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### C Loop L1 (odd level) Bsal type IIS cloning into pCk vectors V.2

Version 1 is forked from Loop L1 (odd level) type IIS cloning - pCk-ye vectors

DOI

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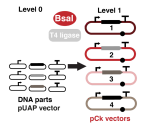
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Protocol status: Working We use this protocol and it's working

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#### Abstract

Protocol based on:

# Pollak B, Cerda A, Delmans M, et al (2019) Loop assembly: a simple and open system for recursive fabrication of DNA circuits. New Phytol 222:628–640

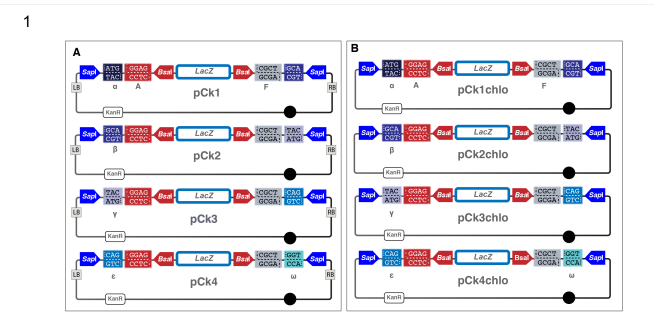
https://doi.org/10.1111/nph.15625

Materials

MATERIALS

- 🔀 Bsal 5,000 units New England Biolabs Catalog #R0535L
- X T4 DNA Ligase 20,000 units New England Biolabs Catalog #M0202S
- 🔀 Sterile water
- 🔀 BSA, molecular biology grade, 20 mg/ml New England Biolabs Catalog # B9000S
- X 10X NEB T4 DNA ligase buffer **New England Biolabs**

#### Loop pCk vectors



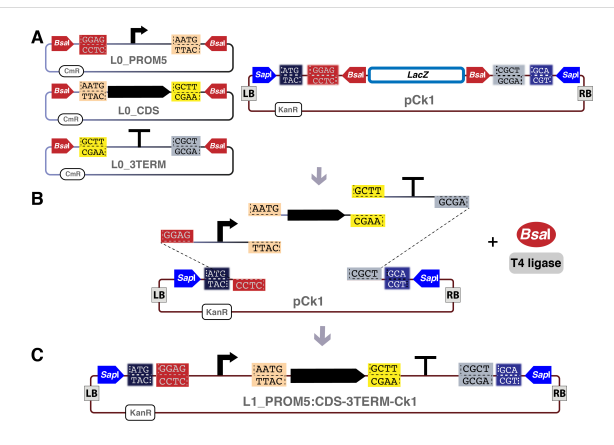
## Loop vectors for nuclear transformation: pCks (A) and for chloroplast transformation pCkchlo (B).

Loop fusion sites in the pCk vectors to assemble different L0 parts into a L1 construct using a pCk vector and Bsal are: A (GGAG) and F (CGCT).

Loop fusion sites in the pCk vectors to assemble different L1 constructs into a L2 construct using a pCs vector and Sapl are: a (ATC), b (GCA), d (TAC), e (CAG) and o (GCT). Left (LB) and right border (RB) repeats from nopaline C58 T-DNA for Agrobacteriummediated nuclear transformation. KanR: kanamycin bacterial resistance cassette. LacZ: lacZα cassette for blue-white screening of colonies.

Example of assembly of L0 parts into a transcription unit (L1)

2



Loop assembly of multiple L0 parts into a transcription unit (L1) using a pCk plasmid and Bsal.

#### Protocol for assembly of L0 parts into a transcription unit (L1)

- Determine the concentrations of each DNA plasmid needed (L0 plasmids and pCk acceptor plasmid) by spectrophotometry (Nanodrop).
  In the example in step 2, determine concentration of plasmids L0\_PROM5, L0\_CDS, L0\_3TERM and pCk1.
- 4 Prepare aliquots for each plasmid at a concentration of 15 nM for the L0 plasmids and of 7.5 nM for the acceptor pCk vector. With this final concentration, 1 μL of each plasmid is added to the plasmids mix (see step 6).

To calculate the concentration in  $ng/\mu L$ :

- For a final concentration of 15 nM, the concentration in [ng/u] equals N (the length in bp of the plasmid) divided by 110. This is an approximation of the formula:

 $15\cdot10^{-9}$ mol/L x ((607.4 x N ) + 157.9)g/mol x 10^{-6}L/µL x 10^9ng/g = concentration (ng/µL)

- For a final concentration of 7.5 nM, the concentration in [ng/ul] equals N divided by 220.

5 Prepare Loop assembly Level 1 reaction master mix (MM) according to Table , if four or less number of L0 parts are assembled into a pCk vector (otherwise see step 8)

Comp onent s	Volu me (µL)
Sterile water	3
10x T4 ligase buffer (NEB)	1
1 mg/m L bovin e serum album in (NEB)	0.5
T4 DNA ligase at 400 U/μL (NEB)	0.25
10 U/µL Bsal (NEB)	0.25
Final volum e	5

- 6
- Prepare plasmids mix by adding in a 0.2 mL tube: 1  $\mu$ L of each L0 plasmid , 1  $\mu$ L of the pCk vector (see step 4), and sterile water up to 5  $\mu$ L. Mix well.

When 4 L0 parts are assembled into a pCk plasmid, the volume of the plasmid mix is 5  $\mu$ L, and thus no volume of water is added.

- 7 Add 5  $\mu$ L of MM (step 5) to the 5  $\mu$ L of plasmids mix (step 6), to a final volume of 10  $\mu$ L. Mix well.
- 8 If more than 4 L0 parts are to be assembled into a pCk vector, reduce the water volume in the MM by 1  $\mu$ L (step 5) for each extra 1  $\mu$ L of DNA part added in the plasmids mix (step 6).
- Place samples in a thermocycler and use the following program:
  Assembly: 26 cycles of 37 °C for 3 min and 16 °C for 4 min.
  Termination and enzyme denaturation: 50 °C for 5 min and 80 °C for 10 min.
- 10 Transform 20 μL of chemically competent E. coli cells (transformation efficiency of 1 × 10<sup>7</sup> transformants/μg plasmid DNA) using 2 μL of the Loop assembly reaction and then plate on LB agar plates containing 50 μg/mL kanamycin and 40 μg/mL of X-gal for blue-white screening.
- 11 Incubate overnight at 37 °C.
- Colonies with white color are likely to contain an L1 insert cloned into the pCk vector (In the example in step 2: PROM5:CDS-3TERM)
  Blue color colonies will contain undigested pCk vector with LacZ
- 13 Confirm the presence of the correct insert with Sanger sequencing using the primers pC\_F (GCAACGCTCTGTCATCGTTAC) and pC\_R (GTAACTTAGGACTTGTGCGACATGTC) for pCk vectors, and pC\_F and pC\_R2 (CAATCTGCTCTGATGCCGCATAGTTAAG) for pCkchlo vectors.