Bacillus subtilisFood Grade Expression System



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Introduction

Gram-positive bacteria are well known for their contributions to agricultural, medical, and food biotechnology, and for the production of recombinant proteins. Among them, **Bacillus subtilis** has been developed as an attractive host because of several reasons:

- It is non-pathogenic and is considered as a GRAS organism (generally regarded as safe), which means it can be used for food production;
- It has no significant bias in codon usage;
- It is capable of secreting functional extracellular proteins directly into the culture medium (at present, about 60 % of the commercially available enzymes are produced by *Bacillus* species):
- A large body of information concerning transcription, translation, protein folding and secretion mechanisms, genetic manipulation, and large-scale fermentation has been acquired, and the complete genome sequence of *B. subtilis* is available.

To increase the yield of heterologous proteins and to simplify operational processes, a wide variety of highly sophisticated plasmid-based expression systems have been developed. The here described *Bacillus* Food Grade Expression System was created to make the advantages of a Bacillus expression system also accessible to areas where antibiotic resistance gene markers are prohibited (e.g., food and feed industry). The *Bacillus* Food Grade Expression System enables stable vector-based large scale heterologous protein production by an alternative selection, without antibiotics.

The Bacillus Food Grade Selection System provides the following features:

- Stable high- or low-level expression without addition of any antibiotics
- All DNA contained in the final expression system is derived from B. subtilis
- No endotoxins are produced
- No inclusion bodies are formed
- Protease-deficient strain for producing secretory enzymes is available

The Food Grade Selection System

The **Bacillus Food Grade Selection System** is based on the interplay of an endogenous **Bacillus toxin EndoA** and its **antitoxin EndoB**.

EndoA (encoded by ydcE) is an endoribonuclease that specifically cleaves mRNA at a five Base U \sqrt{ACAU} sequence (Pellegrini et~al., 2005, Park et~al., 2011). During normal growth conditions EndoA is inactivated by forming a heterohexameric complex with its cognate antitoxin EndoB (encoded by ydcD; Simanshu et~al., 2013). Since the antitoxin is relatively unstable, it is essential for the cell to continuously produce sufficient amounts of EndoB to inactivate the more stable toxin. These characteristics are utilized for the here described selection system.

The mechanism of the selection system

The antitoxin encoding gene *ydcD* is located within the expression vector under the control of a constitutive *Bacillus* promoter whereas the toxin expression cassette is integrated into the genome and is controlled by a xylose-inducible promoter. If the expression of the toxin is induced by xylose but the plasmid with the encoded antitoxin gets lost, no more antitoxin

can be produced, leading to an active endoribonuclease that results in inhibition of protein synthesis, cell growth arrest, and finally dying of the cell.

This expression system has been proven to be very stable, with no plasmid loss observed in 100 generations. Compared with traditional antibiotic-dependent expression systems, this system results in greater biomass and higher titers of the desired products. This has been shown for the expression of the green fluorescent protein and the metabolic product hyaluronan, respectively (Yang *et al.*, 2016).

The expression vectors

Dependent on the particular requirements there can be chosen between two *B. subtilis* expression vectors **pTTB1** and **pTTB2**. These vectors differ in their origin of replication and copy number concerning *B. subtilis*. **pTTB1** is a low copy number vector, replicating via theta replication modus, whereas **pTTB2** is high copy number, replicating by rolling circle mechanism.

Both vectors share the following features:

- For easier handling the vectors are designed as *B. subtilis / E. coli* shuttle vectors. The parts of the vector used for cloning with *E. coli* (*E. coli* origin ColE1 derivative and ampicillin resistance cassette Amp) can be eliminated afterwards by restriction enzyme cleavage and religation of the vector (for details see below). This technique connects the advantage of easy cloning (with *E. coli*) with the food grade property of *B. subtilis*.
- For food grade selection with *B. subtilis* the antitoxin-encoding gene *ydcD* is included under control of a constitutive *Bacillus* promoter (the strong P_{ylxM} promoter in pTTB1 and the weaker P_{aadD} promoter in pTTB2).
- A multiple cloning site for cloning the gene of interest downstream of the constitutive promoter P₄₃ is provided.

The food grade Bacillus strains

For expressing the gene of interest under food grade conditions, two B. subtilis strains are available; TEA and WEA-C. Both strains contain the toxin expression cassette ydcE under control of the xylose-inducible promoter P_{xyl} , whereas the former ydcDE operon (containing the toxin and antitoxin gene) is deleted. The strain B. subtilis TEA is based on B. subtilis 168 Marburg and is recommended for intracellular protein expression and pathway engineering. B. subtilis WEA-C originated from the sixfold extracellular protease-deficient strain B. subtilis WB600 and is particularly constructed for secretory protein production. Both strains are suitable hosts for both the food grade low copy expression vector pTTB1 and the high copy expression vector pTTB2.

TEA: $trpC2 \ ydcDE::P_{xyl}-ycdE$

WEA-C: nprE aprE epr bpr mpr::ble nprB::bsr ydcDE::Pxyl-ycdE; CmR

Please note: this strain carries resistance to chloramphenicol!

Storage and Handling Instructions

Storage and handling of plasmids

Plasmids are supplied lyophilized. Upon receipt, add 100 µl distilled water (final concentration 0.1 µg/µl) and incubate at 50 °C for 5 minutes. Vortex for 1 minute and store at -20 °C. Please note that all plasmids of this system are *E. coli / B. subtilis* shuttle vectors.

Storage and handling of Bacillus strains

The *Bacillus* strains are supplied as frozen cultures and shipped on dry ice. Store the stock at -80 °C. For propagation remove tube from freezer, scratch off some material from the surface of the frozen stock using a sterile loop. Streak onto an LB plate containing 0.5% glucose (see page 7), seal the plate with parafilm, and incubate at 37 °C overnight. *Bacillus* plates can be stored short-time at 4 °C. For preparing competent cells (see page 6) we recommend starting always with a freshly grown single colony. Please save transformed WEA and TEA strains at -80 °C. For protein expression start with one single colony of freshly transformed cells or use cells stored at -80 °C.

We recommend performing a functionality plate test of WEA and TEA before preparing new glycerol stocks: Use one single colony from isolation dilution and transfer it onto 1) LB agar with 1% D-xylose and 2) LB agar with 1% glucose. If there is growth on LB agar with glucose but not on LB agar with xylose the xylose regulated toxin expression is functional.

Protocols

Growth conditions

Detailed protocols for *E. coli* and *B. subtilis* molecular genetic handling (growth, transformation, etc.) can be found in the relevant laboratory manuals such as Sambrook and Russell (2001).

B. subtilis and E. coli can be grown aerobically at 37 °C in 2xYT (Bagyan et al., 1998) or LB medium. When culturing WEA and TEA strains (without vector) it is recommended to add 10 g/L glucose to the medium to keep the xylose promoter of the chromosomal toxin gene tightly closed.

Under optimal conditions the doubling time of *E. coli* is 20 min, of *B. subtilis* 30 min.

For selecting transformed *E. coli* cells, use 100 µg/ml ampicillin. For selecting transformed *Bacillus* TEA and WEA strains, add xylose to a final concentration of 10 g/L to the medium (plates and liquid culture). For protein expression in liquid culture alternatively a Sucrose/Xylose medium can be used (s. page 7).

Vector propagation and cloning the DNA fragment of interest

Since both expression vectors (pTTB1 and pTTB2) are designed as *E. coli / B. subtilis* shuttle vectors, we recommend using *E. coli* for plasmid propagation and for cloning the DNA fragment of interest. Please note: the gene of interest must be cloned together with a ribosome binding site (RBS). The RBS consists of the Shine-Dalgarno (SD) sequence and a spacer region (e.g., 7-8 bp) to the START codon. Statistical analyses revealed that most SD sequences in *B. subtilis* are very close to the consensus sequence AA AGG AGG (Rocha et al, 1999; for more information on RBS of *B. subtilis* see also Guiziou S, *et al* (2016), Popp, P.F. *et al.* (2017))

Follow standard protocols for propagation in *E. coli*, *E. coli* plasmid mini preparation, restriction endonuclease cleavages, ligation of the desired DNA fragment into the vector, and transformation of *E. coli* (Sambrook and Russell, 2001). For selecting successfully transformed *E. coli* cells use 100 µg ampicillin. To retain food grade properties both vectors

provide the option to remove the functional *E. coli* parts (i.e., ampicillin resistance cassette and origin for replication) after finishing the construct. These parts can be removed by cleavage with one single restriction enzyme (for pTTB1 use Spel, for pTTB2 use EcoRI) followed by purification of the larger fragment and religation. The resulting religated vector only contains DNA that is derived exclusively from *B. subtilis* (except for the cloned fragment).

Transformation of Bacillus subtilis

Preparation of competent Bacillus subtilis cells

The following protocol is adopted from Klein *et al.*, 1992.

- Prepare an overnight culture in 5 ml HS medium at 37 °C under vigorous shaking. For inoculation we recommend using one single colony grown on an LB agar plate.
- Measure the OD₆₀₀ (optical density at 600 nm) of the overnight culture and inoculate 50 ml HS medium to an OD₆₀₀ of 0.05; incubate under vigorous shaking at 37 °C.
- Record the growth curve.
- Immediately at transition of exponential to stationary growth phase start taking samples of 10 ml, each 15 min
- Add 1 ml of sterile glycerol (87 %) to each sample, mix, and leave for 15 min on ice.
- Fractionate into 1 ml aliquots, freeze in liquid nitrogen, and store at -80 °C.
- Check one aliquot from each time point with a reference plasmid DNA (see below) to identify the time point(s) yielding high level competent cells; discard the non or low competent aliquots.

Transformation of competent Bacillus subtilis cells

- Thaw one aliquot at 37 °C
- Use these cells to inoculate 20 ml LS medium
- Shake cells slowly in a 30 °C water bath to obtain maximal competence (about 2 h)
- Take 1 ml aliquots into a glass tube or a 2 ml plastic reaction tube, add 10 μl of 0.1 M EGTA, and incubate for 5 min at room temperature
- Add about 1 µg plasmid DNA and incubate for 2 h at 37 °C while well shaking (well mixing is important when using plastic reaction tubes)
- If glass tubes were used, transfer cell suspension into a 2 ml plastic reaction tube
- Centrifuge, discard supernatant carefully, and resuspend the cells in the residual liquid remaining on the pellet
- Plate on selective 2xYT plates containing 1% xylose (see page 7)
- Incubate at 37 °C overnight

Media and Solutions

LB medium: 10 g tryptone

5 g yeast extract

5 g sodium chloride (NaCl)

add distilled water to 1000 ml and autoclave

LB agar plates add 1.5 % agar to the LB medium before autoclaving

2xYT medium: 16 g tryptone 10 g yeast extract

5 g sodium chloride (NaCl)

add distilled water to 1000 ml and autoclave

(121 °C, 15 min)

2xYT / xylose agar plates: add 1.5 % agar to the 2YT medium before

autoclaving, let cool down to 70 °C and add 1 % sterile-

filtered D-xylose before pouring the plates

10x S-base (Spizizen's salt): 2 g (NH₄)₂SO₄

14 g K₂HPO₄

6 g KH₂PO₄

1 g sodium citrate

add distilled water to 100 ml and autoclave add 0.1 ml 1 M MgSO₄ after autoclaving

HS medium: 66.5 ml distilled water

10.0 ml 10x S-base

2.5 ml 20 % (w/v) glucose

5.0 ml 0.1 % (w/v) L-tryptophan

1.0 ml 2 % (w/v) casamino acids

5.0 ml 10 % (w/v) yeast extract (Difco) 10.0 ml 8 % (w/v) arginine, 0.4 % histidine

autoclave all components separately

tryptophan solution: sterile filtration

LS medium: 80.0 ml distilled water

10.0 ml 10x S-base

2.5 ml 20 % (w/v) glucose

0.5 ml 0.1 % (w/v) L-tryptophan

0.5 ml 2 % (w/v) casamino acids

5.0 ml 2 % (w/v) yeast extract (Difco)

0.25 ml 1 mM MgCl₂ 0.05 ml 1 mM CaCl₂

autoclave all components separately

tryptophan solution: sterile filtration

0.1 M EGTA dissolve 3.8 g EGTA in 50 ml distilled water

adjust the pH to 7.2 using 10 N NaOH add distilled water to 100 ml; autoclave

Sucrose / Xylose medium (1L)

20.0 g yeast extract,
1.3 g K₂SO₄,
6.2 g Na₂HPO₄
add 700 ml distilled water and autoclave,
let cool down,
add 100 ml 2% MgSO₄ x 7H₂O (w/v, sterile filtered)
add 100 ml of 70% sucrose (w/v, sterile filtered)
add 100 ml of 10% D-xylose (w/v sterile filtered)

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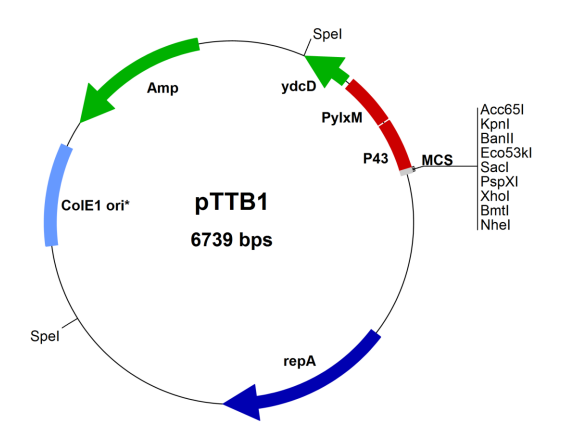
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Vector Maps

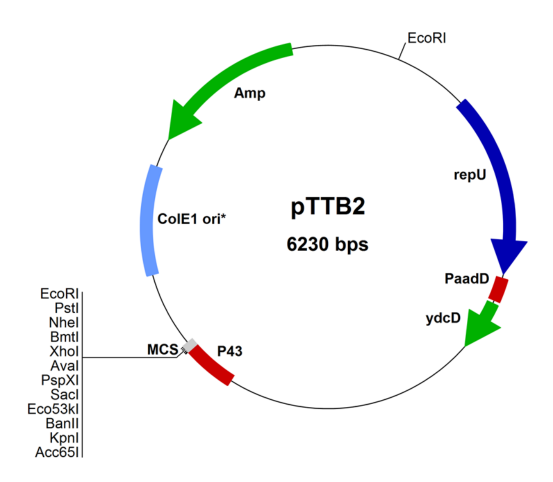
Vector map pTTB1 (low copy)



Туре	Start	End	Name	Description
Selectable genetic marker	724	443	ydcD	Antitoxin encoding gene
Promoter	1054	755	PylxM	Strong constitutive promoter for expression of ydcD
Promoter	1067	1365	P43	Constitutive promoter for expression of gene of interest
MCS	1366	1395	MCS	Multiple Cloning Site
Gene	2388	3422	repA	Gene of replication protein RepA (B. subtilis)
Origin of replication	4918	5517	ColE1 ori*	origin of replication (<i>E. coli</i>); ColE1 incompatibility group
Selectable genetic marker	6539	5679	Amp	Ampicillin resistance (E. coli)

The complete DNA sequence is available at our webpage www.mobitec.com.

Vector map pTTB2 (high copy)



Туре	Start	End	Name	Description
Gene	814	1818	repU	Gene of replication protein RepU (B. subtilis)
Promoter	1841	1969	PaadD	constitutive promoter for expression of ydcD
Selectable genetic marker	1987	2268	ydcD	Antitoxin encoding gene
Promoter	3674	3947	P43	constitutive promoter for expression of gene of interest
MCS	3948	3990	MCS	Multiple Cloning Site
Origin of replication	4409	5008	ColE1 ori*	origin of replication (<i>E. coli</i>); ColE1 incompatibility group
Selectable genetic marker	6030	5170	Amp	Ampicillin resistance (E. coli)

Please consider: EcoRI is no single cutter!

The complete DNA sequence is available at our webpage www.mobitec.com.

Quality Warranty

The vector features and restriction sites specified in this manual are verified by sequencing or checked for being functional. MoBiTec does not give any guarantee for sequence data of nonfunctional parts of the vectors.

Order Information, Shipping, and Storage

Order#	Product		Amount
PBS041	pTTB1		10 μg
PBS042	pTTB2		10 μg
Shipped at F	RT; plasmids are	yophilized from water an	d can be stored at 4 °C. Once
the DNA has	been dissolved	n sterile buffer or water,	store at -20 °C.
PBS043	Bacillus sub	tilis strain TEA	1 ml
PBS045	Bacillus sub	tilis strain WEA-C	1 ml
Shipped on o	dry ice; store at -	20 °C.	

The *Bacillus subtilis* Food Grade Expression System was developed by researchers of the Jiangnan University, China (Yang *et al.*, 2016). All strains and vectors belonging to this system are for academic research and development only.

Commercial enterprises: To obtain strains and plasmids of this system, a license needs to be negotiated. For license agreements please mail to info@mobitec.com.

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