



•





High Efficiency Transformation Protocol (C2987I) V.2

New England Biolabs¹

¹New England Biolabs

1

dx.doi.org/10.17504/protocols.io.bddui26w

New England Biolabs (NEB)

Tech. support phone: +1(800)632-7799 email: info@neb.com



This is the protocol for C2987I cells. If you are using the C2987H cells, please refer to <u>this protocol</u>.

DOI

dx.doi.org/10.17504/protocols.io.bddui26w

https://www.neb.com/protocols/0001/01/01/high-efficiency-transformation-protocol-c2987

New England Biolabs 2022. High Efficiency Transformation Protocol (C2987I). **protocols.io**

https://dx.doi.org/10.17504/protocols.io.bddui26w New England Biolabs

Transformation, Bacterial, C2987I

_____ protocol,

Mar 08, 2020

Feb 14, 2022

33940



Transformation Protocol Variables

Thawing: Cells are best thawed on ice and DNA added as soon as the last bit of ice in the tube disappears. Cells can also be thawed by hand, but warming above 0°C will decrease the transformation efficiency.

Incubation of DNA with Cells on Ice: For maximum transformation efficiency, cells and DNA should be incubated together on ice for 30 minutes. Expect a 2-fold loss in transformation efficiency for every 10 minutes this step is shortened.

Heat Shock: Both the temperature and the timing of the heat shock step are important and specific to the transformation volume and vessel. Using the transformation tube provided, 30 seconds at 42°C is optimal.

Outgrowth: Outgrowth at 37°C for 1 hour is best for cell recovery and for expression of antibiotic resistance. Expect a 2-fold loss in transformation efficiency for every 15 minutes this step is shortened. SOC gives 2-fold higher transformation efficiency than LB medium; and incubation with shaking or rotating the tube gives 2-fold higher transformation efficiency than incubation without shaking.

Plating: Selection plates can be used warm or cold, wet or dry without significantly affecting the transformation efficiency. However, warm, dry plates are easier to spread and allow for the most rapid colony formation.

DNA Contaminants to Avoid

Α	В
Contaminant	Removal Method
Detergents	Ethanol precipitate
Phenol	Extract with chloroform and ethanol precipitate
Ethanol or Isopropanol	Dry pellet before resuspending
PEG*	Column purify or phenol/chloroform extract and ethanol precipitate
DNA binding proteins (e.g. Ligase)	Column purify or phenol/chloroform extract and ethanol precipitate

^{*} Ideally, DNA for transformation should be purified and resuspended in water or TE. However, up to 10 µl of DNA directly from a ligation mix can be used with only a two-fold loss of transformation efficiency. Where it is necessary to maximize the number of transformants (e.g. a library), a purification step, either a spin column or phenol/chloroform extraction and ethanol precipitation should be added.

MATERIALS

SOC Outgrowth Medium - 100 ml New England

Biolabs Catalog #B9020S

№ NEB 5-alpha Competent E.coli (High Efficiency) - 6x0.2 ml New England

Biolabs Catalog #C29871

Please refer to the Safety Data Sheets (SDS) for health and environmental hazards.



1 Thaw a tube of NEB 5-alpha Competent *E. coli* cells § **On ice** until the last ice crystals disappear.

Cells are best thawed on ice and DNA added as soon as the last bit of ice in the tube disappears. Cells can also be thawed by hand, but warming above 0°C will decrease the transformation efficiency.

2

Mix gently and carefully pipette $\blacksquare 50~\mu L$ cells into a transformation tube & On ice.

3

Add $\Box 1 \mu L - \Box 5 \mu L$ containing $\Box 1 pg - \Box 100 ng$ of plasmid DNA to the cell mixture.

4 🔀

Carefully flick the tube **4-5 times** to mix cells and DNA. **Do not vortex.**

5

Place the mixture § On ice for © 00:30:00. Do not mix.

For maximum transformation efficiency, cells and DNA should be incubated together on ice for 30 minutes. Expect a 2-fold loss in transformation efficiency for every 10 minutes this step is shortened.

6 Heat shock at exactly § 42 °C for exactly © 00:00:30. Do not mix.

Both the temperature and the timing of the heat shock step are important and specific to the transformation volume and vessel. Using the transformation tube provided, 30 seconds at 42°C is optimal.

7

Place § On ice for © 00:05:00. Do not mix.

8

Pipette $\mathbf{\square}950~\mu L$ room temperature SOC into the mixture.

9

Place at § 37 °C for © 01:00:00, shaking vigorously (@ 250 rpm) or rotating.

Outgrowth at 37°C for 1 hour is best for cell recovery and for expression of antibiotic resistance. Expect a 2-fold loss in transformation efficiency for every 15 minutes this step is shortened. SOC gives 2-fold higher transformation efficiency than LB medium; and incubation with shaking or rotating the tube gives 2-fold higher transformation efficiency than incubation without shaking.

10 Warm selection plates to § 37 °C.

Selection plates can be used warm or cold, wet or dry without significantly affecting the transformation efficiency. However, warm, dry plates are easier to spread and allow for the most rapid colony formation.

11 🗸

Mix the cells thoroughly by flicking the tube and inverting.

12 Perform several 10-fold serial dilutions in SOC.

13

Spread $\Box 50 \mu L - \Box 100 \mu L$ of each dilution onto a selection plate.

14

Incubate @ Overnight at § 37 °C.

Alternatively, incubate at § 30 °C for 24-36 hours or § 25 °C for (348:00:00).

