Bacillus subtilis pBacTag Tagging Vectors



Page 2

Contents

	1 Introduction	3
	2 The pBacTag Tagging Vectors	3
	2.1 Mechanism of pBacTag Tagging Vectors	4
	3 B. subtilis and E. coli Host Strains	6
	4 Storage and Handling Instructions	6
	5 Chromosomal Integration of pBacTag	7
	5.1 Transformation protocol A	7
	5.2 Transformation protocol B	8
	5.3 Media and solutions	9
	7 Vector Maps	
	7.1 pBacTag Vectors with epitope tag	10
	7.2 pBacTag Vectors with localization tag	
	8 References	16
	9 Order Information, Shipping and Storage	17
•	0 Related Products	17
-	1 Contact and Support	17

1 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 and model organism, because of several reasons:

- Bacillus subtilis is non-pathogenic and awarded GRAS (generally regarded as safe) status from the US Food and Drug Administration.
- There is no significant bias in codon usage.
- It is capable of secreting high levels of functional 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 on *B. subtilis* is already available, greatly facilitating fundamental research experiments or the construction of improved protein production strains. The complete *B. subtilis* genome information is available in addition to many data on transcription, translation, protein folding, secretion mechanisms and gene manipulation results.

However, the functions of the about 4,100 *B. subtilis* genes identified, are still incompletely clarified. The pBacTag Tagging system has been developed to disburden further functional studies. On this, the pBacTag Tagging Vectors enable the directed functional analysis of genes by two different modes of action:

- Specific inactivation of genes of interest within the chromosome (followed by phenotypical analysis)
- Chromosomal expression of the gene of interest as translational fusion with an epitope or localization tag fused to the 3'-end (for selective protein purification, detection by commercially available antibodies or for localization studies).

The tagging or inactivation of the target gene is achieved by chromosomal integration of the pBacTag Tagging Vector into the *B. subtilis* chromosome by homologous recombination.

2 The pBacTag Tagging Vectors

The pBacTag Tagging Vectors enable the directed functional analysis of genes. Tagging or inactivation of target genes is achieved by chromosomal integration of a pBacTag Vector via homologous recombination. All pBacTag Tagging Vectors are derivatives of pMutin vectors (Vagner et al., 1998; Kaltwasser et al., 2002) with the following properties:

- pBacTag Tagging Vectors are able to replicate in *E. coli*, but unable to propagate in *B. subtilis*. The latter enables chromosomal integration with *B. subtilis* (and other bacterial species, in which pBR322 based plasmids are not able to replicate), by homologous recombination, using the erythromycin-resistance gene as selection marker. For propagation in *E. coli*, the β-lactamase gene can be used for selection purpose, causing resistance against ampicillin.
- The IPTG inducible Pspac promoter allows, after chromosomal integration, the controlled expression of genes that are located downstream of the target gene. This

is important, because most of *B. subtilis* genes are organized in multicistronic units, and downstream genes within the same operon may be separated from their native promoter by the integration event. The Pspac is therefore an indispensable tool to avoid polar effects from expression changes of downstream genes.

- For proper cloning, the vectors contain a multiple cloning site downstream of the Pspac with the following unique restriction sites: Kpnl, Eco47III, Clal and Eagl.
- To ensure efficient termination of transcription of the hybrid gene, the vectors contain the *trpA* terminator of the *E. coli* tryptophan operon downstream of the tag.
- The terminators t₁t₂t₀ (t₁ and t₂ of the *E. coli rnb* operon and the lambda terminator λt₀ downstream of the erythromycin resistance gene take care for proper RNA polymerase termination at this place and prevent any "read through" to genes downstream of the Pspac.
- Three of the tagging vectors pBacTag-DYKDDDK (also known as FLAG®), pBacTag-cMyc and pBacTag-HA allow the expression of epitope tagged fusion proteins. These proteins can be detected in immunoblotting experiments by using commercially available antibodies against the respective tag. The fusion proteins can also easily being purified using the tag in affinity chromatography. Since the tags are very short (FLAG®: 7 aa, cMyc: 10 aa, HA (hemagglutinin): 9 aa) protein function is usually not disturbed.
- Localization tags can be fused to the protein of interest, using pBacTag-GFP+, pBacTag-YFP and pBacTag-CFP. The fusion proteins containing a fluorescing tag can be analyzed for their cell compartmental localization. The GFP+ tag (pBacTag-GFP+) is an improved variant of the common GFP, which produces enhanced fluorescence.

2.1 Mechanism of pBacTag Tagging Vectors

The mechanism of pBacTag Tagging Vectors is illustrated with pBacTag-GFP+ as example. This vector can be used for creating a GFP+ fusion protein from any chromosomally located gene of interest, by fusing a gfp+ tag to the chosen gene. In this example the gene of interest is named *orf2*. It is part of an operon, including three genes in total (*orf1*, *orf2* and *orf3*). For getting the gfp+ tag fused to the *orf2*, the 3' part of the gene (*orf2*') has to be inserted into the multiple cloning site of the pBacTag-GFP+ vector. After transforming *B. subtilis* cells with this construct, chromosomal integration of the vector is achieved by selecting for cells with resistance against erythromycin. The integration is facilitated by homologous recombination of both *orf2*' copies (one copy being within the plasmid, the other one within the chromosomal DNA). The mechanism the pBacTag-GFP+ vector (with *orf2*') is integrated into the genome, is displayed in Figure 1.

After vector integration, the complete *orf2* is fused to the *gfp*+ gene and can be transcribed from the native promoter located upstream of *orf1*. Transcription of the tagged gene is terminated at the *trpA* terminator downstream of *gfp*+. *orf3* (formerly within the operon) is no more transcribed from its native promoter. Instead, its transcription is ensured by the IPTG inducible promoter Pspac.

FLAG® is a registered trademark of Sigma-Aldrich Co

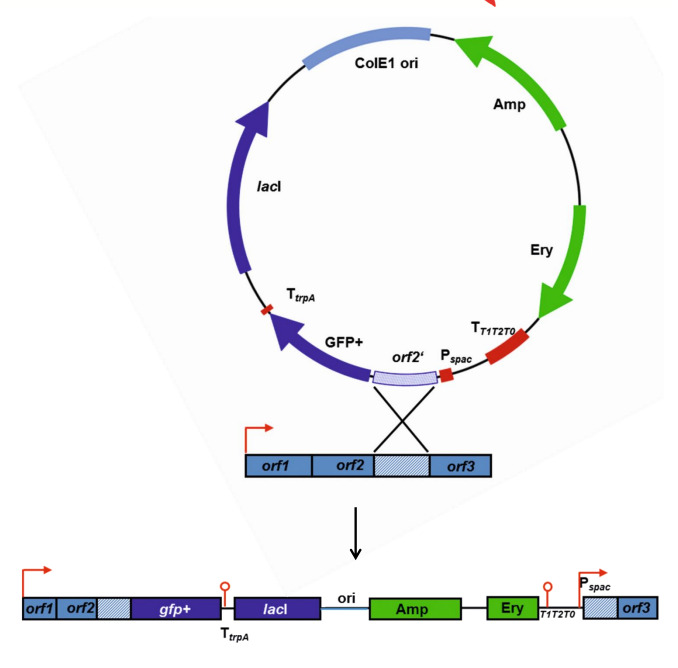


Fig.1: Chromosomal integration of pBacTag-GFP+:

The 3' end of orf2 (orf2') was ligated into the multiple cloning site of the pBacTag-GFP vector. The orf2 belongs to an operon containing a total of three genes (orf1, orf2 and orf3). The pBacTag-GFP+ vector (with orf2') is integrated into the chromosome of B. subtilis via homologous recombination of both orf2' sites, by a single crossing over event. Now, the complete orf2 is fused to the gfp+ gene and can be transcribed from the native promoter localized upstream of orf1. Transcription of the tagged gene is terminated at the trpA terminator downstream of gfp+. Now, orf3 is no more transcribed from the native promoter of the operon. Instead, its transcription can be induced from the Pspac promoter by adding IPTG. Broken arrows denote the promoters of the operon and Pspac. Promoters of lacl and the resistance genes are not depicted. The lollipop strands denote the trpA and the three lambda terminators (T1,T2,T0).

3 B. subtilis and E. coli Host Strains

The following bacterial strains are available from MoBiTec:

- B. subtilis 1012 wild type: leuA8 metB5 trpC2 hsdRM1
- B. subtilis 168 Marburg: trpC2 (Trp⁻)
- B.subtilis AS1: 1012 hrcA::neo (producing strain for enhancing solubility of intracellular protein Schulz and Schumann, 1996, and Phan et al., 2006)
- B. subtilis WB800N: nprE aprE epr bpr mpr::ble nprB::bsr Δvpr wprA::hyg cm::neo: NeoR (eight fold protease deficient strain for heterologous protein secretion)
- E. coli QuickCells F- (chemical competent): recA1 endA1 hsdR17 supE44 thi-1 gyrA96 relA1 Ø80lacZΔM15 Δ(lacZYA-argF)U169
- E. coli RichCells F- (chemical competent): recA1 endA1 hsdR17 supE44 thi-1 gyrA96 relA1 Ø 80lacZ Δ M15 Δ (lacZYA - argF)U169

Note: The neomycin marker of B. subtilis AS1 and WB800N is usually not required. For ordering details see "8. Related Products", p. 10.

4 Storage and Handling Instructions

Storage and handling of plasmids

Plasmids are supplied lyophilized. Upon receipt, add 50 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.

All plasmids of this system can be propagated in *E. coli*.

Storage and handling of B. subtilis and E. coli strains

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 medium (Bagyan et al., 1998). Under optimal conditions the doubling time of E. coli is 20 min, of B. subtilis 30 min.

2xYT

16 g tryptone medium:

10 g yeast extract

5 g sodium chloride (NaCl)

add distilled water to 1000 ml, autoclave at 121 °C for 15 min

Antibiotics: erythromycin (0.3 µg/ml) B. subtilis

> E. coli ampicillin (50 µg/ml)

Phone: +49 551 70722 0 Fax: +49 551 70722 22 MoBiTec GmbH, Germany E-Mail: info@mobitec.com www.mobitec.com

5 Chromosomal Integration of pBacTag

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

5.1 Transformation protocol A

The following transformation protocol is adopted from Klein et al., 1992. Please note that immediate usage may result in higher transformation rates.

Preparation of competent Bacillus subtilis cells

- Culture appropriate recipient cells in 5 ml HS medium at 37 °C overnight
- Inoculate 50 ml HS medium with 0.5 ml of the overnight culture
- Incubate under vigorous shaking at 37 °C
- Record the growth curve
- Take samples of 10 ml each when cells reach the stationary phase at 15 min intervals
- Add 1 ml of sterile glycerol (87%), 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 to identify the time point(s) yielding high level competent cells
- Discard the non- or low competent aliquots

Transformation of competent *Bacillus subtilis* cells and selection for chromosomal pBacTag integration

- 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 glass or 2 ml plastic reaction tubes, add 10 μ l of 0.1 M EGTA (CB-0732-10GAM), and incubate for 5 min at room temperature
- Add pBacTag Vector DNA (5-40 ng) 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 an plastic reaction tube
- Centrifuge, discard supernatant carefully and resuspend the cells into the final supernatant remaining on the pellet
- Plate on selective 2xYT or LB medium (0.3µg/ml erythromycin)
- Incubate at 37°C overnight
- Successful integration of the pBacTag vector as single copy might be controlled by Southern blot hybridization

5.2 Transformation protocol B

Electroporation of B. subtilis (modified from Zhang et al., 2011)

- Culture B. subtilis in 2xYT medium overnight
- Dilute 100-fold with 2xYT medium
- Grow culture to an OD600 of 0.2
- Then supplement culture with 1% DL-threonine, 2% glycine, 0.1% tryptophan and 0.03% Tween 80
- Grow while shaking for 1 h
- Cool on ice for 20 min
- Spin at 5000 x g for 10 min at 4 °C
- Wash twice with electroporation buffer
- Resuspend in electroporation buffer at 1/100 of the original culture volume
- Add 100 µl cell to an ice-cold 2 mm cuvette
- Add 2 µl pBacTag vector DNA (25 ng/ml)
- Shock by a single 12.5 kV/cm pulse (Gene Pulser; Bio-Rad), resistance 200 Ω , capacitance 25 μF
- Immediately add 1 ml 2xYT broth containing 0.5 M sorbitol and 0.38 M mannitol
- Incubate at 37 °C for 3 h
- Plate on selective 2xYT (0.3 µg/ml erythromycin)
- Incubate at 37°C overnight
- Successful integration of the pBacTag vector as single copy might be controlled by Southern blot hybridization

5.3 Media and solutions

S-base (Spizizen's salt): $2 g (NH_4)_2SO_4$

14 g K₂HPO₄ 6 g KH₂PO₄ 1 g sodium citrate

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

HS medium: 66.5 ml distilled water

10 ml 10x S-base

2.5 ml 20% (w/v) glucose 5 ml 0.1% (w/v) L-tryptophan

1 ml 2% (w/v) casein

5 ml 10% (w/v) yeast extract (Difco) 10 ml 8% (w/v) arginine, 0.4% histidine autoclave all components separately tryptophan solution: sterile filtration

LS medium 80 ml distilled water

10 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) casein

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

 $\begin{array}{c} 0.25 \text{ ml 1 M MgCl}_2 \\ 0.05 \text{ ml 1 M CaCl}_2 \end{array}$

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

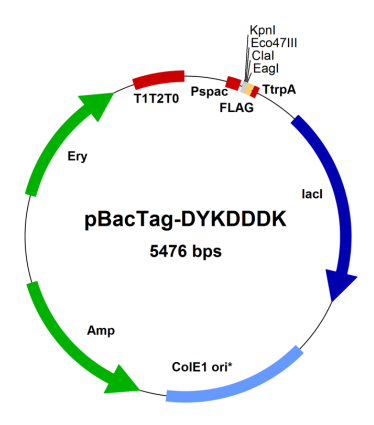
Electroporation buffer 0.5 M trehalose

0.5 M sorbitol 0.5 M mannitol 0.5 mM MgCl2 0.5 mM K2HPO4 0.5 mM KH2PO4 pH 7.4

filter-sterilize and store frozen

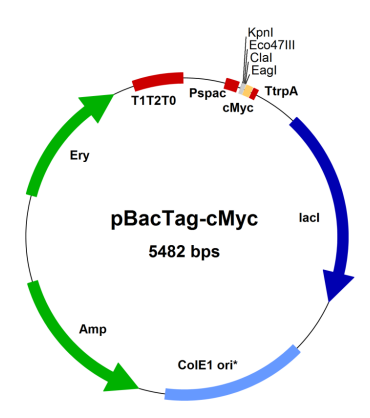
7 Vector Maps

7.1 pBacTag Vectors with epitope tag

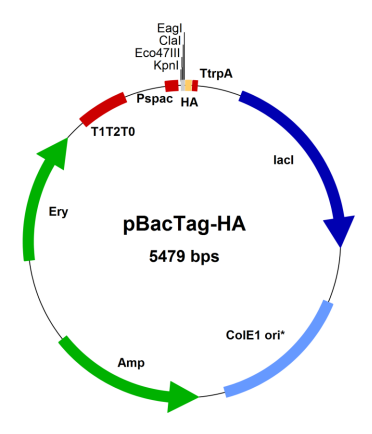


Туре	Start	End	Name	Description	
Promoter	228	298	Pspac	spac promoter	
Region	316	338	MCS	Multiple Cloning Site	
Tag	340	369	FLAG [®]	FLAG [®] tag	
Terminator	376	403	TtrpA	Termination sequence of <i>trpA</i> gene	
Gene	653	1735	lacl	lacl repressor gene	
Replication/ori	2049	2840	ColE1 ori*	Origin, belonging to the ColE1 incompatibility group	
Selectable Genetic Marker	3851	2991	Amp	Ampicillin resistance	
Selectable Genetic Marker	4335	5072	Ery	Erythromycin resistance	
Terminator	5193	5468	T1T2T0	Terminators t_1 , t_2 of \emph{rnb} operon of $\emph{E. coli}$ and λ terminator t_0	

FLAG® is a registered trademark of Sigma-Aldrich Co



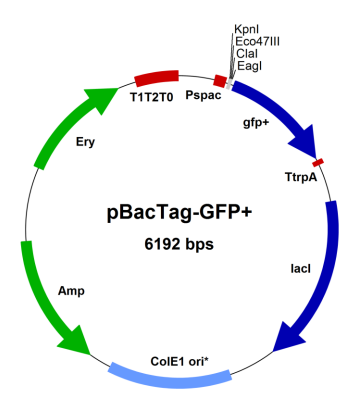
Туре	Start	End	Name	Description
Promoter	228	298	Pspac	spac promoter
Region	316	338	MCS	Multiple Cloning Site
Tag	340	375	сМус	cMyc tag
Terminator	382	409	TtrpA	Termination sequence of <i>trpA</i> gene
Gene	659	1741	lacl	lacl repressor gene
Replication/ori	2055	2846	ColE1 ori*	Origin belonging to the ColE1 incompatibility group
Selectable Genetic Marker	3857	2997	Amp	Ampicillin resistance
Selectable Genetic Marker	4341	5078	Ery	Erythromycin resistance
Terminator	5199	5474	T1T2T0	Terminators t_1 , t_2 of <i>rnb</i> operon of <i>E. coli</i> and λ terminator t_0



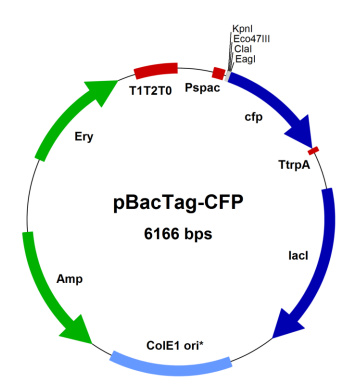
Туре	Start	End	Name	Description
Tag	6	38	НА	Hemmagglutinin tag
Terminator	45	72	TtrpA	Termination sequence of <i>trpA</i> gene
Gene	322	1404	lacl	lacl repressor gene
Replication/ori	1718	2509	ColE1 ori*	Origin belonging to the ColE1 incompatibility group
Selectable Genetic Marker	3520	2660	Amp	Ampicillin resistance
Selectable Genetic Marker	4004	4741	Ery	Erythromycin resistance
Terminator	4862	5137	T1T2T0	Terminators t_1 , t_2 of <i>rnb</i> operon of <i>Ecoli</i> and λ terminator t_0
Promoter	5373	5443	Pspac	spac promoter
Region	5461	5483	MCS	Multiple Cloning Site

Page 13

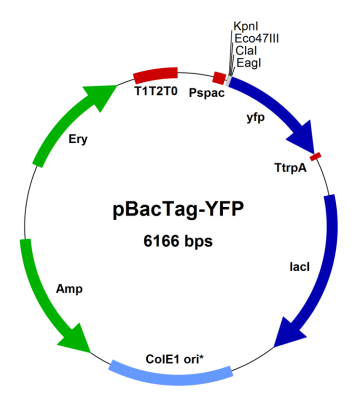
7.2 pBacTag Vectors with localization tag



Type Start		End	Name	Description
Promoter	228	298	Pspac	spac promoter
Region	316	338	MCS	Multiple Cloning Site
Gene	355	1074	gfp+	gfp+ gene (localization tag)
Terminator	1092	1119	TtrpA	Termination sequence of <i>trpA</i> gene
Gene	1369	2451	lacl	lacl repressor gene
Replication/ori	2765	3556	ColE1 ori*	Origin belonging to the ColE1 incompatibility group
Selectable Genetic Marker	4567	3707	Amp	Ampicillin resistance
Selectable Genetic Marker	5051	5788	Ery	Erythromycin resistance
Terminator	5909	6184	T1T2T0	Terminators t_1 , t_2 of <i>rnb</i> operon of <i>E. coli</i> and λ terminator t_0



Туре	Start	End	Name	Description
Promoter	228	298	Pspac	spac promoter
Region	316	338	MCS	Multiple Cloning Site
Gene	340	1059	cfp	cfp gene (localization tag)
Terminator	1066	1093	TtrpA	Termination sequence of <i>trpA</i> gene
Gene	1343	2425	lacl	lacl repressor gene
Replication/ori	2739	3530	ColE1 ori*	Origin belonging to the ColE1 incompatibility group
Selectable Genetic Marker	4541	3681	Amp	Ampicillin resistance
Selectable Genetic Marker	5025	5762	Ery	Erythromycin resistance
Terminator	5883	6158	T1T2T0	Terminators t_1 , t_2 of <i>rnb</i> operon of <i>E. coli</i> and λ terminator t_0



Туре	Start	End	Name	Description
Promoter	228	298	Pspac	spac promoter
Region	316	338	MCS	Multiple Cloning Site
Gene	340	1059	yfp	yfp gene (localization tag)
Terminator	1066	1093	TtrpA	Termination sequence of <i>trpA</i> gene
Gene	1343	2425	lacl	lacl repressor gene
Replication/ori	2739	3530	ColE1 ori*	Origin belonging to the ColE1 incompatibility group
Selectable Genetic Marker	4541	3681	Amp	Ampicillin resistance
Selectable Genetic Marker	5025	5762	Ery	Erythromycin resistance
Terminator	5883	6158	T1T2T0	Terminators t_1 , t_2 of <i>rnb</i> operon of <i>E. coli</i> and λ terminator t_0

8 References

- Anagnostopoulos C. and Spizizen, J. (1961). Requirements for transformation in *Bacillus subtilis*. J. Bacteriol. 81:741-746.
- Chubet R.G. and Brizzard B.L. (1996) Vectors for expression and secretion of FLAG epitope-tagged proteins in mammalian cells. BioTechniques 20:136-141.
- Kaltwasser M., Wiegert T. and Schumann W. (2002) Epitope- and Green Fluorescent Protein-Tagging Integration Vectors for *Bacillus subtilis*. Appl. Environ. Microbiol. 68:2624-2628.
- Klein, C., Kaletta, C., Schnell, N. & Entian, K.D. (1992) Analysis of genes involved in biosynthesis of the lantibiotic subtilin. Appl Environ Microbiol 58: 132–142.
- Margolin W. (2000) Green fluorescent protein as a reporter for macromolecular localization in bacterial cells. Methods 20:62-72.
- Sambrook, J. and Russel, D.W. (2001) Molecular Cloning: A laboratory manual.
- Vagner V., Dervyn E. and Ehrlich S.D. (1998) A vector for systematic gene inactivation in *Bacillus subtilis*. Microbiology 144:3097-3104.
- Zhang,G., Bao,P., Zhang,Y., Deng,A., Chen,N. and Wen,T. (2011) Enhancing electro-transformation competency of recalcitrant *Bacillus amyloliquefaciens* by combining cell-wall weakening and cell-membrane fluidity disturbing. Anal. Biochem., 409, 130–137.

9 Order Information, Shipping and Storage

Order#	Product	Quantity
PBT001	pBacTag-DYKDDDDK vector DNA	5 µg
PBT002	pBacTag-cMyc vector DNA	5 µg
PBT003	pBacTag-HA vector DNA	5 µg
PBT004	pBacTag-GFP+ vector DNA	5 µg
PBT005	pBacTag-CFP vector DNA	5 µg
PBT006	pBacTag-YFP vector DNA	5 µg
	om water, shipped at RT. Lyophilized plasmid DNA can be stored at 4	
the DNA has	been dissolved in sterile water or buffer we recommend storage at -20	°C.

10 Related Products

Order#	Product	Quantity
PBS020	Bacillus subtilis strain 1012wt	1 ml
PBS021	Bacillus subtilis strain 168 Marburg	1 ml
PBS022	Bacillus subtilis strain WB800N	1 ml
	(for secretion vectors)	
PBS026	Bacillus subtilis strain AS1	1 ml
PBS020	Bacillus subtilis strain 1012wt	1 ml
PBS021	Bacillus subtilis strain 168 Marburg	1 ml
Shipped on dry ice; s	tore at -80 °C	
CB-J902-100GAM	2xYT medium broth	100 g
CB-0339-25GAM	ampicillin sodium salt	25 g
CB-J859-100GAM	tryptone	100 g
CB-J851-100GAM	casamino acids	100 g
CB-0241-1KGAM	sodium chloride (NaCl)	1 kg
Shipped at RT		

11 Contact and Support

MoBiTec GmbH ◆ Lotzestrasse 22a ◆ D-37083 Goettingen ◆ Germany

Customer Service – General inquiries & orders **Technical Service** – Product information

phone: +49 (0)551 707 22 0 +49 (0)551 707 22 22 e-mail: order@mobitec.com

phone: +49 (0)551 707 22 70 +49 (0)551 707 22 77 e-mail: info@mobitec.com

MoBiTec in your area: Find your local distributor at www.mobitec.com