should use.
- 1/2xL1: cut all N, P, Si and vitamin volumes from the kit in half.
- I had success with Bacto Agar and Sigma's high purity agar.

1/2xL1 1% agar plates w/ 100 μg/mL nourseothricin

Same as above except the LB part; add nourseothricin when cooled down to ~ 50 $^{\circ}$ C.

NOTE:

I observed a lot of background - false positive - colonies on plates with only 50 μ g/mL nourseothricin so I switched to 100 μ g/mL. You can read more about nourseothricin on the <u>following website</u> which I found very useful.

Protocol

Growth and preparation of E. coli donor

Step 1.

Pick colonies from your Gent+Kan plates & inoculate 10 mL LB medium. Grow overnight.

NOTES

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Start growing colonies early in the afternoon the day before conjugation to get enough biomass. \sim 16-20 hrs should suffice. Expect OD₆₀₀ \sim 2 next morning.

Growth and preparation of E. coli donor

Step 2.

Measure OD₆₀₀ and start a 150 mL LB subculture (starting OD₆₀₀ either 0.05 or 0.1).

Growth and preparation of E. coli donor

Step 3.

Grow at 37 C until OD₆₀₀ reaches 0.3-0.4.

Growth and preparation of E. coli donor

Step 4.

Centrifuge at 4,000 rpm, 10 °C, for 10 min.

O DURATION

00:10:00

Growth and preparation of E. coli donor

Step 5.

Decant supernatant and resuspend in 800 µL SOC.

Growth and preparation of T. pseudonana cells

Step 6.

T. pseudonana was cultured in L1 medium with moderate shaking (130 rpm), constant light and 18 °C. L1 medium described in 'Before start' section.

NOTES

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- I've never tested if cell density before spinning cells down matters with respect to final conjugation outcome same as the Alverson Lab did for P. tricornutum. They observe $\sim 8 \times 10^6$ cells/mL to be a sweet spot. In my hands spinning cells down at $\sim 4-8 \times 10^6$ cells/mL seemed to work fine. According to my information harvesting T. pseudonana at $\sim 0.8 \times 10^5$ cell/mL works best for biolistic experiments.
- I counted cells with BioRad's TC20 automatic cell counter.

Growth and preparation of T. pseudonana cells

Step 7.

Spin down 2 x 10⁸ cells at 4000 rpm, 10 °C, for 10 min.

O DURATION

00:10:00

Growth and preparation of T. pseudonana cells

Step 8.

Decant supernatant and resuspend pellet in 1 mL L1 medium.

Conjugation

Step 9.

Mix 200 μ L T. pseudonana cells and 200 μ L E. coli cells in a 1.5 mL tube.

Conjugation

Step 10.

Pipette up and down a few times.

Conjugation

Step 11.

Plate on 1/2xL1 1% agar plates w/ 5% LB.

NOTES

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Make sure the plates are dry. Leave them open at 37 $^{\circ}$ C for an hour or so before plating diatom-bacteria co-cultures.

Conjugation

Step 12.

Incubate in dark at 30 °C for 90 minutes

O DURATION

01:30:00

Conjugation

Step 13.

Move plates to standard diatom growth conditions - 18 °C and constant light for 4 hours

O DURATION

04:00:00

Conjugation

Step 14.

Add 1 mL L1 medim and scrape with a <u>cell scraper</u> or <u>L spreader</u>.

Selection

Step 15.

Expect to recover 500 µL co-culture suspension after scraping.

Selection

Step 16.

Plate 200 μL of the resulting suspension on pre-dried 1/2xL1 1% agar plates w/ 100 $\mu g/mL$ nourseothricin.

Selection

Step 17.

Leave at 18 °C and constant light until colonies appear - 2 weeks.

Screening

Step 18.

Here are a few ways to confirm the presence and expression of your heterologous gene in resulting colonies:

1. Growth under selection pressure.

Make sure colonies are able to grow in liquid L1 medium with 100 μ g/mL nourseothricin (Nou100). Pick colonies with a small tip or better a toothpick and inoculate 0.5 mL L1 medium. Once you observe growth subculture in larger volume.

2. PCR

- Use 1 µL of diatom culture as a template to amplify your expression cassette.
- Confirm the absence of donor DNA by amplifying *E. coli*-specific genes.
- Confirm the absence of live donor cells by plating some diatom culture on LB plates w/o antibiotics.

NOTE:

Any remaining donors cells and donor DNA are gone after a few liquid subcultures.

3. RT-PCR

- Purify total RNA from Nou100 diatom culture, convert it to cDNA and use it to run a PCR with heterologous gene-specific primers.
- Example of a result with controls can be found here.

4. Episome	recovery
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- Perform a diatom miniprep as described by Karas et al.
- Transform E. coli with diatom-derived DNA.
- Select on LB agar plates with 50 μg/mL kanamycin.
- Miniprep, digest and analyze on a gel.
- 5. Western blot
- 6. Protein-specific assays
- enzymatic assay
- microscopy

- ...

I've had success with 1, 2 and 3, tried 4, 5 and 6 without success.