

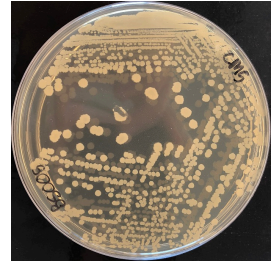
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# Transformation of heterolous DNA in *Bacillus subtilis*

 In 1 collection

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**Protocol status:** Working

**We use this protocol routinely and it works for us**

**Created:** March 13, 2020

**Last Modified:** October 28, 2020

**Protocol Integer ID:** 34195

**Keywords:** B. subtilis, Bacillus, Bacillus subtilis, Transformation, Natural competence



## Abstract

*B. subtilis* is a gram-positive bacteria used by both academia and industry as a protein production workhorse. This is due to its' excellent fermentation properties, high production titers, and capacity to secrete proteins into the extracellular medium.

This protocol describes transformation of *B. subtilis* by natural competence. The method utilizes the natural stress-induced competence of *B. subtilis* to take up heterologous DNA. The protocol requires the cells to be grown for a specific amount of time in starvation media (SM). The protocol is adapted from Vojcic, L., Despotovic, D., Martinez, R., Maurer, K., & Schwaneberg, U. (2012). An efficient transformation method for *Bacillus subtilis* DB104. *Applied Microbiology and Biotechnology*, 94(2), 487–493. <https://doi.org/10.1007/s00253-012-3987-2>.

## Guidelines

This protocol works for a range of *B. subtilis* strains (e.g. 168, WB800, PY79, KO7, KO7-S, and derivatives), although the protocol might need optimization for heavily growth impaired strains. For protein production, it is generally recommended to use a protease deficient strains such as WB700, WB800, KO7 or KO7-S.

## Materials

### MATERIALS

⊗ Glucose **P212121 Catalog #**Glucose

⊗ Sodium citrate **P212121**

⊗ Ammonium Sulfate **P212121**

⊗ Potassium phosphate (dibasic) **P212121**

⊗ Calcium Chloride

⊗ Potassium dihydrogen phosphate **Merck MilliporeSigma (Sigma-Aldrich) Catalog #**NIST200B

⊗ Magnesium Sulfate **Amresco Catalog #**0338

⊗ Yeast extract

⊗ Bacto&trade; Casamino Acids **Thermo Fisher Catalog #**223020

## Safety warnings



This protocol describes the construction of GMO classified organisms. Make sure that the local GMO and safety legislations are respected.

## Before start

Make sure you have your recipient strain freshly streaked on an agar plate.

## Preparation of stock solutions

### 1 Mix 10xSM1 stock

#### 1.1 Weigh the following in a 100mL blue cap bottle:

- 2 g ammonium sulphate
- 14 g dipotassium hydrogen phosphate
- 6 g potassium dihydrogen phosphate
- 0.7 g sodium citrate
- 0.2 g magnesium sulfate heptahydrate
- 2 g yeast extract
- 0.25 g casamino acids

#### 1.2 Add MQ water to 100 mL and mix to dissolve the ingredients

### 2 Mix 1xSM2 stock

#### Note

SM2 is made as a 1x stock since it otherwise precipitates

#### 2.1 Weigh the following in a 1L blue cap bottle:

- 2 g ammonium sulfate
- 14 g dipotassium hydrogen phosphate
- 6 g potassium dihydrogen phosphate
- 0.7 g sodium citrate
- 0.8 g magnesium sulfate heptahydrate
- 1 g yeast extract
- 0.1 g casamino acids

#### 2.2 Add MQ water to 1 L and mix to dissolve the ingredients

### 3 Autoclave the stocks and store at Room temperature or 4 °C



### 4 Prepare a 200g/L glucose and a 1M CaCl<sub>2</sub> stock, separately




### Note

Glucose is prepared separately to avoid Millard's reaction.  $\text{CaCl}_2$  is prepared separately since it makes the SM2 stock precipitate.







## Overnight culture

- 5 Prepare fresh SM1 media from the stock by mixing 1 part 10xSM1 media stock with 9 parts sterile MQ water and 25uL 200g/L glucose stock per mL media (to final concentration of 5g/L).
- 6 Inoculate 10mL SM1 media in a 50mL falcon tube with a single colony from a freshly streaked plate
- 7 Incubate  Overnight at  37 °C with 250RPM shaking


### Note

Make sure to not incubate the overnight culture for longer than  16:00:00 . Using an overnight culture that has been incubating for longer than this, often results in low transformation efficiencies

## Transformation



- 8 Measure OD<sub>600</sub> of the O/N culture
- 9 Dilute the O/N culture in  10 mL SM1 media to a final OD<sub>600</sub> of 0.5 in a  250 mL Erlenmeyer flask
- 10 Grow culture at  37 °C at 250 RPM shaking for  03:00:00
- 11 Add  10 mL SM2 medium to the culture
- 12 Add  250 µL 200g/L glucose stock to the culture



- 13 Swirl the culture around, and add  45  $\mu\text{L}$  1M  $\text{CaCl}_2$  stock while the media is still in motion

#### Note

The swirling is done since high local concentrations of  $\text{CaCl}_2$  makes the media precipitate. This way the  $\text{CaCl}_2$  is mixed in the media before it precipitates




- 14 Grow culture at  37  $^{\circ}\text{C}$  at 250 RPM shaking for  02:00:00

#### Note

After this step the cells are competent for approximately  01:00:00

- 15 Distribute the cells in  500  $\mu\text{L}$  aliquotes in  2 mL eppendorf tubes

#### Note



The protocol can be paused at this point, by adding  250  $\mu\text{L}$   50 % volume glycerol to the aliquotes and freezing them in a  -80  $^{\circ}\text{C}$  freezer. The protocol can be restarted by thawing the cells on ice and moving on the next step. This can also be used to prepared stocks of strains that are used often




- 16 Add 250ng - 1pg of plasmid DNA to the aliquotes

#### Note

Some protocols call for linearized plasmids or PCR products, although we have found that circular plasmids work as well




17 Incubate the aliquotes in a thermoblock for  00:30:00 at  37 °C with 800 RPM shaking


18 Add  300 µL LB media and recover the cells for at least  02:00:00 at  37 °C with 800 RPM shaking

#### Note

The longer duration of incubation the better

19 Plate up to  200 µL of each aliquot on LB agar plates with appropriate antibiotics

#### Note

In order to increase the number of colonies on the transformation plates, the aliquotes can be centrifuged for  6000 x g, 00:02:00 and reinoculated in a smaller volume.

20 Incubate the plates  Overnight at  37 °C