SynBac™

Ultra-Stable Baculovirus System for Expression Scale Up



User Manual

Version 2.0 (August 2024)

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SynBac™ Kit Plus Competent Cells Contents.

Plasmid acceptor vectors

pACEBac1, pACEBac2; approx. 5 μg DNA per vial (in buffer solution)

keep at 4°C for short-term storage and in a freezer at -20°C or lower for medium- and long-term storage (take care to avoid repeated freeze-thaw cycles, e.g. by aliquotting DNA prior to freezing)

Plasmid donor vectors

pIDC, pIDK, pIDS; approx. 5 μg DNA per vial (in buffer solution)

keep at 4°C for short-term storage and in a freezer at -20°C or lower for medium- and long-term storage (take care to avoid repeated freeze-thaw cycles, e.g. by aliquotting DNA prior to freezing)

• E. coli strains as chemical competent cells

a) *E.coli* strain harboring DH10SynBac bacmid (12 aliquots of 50μl each of chemical competent cells)

Please see section E.3.4 for a schematic description of the genome.

c) pirHC cells[†] (5 aliquots 100µl each chemical competent cells)

For propagation and amplification of donor vectors, donor multigene expression constructs or donor-donor fusions

Keep competent cells at -80°C do not store at -20!

[†] E. coli strains expressing the *pir* gene for propagation of donor vectors (any other strain with pir^{\dagger} background can be used as well). HC: high copy number propagation of plasmids with R6K γ origin.

Reagents to be supplied by user (see also section D. Protocols)

- Restriction enzymes
- Homing endonucleases PI-Scel and I-Ceul
- Insect cells, e.g. Sf9, Sf21 or High-Five™
- T4 DNA ligase
- Cre recombinase
- Standard *E. coli* strains for cloning (such as TOP10, DH5 α , HB101 etc.)
- Standard laboratory buffers, solutions, media and equipment for bacterial and insect cell culture, transformation etc.
- Commercially available transfection reagents, e.g. FuGENE® (Roche), jetPEI™ (Polyplus transfection), GeneJuice, etc.; alternatively, equipment for electroporation
- Antibiotics, chemicals (e.g. Kan, Tet, Gent, IPTG, X-Gal, etc.)

NOTES

B. Overview

This manual introduces a set of novel baculovirus transfer vectors that specifically enable multigene applications (Trowitzsch et al., 2010; Vijayachandran et al, 2011). These transfer vectors are accompanied by modified recipient baculovirus DNA that has been engineered towards improved protein production. This manual also presents a simple and rapid transposition method for integrating your gene(s) of interest into the baculoviral genome.

The role of protein interaction networks (the so-called **interactome**; reviewed e.g. in Figeys, 2008; Charbonnier et al., 2008) has become an intense focus of biological research efforts in the post-genomic era as most proteins have been shown to work together structurally and/or functionally in complexes for most basic cellular functions (transcription, translation, DNA replication and repair, cell cycle, protein quality control, etc.) but also dynamically in response to internal or environmental stimuli (inflammasome, signaling cascades, etc.).

Some of the identified multi-protein complexes are expressed at only low abundance in their native cells or such complexes exist only for brief periods (i.e. they are transitory in nature). This makes analysis of their structure-function difficult but this can be remedied by using recombinant technologies to facilitate large-scale heterologous protein production. Currently, recombinant expression methods require a disproportionate investment in both labor and materials prior to multi-protein expression, and, once expression has been established, provide little or no flexibility for rapidly altering the multiprotein components, which is a prerequisite for revising expression studies.

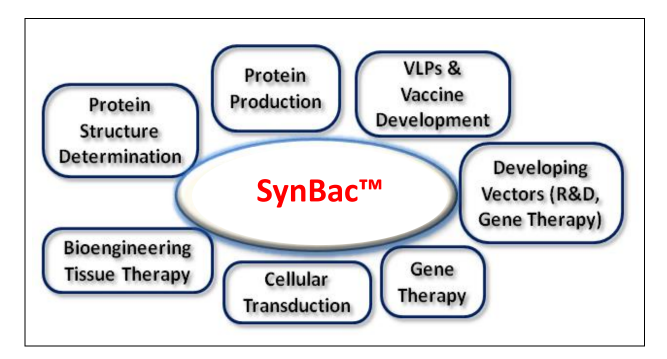


Figure 1. Applications of baculoviral expression systems (BEVS). Some of these are closely intertwined, e.g. transducing cells opens up the path for gene therapy and tissue therapy in mammalian cells.

The baculoviral expression system (BEVS) in its various shapes and forms has become a popular eukaryotic expression system, especially for multiple proteins. Yet, it has been adapted for other applications as well (see Figure 1), e.g. the generation of virus-like particles, bioengineering and tissues therapy research, transduction of mammalian cells, etc. The baculovirus expression system introduced here (see also Trowitzsch et al., 2010) boasts three major advances that are instrumental in fully exploiting the potential of this heterologous protein production system:

1. New transfer vectors (pACEBac1, pACEBac2, pIDC, pIDK, pIDS; see Figure 2) that contain a homing endonuclease-based multiplication module. These vectors greatly facilitate modular combination of heterologous genes (in their respective gene expression cassettes) with a minimum requirement for unique restriction sites (BstXI). Baculoviral promoters (currently p10 and polh very late promoters) can be exchanged in our vectors for other promoter sequences (early and late viral, mammalian) if desired. Likewise, terminator sequences (currently SV40, HSVtk) can be substituted as required.

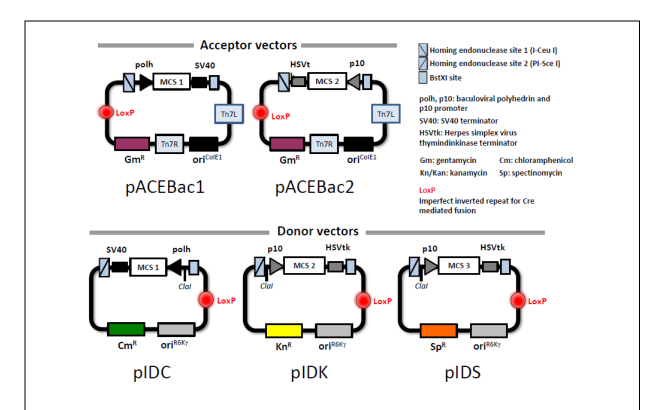


Figure 2. Schematic representation of the SynBac[™] acceptor and donor vectors. More detailed vectors maps and sequence information can be found in chapter C (p. 10-11) and in the appendix (chapter E, p. 32).

- 2. The baculovirus genome has been engineered to improve its protein production properties. Disruption of two baculoviral genes improves the integrity of cellular compartments during infection and protein production (see below). The quality of proteins produced by this system is significantly improved by curtailing virus-dependent proteolytic activity and reducing cell lysis.
- **3.** New protocol for rapid generation of multigene expression constructs via Cre-LoxP recombineering. The resulting multigene fusion is then incorporated into baculovirus DNA by accessing the viral genome via a specific site (see Figure 3). This protocol can be used to integrate multigene cassettes with coding sequences for multiprotein complex subunits into SynBac™, but also to integrate specific enzymes (kinases, acetylases etc.) for modifying the proteins under investigation.

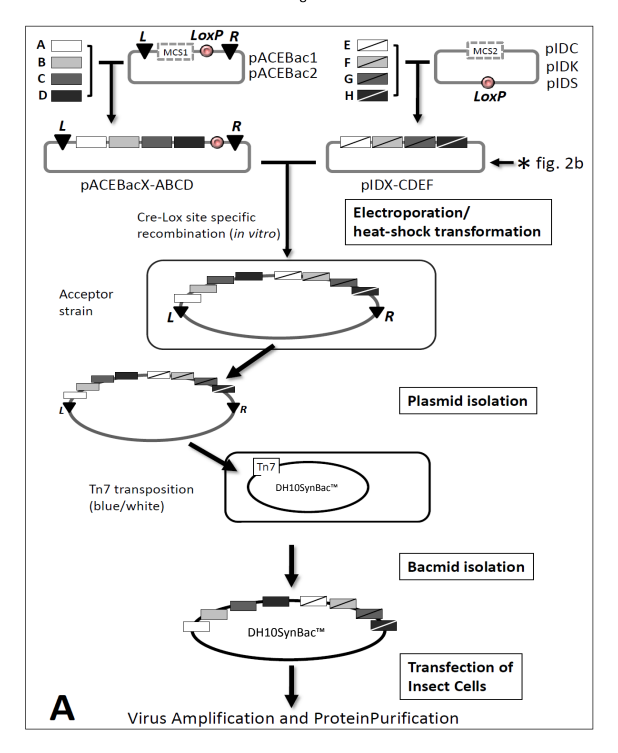


Figure 3. Schematic overview of the SynBac[™] system and its application. Genes of interest are assembled into multigene expression cassettes using the multiplication module present on the donor (pIDC, pIDK, pIDS) and acceptor vectors (pACEBac1, pACEBac2). Acceptor-donor fusions can then be generated by Cre-LoxP recombination. These multigene fusions contain one Acceptor and one to several Donor vectors, each with one or several genes of interest (here A-H). Desired Acceptor-Donor combinations are identified by transforming in E.coli and subsequent antibiotic selection. Acceptor vectors contain the DNA elements required for integration into the baculovirus via Tn7 transposition. DH10SynBac[™] E. coli cells contain the recipient baculovirus and also the transposase required for Tn7 transposition for integrating acceptor and acceptor-donor

derivatives that are transformed into these cells. Colonies containing bacmid carrying integrated multigene cassettes are identified by blue/white screening (Tn7 transposition disrupts expression of the $lacZ\alpha$ peptide) plus gentamycin resistance, and optionally through selective resistance markers introduced via Cre-catalyzed integration and hosted on the acceptor, donor or acceptor-donor derivatives). Bacmid DNA is prepared from selected clones and used to transfect insect cells for protein production. LoxP sites in the acceptor-donor fusions have been omitted for reasons of clarity.

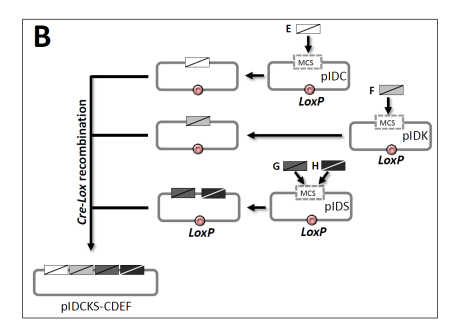


Figure 4. Generation of multigene donor constructs through Cre-Lox fusion. As indicated in Figure 3, donor multigene expression cassette constructs can also be generated by Cre-Lox recombination. Individual or multiple gene cassettes are cloned into the multiple cloning site via standard restriction-ligation cloning or, when introducing multiple gene cassettes, homing endonuclease /BstXI cloning. The gene cassettes harbored on different donor vectors are then merged into a single vector construct via Cre-Lox recombination. This construct will differ from the multigene constructs in Figure 3 with respect to selective markers. While the multigene construct in Figure 3 carries only one antibiotic resistance marker, the construct in this scheme will carry three, one from each donor vector. This will allow selection of multigene constructs with higher stringency by subjecting the constructs to a multiantibiotic selection regimen (refer to protocol 2). LoxP sites in the donor fusion have been omitted for reasons of clarity.

C. New Baculovirus Tools for Multigene Applications

C.1. Transfer vectors: the Acceptor-Donor recombineering system.

The **Acceptor vectors** pACEBac1 and pACEBac2 contain multiple cloning sites (see Appendix) flanked by either a polh or p10 promoters and SV40 or HSVtk polyA signal sequences, respectively. A multiplication module M – defined by the homing endonuclease site I-CeuI and a corresponding BsXI site (see Figure 5) – allows integration of multiple gene cassettes (ORFs and associated regulatory regions). The sequences used for Tn7 transposition (Tn7L and Tn7R) encompass the expression cassettes and a gentamycin resistance marker.

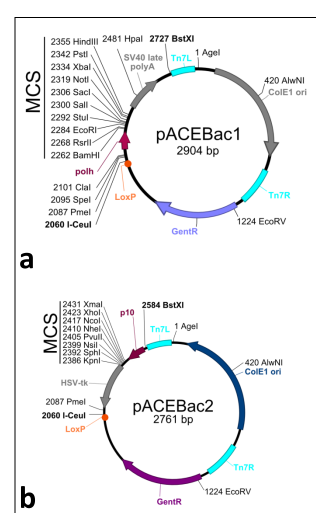


Figure 5. Circle map representation of Acceptor vectors

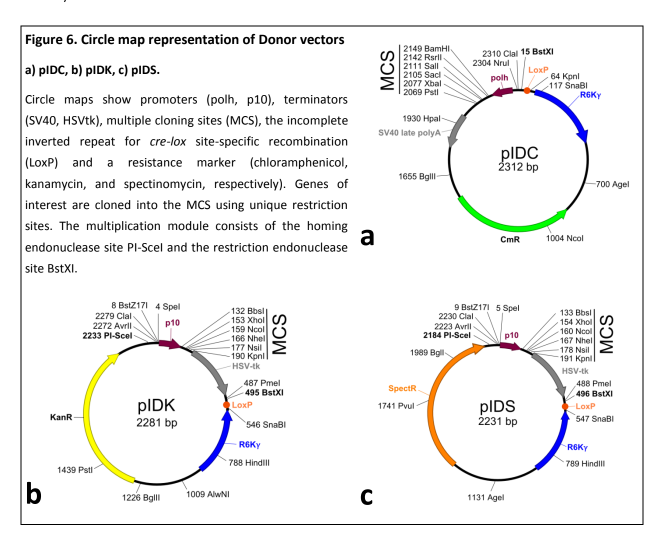
(a) pACEBac1 (2904 bp),

(b) pACEBac2 (2761 bp)

Both vectors carry a ColE1 origin of replication for maintenance of high plasmid copy number. Acceptor vectors also host polh (pACEBac1) and p10 (pACEBac2) promoters, SV40 and HSVtk terminators, multiple cloning sites (MCS), transposition elements (Tn7L, Tn7R) and a gentamycin resistance marker. Genes of interest are cloned into the unique restriction sites in the multiple cloning site.

The multiplication module flanks the MCS on either side and is defined by the restriction sites for the homing endonuclease *I-Ceu* and the restriction endonuclease BstXI, respectively. Genes or gene cassettes can also be recombined via Cre-Lox recombination making use of the incomplete inverted LoxP sites hosted on the vectors.

The **Donor vectors** pIDC, pIDS, pIDK are similar to the acceptor vectors with respect to their over-all design. The multiple cloning site is bracketed by a multiplication element (in this case, PI-Scel /BstXI) to enable concatenation of inserts between the different donor vectors. Vectors also contain a LoxP incomplete inverted repeat to create acceptor-donor or donor-donor fusions. The vectors contain "tell-tale" resistance markers (pID**C**: chloramphenicol, pID**K**: kanamycin, pID**S**: spectinomycin) and, importantly, a conditional R6Kγ origin of replication which makes its propagation dependent on the expression of the *pir* gene in the prokaryotic host (such as the pirLC and pirHC cells contained in the kit).



Currently, the SynBac[™] vectors do not contain DNA sequences coding for affinity tags that will facilitate purification or solubilization of the protein(s) of interest. Tags that are typically used are C- or N-terminal oligohistidine tags, with or without protease sites for tag removal. They can be introduced by designing the respective PCR primers used for amplification of the genes of interest. We recommend outfitting Donors or Acceptors of choice with any custom tag that is favored in individual

user laboratories prior to inserting recombinant genes of interest. This is best done by using a design that will, after tag insertion, still be compatible with the recombination-based principles of $SynBac^{TM}$ system usage.

C.2. Generating multigene expression cassettes

C.2.1. Multigene construction via homing endonuclease/BstXI multiplication

The acceptor and donor vectors are suited for generating multigene expression cassettes from individual gene expression cassettes (complete with regulatory regions such as promoter and terminator) via a multiplication module bracketing the multiple cloning site (MCS). All SynBac™ vectors contain a homing endonuclease (HE) site and a matching designed *BstXI* site that envelop the MCS. Homing endonucleases have long recognition sites (20-30 base pairs or more). Although not all equally stringent, homing endonuclease sites are most probably unique in the context of even large plasmids, or, in fact, entire genomes.

The logic of multiplication is illustrated below. The homing endonuclease site can be used to insert entire expression cassettes into a vector already containing one gene or several genes of interest as separate expression cassettes. The only prerequisite for assembling multigene expression cassettes is that the homing endonucleases and restriction enzymes used for multiplication (*I-CeuI/PI-SceI* and *BstXI*) are unique, which can be easily accomplished for instance by site directed mutagenesis prior to multigene cassette assembly. First, individual genes are cloned into the multiple cloning sites of the acceptor and donor vectors. The entire expression cassette, including promoter and terminator, is then excised by *I-CeuI* and *BstXI* (acceptors) or *PI-SceI* and *BstXI* (donors) digestion. The resulting fragment is placed into the multiplication module of another acceptor or donor vector containing single or multiple gene cassettes. The restriction sites involved are eliminated in the process and multiplication can be repeated iteratively using the module present in the inserted cassette. Moreover, promoter and terminator sequences can be easily modified if desired using appropriate restriction sites in our vectors

Please note that multiplication cannot be accomplished from donors to acceptors and vice versa since the overhangs generated by endonuclease digestion are incompatible.

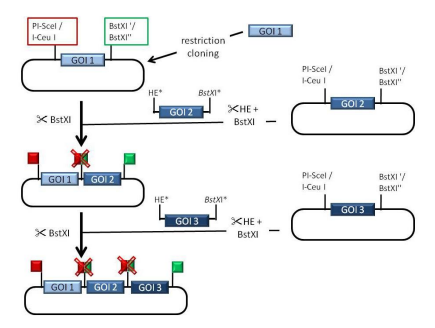


Figure 7. Assembling individual gene cassettes into multigene expression cassettes. The logic of multiplication is shown schematically. The expression cassette containing the gene of choice (denoted as GOI2 in this case) is excised by digestion with the homing endonuclease (red box) and BstXI (green box). For acceptors vectors, I-CeuI is the homing endonuclease of choice, and for donor vectors PI-SceI. The plasmid vector harboring the GOI1-cassette only needs to be linearized with BstXI. The homing endonucleases produce cohesive ends that are compatible with the ends generated by the BstXI digest. Upon insertion of GOI2 into the target vector, a homing endonuclease/BstXI hybrid restriction site is created that can then cannot be cut by either enzyme (crossed-out red/green box) while the 3'-BstXI site is regenerated. The same procedure can be repeated over and over as exemplified by the integration of GOI3. This cycling logic can be used to generate multigene assemblies. Note that the promoters and terminators are not explicitly shown for reasons of clarity.

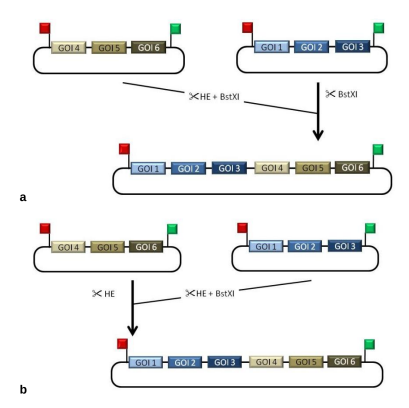


Figure 8. Combining multigene expression cassettes. Different multigene expression cassettes can be combined into one expression construct following the same logic that applies to the generation of multigene expression cassettes from individual gene cassettes (Figure 4). The 5' homing endonuclease recognition site (filled red box) will be preserved if GOI1 has been introduced by conventional restriction cloning into the MCS. Promoters and terminators are not explicitly shown for reasons of clarity but flank the GOIs in every individual gene expression cassette.

C.2.2. Multigene construction using Cre-Lox recombination

Cre recombinase is a member of the integrase family (Type I topoisomerase from bacteriophage P1). It recombines a 34 bp LoxP site in the absence of accessory protein or auxiliary DNA sequence. The LoxP site is comprised of two 13 bp recombinase-binding elements arranged as inverted repeats which flank an 8 bp central region where cleavage and the ligation reaction occur.

The site-specific recombination mediated by Cre recombinase involves the formation of a Holliday junction (HJ). The recombination events catalyzed by Cre recombinase depend on the location and relative orientation of the LoxP sites. Two DNA molecules, for example an acceptor and a donor plasmid, containing single LoxP sites will be fused. Furthermore, the Cre recombination is an equilibrium reaction with 20-30% efficiency in recombination. This provides useful options for multigene combinations for multi-protein complex expression.

13bp	8bp	13bp		
5'ATAACTTCGTATA	GCATACAT	TATACGAAGTTAT3'		
3'TATTGAAGCATAT	CGTATGTA	ATATGCTTCAATA5'		
inverted repeat	Spacer	inverted repeat		
Figure 9.: LoxP imperfect inverted repeat				

In a reaction where several DNA molecules such as donors and acceptors are incubated with Cre recombinase, the fusion/excision activity of the enzyme will result in an equilibrium state where single vectors (educt vectors) and all possible fusions coexist. Donor vectors can be used with acceptors and/or donors, and vice versa. Higher order fusions are also generated where more than two vectors are fused. This is shown schematically in Illustration 6.

The fact that Donors contain a conditional origin of replication that depends on a pir^{+} (pir positive) background now allows for selecting out from this reaction mix all desired Acceptor-Donor(s) combinations. For this, the reaction mix is used to transform pir negative strains (TOP10, DH5 α , HB101 or other common laboratory cloning strains). Then, Donor vectors will act as suicide vectors when plated out on agar containing the antibiotic corresponding to the Donor encoded resistance marker, unless fused with an Acceptor. By using agar with the appropriate combinations of antibiotics, all desired Acceptor-Donor fusions can be selected for.

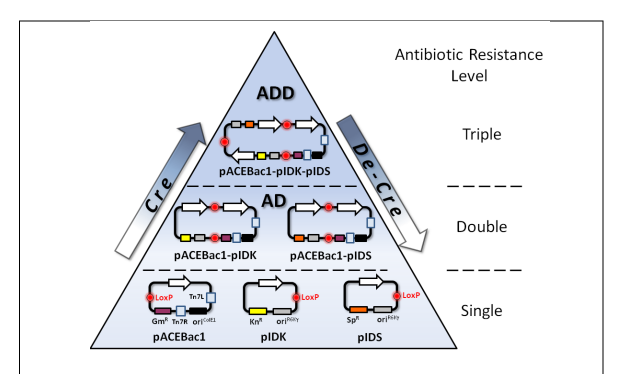


Figure 10.: Cre and De-Cre reaction pyramid

Cre-mediated assembly and disassembly of pACEBac1, pIDK, and pIDS vectors are shown in a schematic representation (left). LoxP sites are shown as red circles, resistance markers and origins are labelled. White arrows stand for the entire expression cassette (including promoter, terminator and gene integration/multiplication elements) in the SynBac™ vectors. Not all possible fusion products are shown for reasons of clarity. Levels of multiresistance are indicated (right column).

C.2.3. Combining HE/BstXI cycling and Cre-Lox recombination

Of course, both methods can also be combined to generate multiple gene-expression cassette constructs. To this end, you can introduce multiple gene cassettes with the homing endonuclease/BstXI protocol into different Acceptor/Donor vectors and then fuse these using the Cre-Lox modules.

C.3. Baculovirus engineered for improved protein production.

During heterologous protein production using other commercially available baculovirus expression system, viral-dependent proteolytic breakdown consistent with the action of a cysteine protease can be observed. The SynBac™ baculovirus genome (schematically shown in Figure 11) was modified to yield improved protein production properties. Two baculoviral genes, *v-cath* and *chiA*, have been disrupted which leads to improved maintenance of cellular compartments during infection and protein production. The *v-cath* gene (Slack et al., 1995) encodes for a viral cathepsin-type cysteine protease, V-CATH, which is activated upon cell death by a process that depends on a juxtaposed gene on the viral DNA, chiA, which encodes for a chitinase (Hawtin et al., 1995; Hom and Volkman, 2000). Both are involved in the liquefaction of the host insect cells (Slack et al., 1995; Hawtin et al., 1997).

Disruption of both genes served to a) eliminate V-CATH activity and b) to enable chitin-affinity chromatography for purification without interference from the *chiA* gene product. The quality of proteins produced by the SynBacTM baculovirus is significantly improved through a reduction of virus-dependent proteolytic activity and reduced cell lysis. The disrupted viral DNA sequence was replaced with a LoxP sequence for *cre-lox* site-specific recombination. Note that this LoxP site is not used for introduction of the target constructs into SynBacTM. Instead, the genes of interest are transferred to the bacmid via transposition into the mini Tn7 attachment site. Successful integration results in disruption of the lacZ α subunit-coding sequence. As a consequence, clones carrying inserted DNA will appear white.

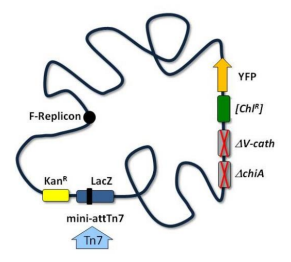


Figure 11.: DH10SynBac™ baculoviral DNA. The modified viral genome is shown in a schematic representation. The Tn7 attachment site is located within a LacZα gene; insertion of Tn7 elements from pACEBac derivatives therefore produces a white phenotype when plated on agar containing BluoGal (or Xgal) and IPTG. The viral genes *v-cath* and *chiA* are disrupted by replacement with an antibiotic marker and an eYFP expression cassette. Other regions of DNA are minimized, with a number of non-essential genes deleted

C.4. Introducing additional control elements

Protein expression requires certain genetic elements at the DNA and RNA level for it to work efficiently, as exemplified by the Shine-Dalgarno or Kozak sequence in bacterial and vertebrate mRNAs, respectively. Without this ribosome-binding motif, efficient translation of the corresponding mRNAs is compromised. Yet, many genes/mRNAs lack such a consensus sequence motif which is not an absolute must for successful translation.

While the polyhedrin promoter already is one of the strongest promoters known, protein production may in some cases be enhanced by substituting the intrinsic leader with alternative leader sequences in the gene expression cassettes. While there is no guarantee that this will work equally well for all proteins under investigation, several reports suggest the inclusion of such motifs into gene design, e.g. the L21 leader from a lobster tropomyosin cDNA (Sano et al., 2002).

In most cases, though, protein production should work just fine with the promoters provided in the transfer vectors.

- 20 -

D. Protocols

D.0 Introductory remarks

Please note that the bacteria in agar stabs have not been made competent for transformation. If

you wish to use them to transform your constructs, you will have to prepare competent cells. This

applies specifically to the pirHC and pirLC strains used to maintain and amplify donor constructs. You

may follow your preferred protocol for preparing chemically or electrocompetent cells, e.g. Inoue et

al. (1990) or variations of this protocol, or standard protocols as described in Current Protocols in

Molecular Biology or Sambrook and Russell: Molecular Cloning (3rd edition, 2001, or older versions).

D.1 Cloning into pACEBac or pIDx transfer vectors

Reagents:

Restriction endonucleases

DNA ligase

E. coli competent cells

Antibiotics: Chloramphenicol, Gentamycin, Kanamycin, Spectinomycin

The genes of choice are cloned using standard cloning procedures into the multiple cloning sites MCS

(see Supplementary Information) of pACEBac1/2 and pIDC, pIDK, pIDS. Ligation reactions for pACEBac

derivatives are transformed into standard *E. coli* cells for cloning (such as TOP10, DH5 α , HB101) and

plated on agar containing gentamycin (7 µg/ml). Ligation reactions for pIDx derivatives are

transformed into E. coli cells expressing the pir gene (pirHC and pirLC from this kit - in this case you

will need to make the cells electro- or chemically competent first; other strains, e.g. BW23473,

BW23474) and plated on agar containing chloramphenicol (pIDC; 25 μg/ml), kanamycin (pIDK; 50

μg/ml) or spectinomycin (pIDS; 50 μg/ml). Correct clones are selected based on specific restriction

digestion patterns and DNA sequencing of the inserts.

D.2 Multiplication by using the HE and BstXI sites

SynBac™ donor vectors contain a recognition site for the homing endonuclease PI-SceI (fig. 3). Upon

cleavage, this HE site yields a 3' overhang with the sequence -GTGC. Acceptor vectors contain the

homing endonuclease site I-CeuI (see fig. 2), which upon cleavage will result in a 3' overhang of -CTAA. On acceptors and donors, the respective HE site precedes the MCS (see fig. 2 and 3). The 3' end of the MIE contains a specifically designed BstXI site, which upon cleavage will generate a matching overhang. The basis of this is the specificity of cleavage by BstXI. The recognition sequence of BstXI is defined as CCANNNNN'NTGG (the apostrophe marks the position of the phosphodiester link cleavage). The residues denoted as N can be chosen freely. Donor vectors thus contain a BstXI recognition site with the sequence CCATGTGC'CTGG, and Acceptor vectors contain CCATCTAA'TTGG. The overhangs generated by BstXI cleavage in each case will match the overhangs generated by HE cleavage. Note that Acceptors and Donors have different HE sites.

The recognition sites are not symmetric. Therefore, ligation of a HE/BstXI digested fragment into a HE site of an SynBac™ vector will be (1) directional and (2) result in a hybrid DNA sequence where a HE half site is combined with a BstXI half site (see Figure 5). This site will be cut by neither the HE nor BstXI. Therefore, in a construct that has been digested with a HE, insertion by ligation of HE/BstXI digested DNA fragment containing an expression cassette with one or several genes will result in a construct which contains all heterologous genes of interest, enveloped by an intact HE site in front, and a BstXI site at the end. Therefore, the process of integrating entire expression cassettes by means of HE/BstXI digestion and ligation into a HE site can be repeated iteratively.

D2.1 Protocol 1. Multiplication by using homing endonuclease/BstXI.

Reagents required:

Homing endonucleases PI-Scel, I-Ceul
10x Buffers for homing endonucleases
Restriction enzyme BstXI (and 10x Buffer)
T4 DNA ligase (and 10x Buffer)
E. coli competent cells
Antibiotics

Step 1: Insert preparation

Restriction reactions are carried out in 40 μ l reaction volumes, using homing endonucleases PI-Scel (Donors) or I-Ceul (Acceptors) as recommended by the supplier.

Acceptor or donor plasmid (\geq 0.5 µg) in ddH₂O 32 µl 10x restriction enzyme buffer 4 µl 10 mM BSA 2 µl PI-Scel (Donors) or I-Ceul (acceptors) 2 µl ca. 10 U

Reactions are then purified using a PCR extraction kit or by acidic ethanol precipitation, and subsequently digested with BstXI according to the supplier's recommendations.

HE digested DNA in ddH $_2$ O 32 μ l 10x restriction enzyme buffer 4 μ l 2 μ l BstXI 2 μ l ca. 20 U

Gel extraction of insert(s):

Processed insert is then purified by agarose gel extraction using commercial kits (Qiagen, Macherey Nagel etc.). Elution of the extracted DNA in the minimal volume defined by the manufacturer is recommended.

Step 2: Vector preparation

Restriction reactions are carried out in 40 μ l reaction volumes, using homing endonucleases PI-Scel (Donors) or I-Ceul (Acceptors) as recommended by the supplier.

Acceptor or donor plasmid ($\geq 0.5 \mu g$) in ddH2O	33 µl	
10x Restriction enzyme buffer	4 μΙ	
10 mM BSA	2 μΙ	
BstXI	2 μΙ	ca. 20 U

Reactions are then purified by PCR extraction kit or acidic ethanol precipitation, and next treated with intestinal alkaline phosphatase according to the supplier's recommendations. Dephosphorylation is performed to minimize vector re-annealing and to increase integration of the insert.

HE digested DNA in ddH2O	17 µl	
10x Alkaline phosphatase buffer	2 μΙ	
Alkaline phosphatase	1 μΙ	ca. 10 U

Gel extraction of vector:

Processed vector is then purified by agarose gel extraction using commercial kits (Qiagen, Macherey Nagel etc). Elution of the extracted DNA in the minimal volume defined by the manufacturer is recommended.

Step 3: Ligation

Ligation reactions are carried out in 20 μ l reaction volumes:

HE/Phosphatase treated vector (gel extracted)	4 μΙ	
HE/BstXI treated insert (gel extracted)	14 μΙ	
10x T4 DNA Ligase buffer	2 μΙ	
T4 DNA Ligase	0.5 μΙ	150 - 200 c.e. U or 2-4 Weiss U

Ligation reactions are performed at 25°C for 1h or at 16°C overnight.

Step 4: Transformation

Mixtures are next transformed into competent cells following standard transformation procedures.

Ligation reactions for pACEBac1 and pACEBac2 derivatives are transformed into standard *E. coli* cells for cloning (such as TOP10, DH5 α , HB101) and, after recovery, are plated on agar containing gentamycin (7 μ g/ml).

Reactions for Donor derivatives are transformed into *E. coli* cells expressing the *pir* gene (such as BW23473, BW23474, or PIR1 and PIR2 from Invitrogen and, of course, pirLC and pirHC in this kit) and plated on agar containing chloramphenicol (25 μ g/ml, pIDC), kanamycin (50 μ g/ml, pIDK), or spectinomycin (50 μ g/ml, pIDS).

Step 5: Plasmid analysis

Plasmids are cultured and correct clones selected based on specific restriction digestion and DNA sequencing of the inserts.

D.3 Cre-LoxP reaction of Acceptors and Donors

D 3.1. Protocol 2: Cre-LoxP fusion of Acceptors and Donors

This protocol is designed for generating multigene fusions from Donors and Acceptors by Cre-LoxP reaction.

Reagents:

Cre recombinase (from NEB or self-made)

Standard E. coli competent cells (pir strain, such as Top10)

Antibiotics

96 well microtiter plates

12 well tissue-culture plates (or petri dishes) w. agar/antibiotics

LB medium

- 1. For a 20 μ l Cre reaction, mix 1-2 μ g of each educt in approximately equal amounts. Add ddH₂O to adjust the total volume to 16-17 μ l, then add 2 μ l 10x Cre buffer and 1-2 μ l Cre recombinase (1-2 U).
- 2. Incubate Cre reaction at 37°C (or 30°C) for 1 hour.
- 3. Optional: load 2-5 µl of Cre reaction on an analytical agarose gel for examination.

Heat inactivation at 70°C for 10 minutes before gel loading is strongly recommended.

4. For chemical transformation, mix 10-15 μ l Cre reaction with 200 μ l chemically competent cells. Incubate the mixture on ice for 15-30 minutes. Then perform heat shock at 42°C for 45-60 s.

Up to 20 μ l Cre reaction (0.1 volumes of the chemically competent cell suspension) can be directly transformed into 200 μ l chemical competent cells.

For electrotransformation, up to 2 μ l Cre reaction can be directly mixed with 100 μ l electrocompetent cells, and transformed by using an electroporator (e.g. BIORAD E. coli Pulser) at 1.8-2.0 kV.

Larger volumes of Cre reaction must be desalted by ethanol precipitation or via PCR purification columns before electrotransformation. The desalted Cre reaction mix should not exceed 0.1 volumes of the electrocompetent cell suspension.

The cell/DNA mixture can be immediately used for electrotransformation without prolonged incubation on ice.

5. Add up to 400 μ l of LB (or SOC) medium per 100 μ l of cell/DNA suspension immediately after the transformation (heat shock or electroporation).

6. Incubate the suspension in a 37°C shaking incubator overnight or for at least 4 hours (recovery period).

To recover multifusion plasmid containing more than 2 resistance markers, it is strongly recommended to incubate the suspension at 37°C overnight.

- 7. Plate out the recovered cell suspension on agar containing the desired combination of antibiotics. Incubate at 37°C overnight.
- 8. Clones from colonies present after overnight incubation can be verified by restriction digestion at this stage (refer to steps 12-16).

This quality control step should be carried out especially in the case that only one specific multifusion plasmid is desired.

For further selection by single antibiotic challenges on a 96 well microtiter plate, continue to step 9.

Several to many different multifusion plasmid combinations can be processed and selected in parallel on one 96 well microtiter plate.

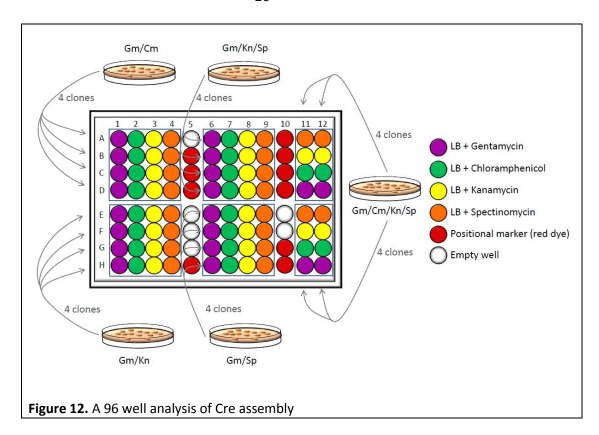
- 9. For 96 well antibiotic tests, inoculate four colonies from each agar plate with different antibiotic combinations into approx. 500 μ l LB medium without antibiotics. Incubate the cell cultures in a 37°C shaking incubator for 1-2 hours.
- 10. While incubating the colonies, fill a 96 well microtiter plate with 150 μ l antibiotic-containing LB medium (following Illustration 7). It is recommended to add colored dye (positional marker) in the wells indicated.

A typical arrangement of the solutions, which is used for parallel selections of multifusion plasmids, is shown in Illustration 10. The concept behind the 96 well plate experiment is that every cell suspension from single colonies needs to be challenged by all four single antibiotics for unambiguous interpretation.

11. Add 1 μ l aliquots of pre-incubated cell culture (Step 9) to the corresponding wells. Then incubate the inoculated 96 well microtiter plate in a 37°C shaking incubator overnight at 180-200 rpm.

Recommended: use parafilm to wrap the plate to avoid drying out.

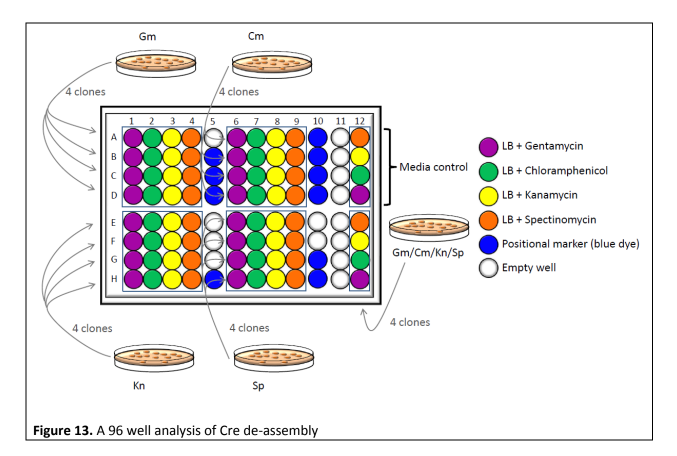
The remainder of the pre-incubated cell cultures can be kept at 4°C for further inoculations if necessary.



- 12. Select transformants containing desired multifusion plasmids based on antibiotic resistance, according to the combination of dense (positive) and clear (no growth) cell microcultures from each colony. Inoculate 10-20 μ l cell culture into 10 ml LB media with corresponding antibiotics. Incubate in a 37°C shaking incubator overnight.
- 13. Centrifuge the overnight cell cultures at 4000g for 5-10 minutes. Purify plasmid from the resulting cell pellets with common plasmid miniprep kits, according to manufacturers' recommendation.
- 14. Determine the concentrations of purified plasmid solutions by using UV absorption spectroscopy (e.g. by using a NanoDrop[™] 1000 machine).
- 15. Digest $0.5-1~\mu g$ of the purified plasmid solution in a 20 μl restriction digestion with appropriate endonuclease(s). Incubate under recommended reaction condition for approx. 2 hours.
- 16. Use 5-10 μl of the digestion for analytical agarose (0.8-1.2%) gel electrophoresis. Verify plasmid integrity by comparing the experimental restriction pattern to a restriction pattern predicted *in silico* (e.g. by using program VectorNTI, SnapGene or similar programs).

D 3.2. Protocol 3. Deconstruction of fusion vectors by Cre

The following protocol is suitable for releasing any single educt from multifusion constructs (deconstruction). This is achieved by Cre-LoxP reaction, transformation and plating on agar with appropriately reduced antibiotic resistance level (cf. Figure 13). In the liberated educt entity, encoding genes can be modified and diversified. Then, the diversified construct is resupplied by Cre-LoxP reaction (C3.1).



Reagents:

Cre recombinase (and 10x Buffer)

E. coli competent cells

 $(pir^{+}$ strains, pir^{-} strains can be used only when partially deconstructed Acceptor-Donor fusions are desired).

Antibiotics

- 1. Incubate approx. 1 μ g multifusion plasmid with 2 μ l 10x Cre buffer, 1-2 μ l Cre recombinase, add ddH₂O to adjust the total reaction volume to 20 μ l.
- 2. Incubate this Cre deconstruction reaction mixture at 30°C for 1 to 4 hour(s).

3. Optional: load 2-5 µl of the reaction on an analytical agarose gel for examination.

Heat inactivation at 70°C for 10 minutes before gel loading is strongly recommended.

4. For chemical transformation, mix 10-15 μ l De-Cre reaction with 200 μ l chemically competent cells. Incubate the mixture on ice for 15-30 minutes. Then perform heat shock at 42°C for 45-60 s.

Up to 20 μ l De-Cre reaction (0.1 volumes of the chemical competent cell suspension) can be directly transformed into 200 μ l chemically competent cells.

For electrotransformation, up to 2 μ l De-Cre reaction can be directly mixed with 100 μ l electrocompetent cells, and transformed by using an electroporator (e.g. BIORAD *E. coli* Pulser) at 1.8-2.0 kV.

Larger volume of De-Cre reaction must be desalted by ethanol precipitation or PCR purification column prior to electrotransformation. The desalted De-Cre reaction mix should not exceed 0.1 volumes of the electrocompetent cell suspension.

The cell/DNA mixture can be immediately used for electrotransformation without prior incubation on ice.

- 5. Add up to 400 μ l of LB media (or SOC media) per 100 μ l of cell/DNA suspension immediately after the transformation (heat shock or electroporation).
- 6. Incubate the suspension in a 37°C shaking incubator (recovery).

For recovery of partially deconstructed double/triple fusions, incubate the suspension in a 37°C shaking incubator for 1 to 2 hours.

For recovery of individual educts, incubate the suspension in a 37°C shaking incubator overnight or for at least 4 hours.

- 7. Plate out the recovered cell suspension on agar containing the desired (combination of) antibiotic(s). Incubate at 37°C overnight.
- 8. Colonies after overnight incubation can be verified directly by restriction digestion at this stage (refer to steps 12-16).

This is especially recommended in cases where only a single educt or partially deconstructed multifusion plasmid is desired.

For further selection by single antibiotic challenge on a 96 well microtiter plate, continue with step 9.

Several different single educts/partially deconstructed multifusion plasmids can be processed and selected in parallel on one 96 well microtiter plate.

9. For 96 well analysis, inoculate four colonies each from agar plates containing a defined set of antibiotics into approx. 500 μ l LB medium without antibiotics. Incubate the cell cultures in a 37°C shaking incubator for 1-2 hours.

- 10. While incubating the colonies, fill a 96 well microtiter plate with 150 μ l antibiotic-containing LB medium or dye (positional marker) in the corresponding wells.
 - Refer to Figures 12 and 13 for the arrangement of the solutions in the wells, which are used for parallel selection of single educts or partially deconstructed multifusion plasmids. The concept is that every cell suspension from a single colony needs to be challenged by all four antibiotics separately for unambiguous interpretation.
- 11. Add 1 μ l aliquots from the pre-incubated cell cultures (Step 9) into the corresponding wells. Incubate the 96 well microtiter plate in a 37°C shaking incubator overnight at 180-200 rpm.
 - Recommended: use parafilm to wrap the plate to prevent desiccation.
 - The remainder of the pre-incubated cell cultures can be kept at 4°C in a refrigerator for further inoculations if necessary.
- 12. Select transformants containing desired single educts or partially deconstructed multifusion plasmids according to the combination of dense (growth) and clear (no growth) cell cultures from each colony. Inoculate 10-20 μ l from the cell cultures into 10 ml LB media with corresponding antibiotic(s). Incubate in a 37°C shaking incubator overnight.
- 13. The next day, centrifuge the overnight cell cultures at 4000g for 5-10 minutes. Purify plasmid from cell pellets with common plasmid miniprep kits, according to manufacturers' protocols.
- 14. Determine the concentrations of purified plasmid solutions by using UV absorption spectroscopy (e.g. NanoDrop[™] 1000).
- 15. Digest 0.5-1 μ g of the purified plasmid solution in a 20 μ l restriction digestion (with 5-10 units endonuclease). Incubate under recommended reaction condition for approx. 2 hours.
- 16. Use 5-10 μl of the digestion for analytical agarose gel (0.8-1.2%) electrophoresis. Verify plasmid integrity by comparing the *de facto* restriction pattern to the *in silico* predicted restriction pattern (e.g. by using VectorNTI, Invitrogen, or any other similar program).
- 17. Optional: Occasionally, a deconstruction reaction is not complete but yields partially deconstructed fusions which still retain entities to be eliminated. In this case, we recommend to pick these partially deconstructed fusions containing and perform a second round of Cre deconstruction reaction (repeat steps 1-8) by using this construct as starting material.

D.4. Transposition protocol for pACEBac derivatives (electrocompetent cells or chemical competent).

Reagents:

Electrocompetent or chemical competent DH10SynBac™ cells

Kanamycin

Tetracycline

Gentamycin

BluoGal

IPTG

Electroporation: Approximately 5-10 ng of the sequenced pACEBac derivative or of an acceptor-donor fusion is incubated on ice (15 min) with 50-100 μ l electro-competent DH10SynBacTM cells. Optimum electroporation settings need to be determined for the equipment used.

Transformation of chemical competent cells: Approximately 100 ng ($\underline{maximally\ 10\ \mu l\ volume}$) of the sequenced pACEBac derivative or of an acceptor-donor fusion in is incubated on ice (30 min) with 50-100 μ l chemical-competent DH10SynBacTM cells. Heat shock is carried out at 42 °C for 15 seconds and the cells are placed again quickly on ice.

Following electroporation or transformation using chemical competent cells, 500 microliters prewarmed SOC media is added to cells. Cells are incubated at 37 °C for 4 hours and plated on agar plates containing kanamycin (50 μ g/ml), gentamycin (7 μ g/ml), tetracycline (10 μ g/ml), BluoGal (100 μ g/ml) and IPTG (40 μ g/ml). We recommend plating a dilution series in SOC or 2XYT media, plating 300 microliters transformed cells on one plate, and plating 30 microliters transformed cells on a second plate.

White colonies are selected after incubation at 37 °C (24 hours). Deeper blue and white color colonies become more visible after leaving the plates for an additional day on the bench at room temperature. Proceed to bacmid preparation for insect cell infection (D.5.).

D.5. Bacmid preparation and infection of insect cells.

Preparation of bacmid DNA, infection of insect cells and protein expression is carried out according to established protocols, e.g. O'Reilly, D.R., Miller, L.K. & Luckow, V.A. "Baculovirus expression vectors. A laboratory manual." Oxford University Press, New York - Oxford, 1994 or David W. Murhammer (ed.). Baculovirus and Insect Cell Expression Protocols, 2nd edition, Methods in Molecular Biology™ 388, Humana Press, Totowa 2007. You may also refer to Fitzgerald et al. (2006), especially pp. 1025-27 and Bieniossek et al. (2008) for protocols on insect cell culture and bacmid preparation. These publications are available for download on the geneva-biotech.com website.

Large plasmid kits (e.g. Qiagen) can also be used to extract recombinant bacmids, but we prefer precipitation of crude DNA with isopropanol from cleared lysate as described in Fitzgerald et al. (2006), especially pp. 1025-27 and Bieniossek et al. (2008) over using kits.

D.6. Determining baculoviral expression

As protein expression in the baculoviral system is driven by the late to very late polyhedrin and p10 promoters, onset of protein expression will show some lag and should be expected to start yielding significant amounts at 15 to 24 hrs. post-infection. Expression will likely peak at around 40 hrs. post-infection but this may vary. Therefore, rapid and reliable methods to determine peak expression are needed. Approaches in use include immunoblot analysis, protein staining techniques or metabolic labeling to chart protein accumulation. Most of these come with a delay due to the necessary handling, e.g. casting gels, preparing protein samples etc.

Expression of target genes from the SynBac[™] bacmid can be determined by SDS-PAGE with or without subsequent Western Blot. Incorporation of tags such as 6xHis (Hexa-His) or FLAG allow purification of proteins and protein complexes. Additionally, you may equip any of your expressed proteins with a reporter protein, e.g. fluorescent proteins such as GFP or YFP.

E. Appendix

E.1. Verifying bacmid integrity in DH10SynBac™ cells

If you create your own glycerol stock or competent cells, to verify bacmid integrity, streak DH10SynBacTM cells on LB plates containing kanamycin (50 μ g/ml), tetracycline (10 μ g/ml), BluoGal (100 μ g/ml) or X-Gal (20 μ g/ml) and IPTG (40 μ g/ml) and incubate overnight at 37°C. All colonies should turn blue. Note that the dye reaction can take >24 hrs. to develop completely. Working stock of bacmid-containing *E.coli* can then be prepared from individual colonies. If, after careful plating, a significant number of colonies show up white, then the stock is likely contaminated and you are advised to re-order competent cells.

E.2. Preparing competent DH10SynBac™ cells

E.2.1 Protocol for electrocompetent cells

Reagents required:

Kanamycin: 50mg/ml stock solution in water (1000x)

Tetracycline: 12 mg/ml stock solution in 70% EtOH (500x)

BluoGal: 100mg/ml stock (1000x)

IPTG: 0.5M (or 1M) stock (1000x)

Sterile 10% glycerol (mix glycerol with MilliQ water, autoclave)

Protect plates from light, store at 4°C

 Streak out DH10SynBac™ cells onto an agar plate with antibiotics (Kan, Tet), X-Gal or BluoGal and IPTG.

All colonies must turn BLUE (can take 24 to 30hrs, sometimes longer if the stabs are old). If there are white colonies, the frozen stock is contaminated.

- 2. Pick one to three blue colonies, prepare a 100mL preculture (make 2 glycerol stocks 30% from preculture) and inoculate 1L LB or 2xTY culture medium adding Kan and Tet as antibiotics. Shake over night at 37 °C or RT.
- 3. Let culture grow to OD600 = 0.5.

- 4. Chill 15 min on ice. Pellet by centrifuging at 4000rpm, 4°C, 15 min.
- 5. Resuspend in 500 ml ICE COLD STERILE 10% Glycerol-solution.
- 6. Centrifuge at 4000rpm, 4°C, 15 min.
- 7. Resuspend in 250 ml ICE COLD STERILE 10% Glycerol-solution.
- 8. Centrifuge at 4000rpm, 4°C, 15 min.
- 9. Resuspend in 50 ml ICE COLD STERILE 10% Glycerol-solution.
- 10. Centrifuge at 4000rpm, 4°C, 15 min (in 50ml Falcon tube, table top centrifuge).
- 11. Resuspend in 2 ml ICE COLD STERILE 10% Glycerol-solution.
- 12. Prepare 100 μl aliquots (in sterile Eppendorf tubes, labeled either MBT or MBY)
- 13. Flash freeze in liquid nitrogen, store at -70°C
- 14. Test cells by streaking out (!!) from one frozen aliquot on agar plate containing antibiotics Kan,

 Tet plus X-Gal or BluoGal/ IPTG.

All colonies must be BLUE (can take 24 to 30 hrs.). If there are white colonies, the competent cells are unusable and must be made anew.

15. Test one frozen aliquot by "blank" electroporation (1.8kV) – actual kV should be 4-4.5 or more.

You may also want to take a look at other protocols, e.g. the protocol in Sambrook and Russell's Molecular Cloning: A Laboratory Manual or the protocol for enhanced electrotransformation efficiency by Wu et al. in the Electronic Journal of Biotechnology.

E.2.2 Protocol for chemically competent cells

Aside from the protocol provided below, there are numerous protocols for preparing competent cells for heat-shock transformation, e.g. the classical CaCl₂ method, the TSS protocol (originally by Chung et al., 1989, PNAS USA, Vol. 86, pp. 2172-2175), etc.

You will find various protocols online, e.g.

http://francois.schweisguth.free.fr/protocols/TSS_competent_cell_preparation.pdf;

http://www.personal.psu.edu/dsg11/labmanual/DNA_manipulations/Comp_bact_by_TSS.htm;

http://openwetware.org/wiki/Preparing chemically competent cells

or in Sambrook and Russell.

Preparing competent E.coli (TOP10) cells with the CaCl₂ method

- Grow E.coli strain by streaking on LB agar plates; incubate o/n at 37 °C
- 2. Grow a few colonies in 20ml LB at 37 ° C o/n (to OD 600=1.8)
- 3. Culture 13ml into 100ml freshly prepared LB (to OD 600 = 0.25)
- 4. Incubate at 37 ° C on a shaker (180 rpm) for 1 hour (OD600=0.5-0.6, depending on the strain)
- 5. Distribute the culture into 50ml centrifuge tubes and centrifuge at 8000 rpm, 4 ° C for 5 min (or, alternatively, in 50 ml Falcon tubes, spin at -4000rpm for 6 min)
- 6. Discard the supernatant completely; then add 10ml of ice cold 0.1M CaCl₂ (keep the cells on ice and resuspend gently)
- 7. Incubate the solution on ice for 15 min, then centrifuge at 8000 rpm at 4 ° C for 5 min (or, alternatively, in 50 ml Falcon tubes, spin at -4000rpm for 6 min)
- 8. Remove the supernatant completely and let the pellet air dry
- 9. Add 2.8ml ice cold 0.1M CaCl₂ and mix gently. Keep on ice till use **OR**
- 10. Add 1.2 ml of sterile ice cold 80% glycerol to a tube with 2.8ml cells each and mix gently; transfer 100 μ l aliquots into sterile 1.5 ml Eppendorf microcentrifuge tubes (use gloves and Bunsen burner flame); keep the Eppendorfs in ice until snap-freezing the bacterial cells in liquid N₂; store at -80°C. **Alternatively**, transfer 100 μ l aliquots into sterile 1.5 ml Eppendorfs, add 32 μ l of 80% glycerol, freeze in dry ice/EtOH bath or liquid N₂ and store at -80°C.

E.3. Preparing bacterial stocks from agar stabs

We recommend preparing your personal bacterial stock from the agar stabs you received in the kit or transforming your laboratory strain of choice with the vectors (please note that for the donor vectors this needs to be a pir+ strain). This is advisable since agar stabs only have a limited shelf life (3-12 months depending on storage conditions). Permanent cultures in glycerol or DMSO, on the other side, should yield viable colonies even after 1 year and beyond if properly stored.

To generate your bacterial stock for long-term storage, streak bacteria from the agar stab onto an appropriate selective plate (refer to the vector maps for acceptor and donor vectors) or plates without antibiotics (pirHC and pirLC strains; we recommend to test these strains against a panel of antibiotics to be on the safe side; no growth of colonies should be observed under conditions of antibiotic selection). Incubate the plates over night at 37°C and then proceed to prepare stocks from

individual colonies for long-term storage according to your protocol of choice (glycerol, DMSO, etc.), as described, for example, in Inoue et al. (1990), Molecular Cloning (Sambrook and Russell, 2000), Current Protocols in Molecular Biology (Ausubel et al., 1994), etc.

E.4. SynBac™ vectors: maps, sequences, restriction

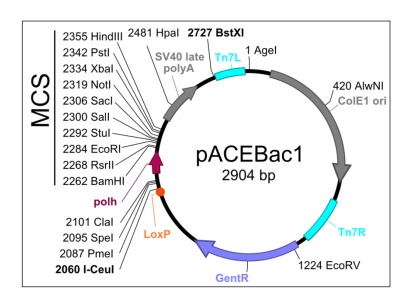
Note: All acceptor and donor vector sequences can be provided in electronic format. These sequences contain all relevant information such as unique restriction sites, oris, resistance markers, etc. that is also shown in the circle maps. Request your set of vector files and accompanying files from Geneva Biotech at contact@geneva-biotech.com.

In the next section, the acceptor and donor vectors are presented as circle maps and, in addition, the multiple cloning site (MCS) of each vector is shown featuring important unique restriction sites. Moreover, you will find, for the purposes of designing a restriction strategy, a non-exhaustive list of restriction endonucleases that cut once, twice or not at all. Additional restriction sites can be identified with any sequence analysis software, e.g. VectorNTI, ApE, etc. or by using online tools such as WebCutter 2.0 (http://rna.lundberg.gu.se/cutter2) or the NEB cutter V2.0 (http://rna.lundberg.gu.se/cutter2).

Finally, for verifying vector identity, representative digestion patterns are listed in a table.

E.4.1 Acceptor vectors

E.4.1.1 pACEBac1: 2904 bp



Multiple Cloning Site

AatI

BamHI RsrII BssHII EcoRI StuI SalI SacI

 $\texttt{CGGGCGC} \underbrace{\texttt{GGATCCCGGTCCGAAGCGCGCGGAATTCAAAGGCCTACGTCGACGAGCTCACTTGT}$

NotI BstBI XbaI PstI HindIII
CGCGGCCGCTTTCGAATCTAGAGCCTGCAGTCTCGACAAGCTTGTCGAGAAGTACTAGAGGA

Enzymes that cut pACEBac1 once

2294	Aatl	1	Age I	520	AlwNI	2127	Asel, Asnl
1629	Aspl	2715	AvrII	2262	BamHI, Bstl	2310	Banll
2619	Bcll	1307	BsaAI	2102	Bsp106I, BspXI	2277	BssHII
2329	BstBI	2744	BstXI	2075	Bsu36I	2102	ClaI
2284	EcoRI	1226	EcoRV	890	FspI	2355	HindIII
2483	Hpal	2470	Mfel	1111	MscI, MluNI	2320	Notl
2109	Nrul	2090	Pmel	2346	Pstl	114	Rcal
2269	RsrII	2310	SacI	1169	SacII	2300	Sall
2370	Scal	2095	Spel	2294	Stul	2334	Xbal

Bold type: restriction enzymes cutting in the MCS

Enzymes that cut pACEBac1 twice

2301, 2722	Accl	949, 1418	BgIII	2118, 2228	BseAl, BsiMl
2470, 2569	Bsml	1169, 2320	BstZI	2118, 2228	Mrol
2125, 2225	Sspl				

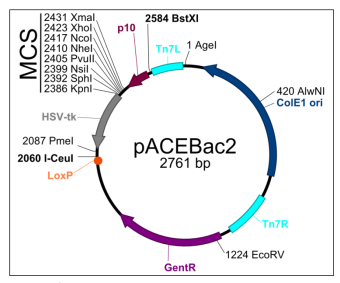
Enzymes that do not cut pACEBac1

Acc65I	AfIII	Apal	Bcll	Bgll	BstEl	EcoNI	Mlul
Nael	Narl	Ncol	Ndel	Nhel	Nsil	Pvul	Sfil
Smal/Xmal	SnaBl	SphI	Xcml	XhoI	XmnI		

Sequence

5**′** –

 $\verb|ttaatttaaaaggatctaggtgaagatcctttttgataatctcatgaccaaaatcccttaacgtgagttttc||$ $\verb|ttttccgaaggtaactggcttcagcagagcgcagataccaaatactgttcttctagtgtagccgtagttagg|\\$ ccaccacttcaagaactctgtagcaccgcctacatacctcgctctgctaatcctgttaccagtggctgctgc $\verb|cagtggcgata| agtcgtgtctta| ccgggttggactca| agacgatagtta| ccggata| aggcgcagcggtcggg| cgggtcggg| cggg| cggg|$ $\verb|ctgaacgggggttcgtgcacacagcctaggagcgaacgacctacaccgaactgagatacctacagcg| \\$ tgagctatgagaaagcgccacgcttcccgaagggagaaaggcggacaggtatccggtaagcggcagggtcgg aa caggagag cgcacgagggagcttccagggggaaacgcctggtatctttatagtcctgtcgggtttcgccaggcctttttacggttcctggccttttgctggcctttttgctcacatgttctttcctgcgttatcccctgattg $\verb|acttgggtcgctcttcctgtggatgcgcagatgccctgcgtaagcgggtgtggggcggacaataaagtcttaa||$ actgaacaaaatagatctaaactatgacaataaagtcttaaactagacagaatagttgtaaactgaaatcag tccagttatgctgtgaaaaagcatactggacttttgttatggctaaagcaaactcttcattttctgaagtgc $\verb|ttaccgaacaactccgcggccgggaagccgatctcggcttgaacgaattgttaggtggcggtacttgggtcg|$ atatcaaagtgcatcacttcttcccgtatgcccaactttgtatagagagccactgcgggatcgtcaccgtaa tctgcttgcacgtagatcacataagcaccaagcgcgttggcctcatgcttgaggaggattgatgagcgcggtggcaatgccctgcctccggtgctcgccggagactgcgagatcatagatatagatctcactacgcggctgctca aacttgggcagaacgtaagccgcgagagcgccaacaaccgcttcttggtcgaaggcagcaagcgcgatgaat $\verb|gtcttactacggagcaag| tcccgaggtaatcggagtccggctgatgttgggagtaggtggctacgtctccg| |$ aactcacgaccgaaaagatcaagagcagcccgcatggatttgacttggtcagggccgagcctacatgtgcga atgatgcccatacttgagccacctaactttgttttagggcgactgccctgctgcgtaacatcgttgctgctgcgtaacatcgttgctgctccataacatcaaacatcgacccacggcgtaacgcgcttgctgcttggatgcccg aggcatagactgtacaaaaaacagtcataacaagccatgaaaaccgccactgcgccgttaccaccgctgcg ttcggtcaaggttctggaccagttgcgtgagcgcatacgctacttgcattacagtttacgaaccgaacaggcatcgcattacaggttacgaaccgaacaggcatacgctacttgcattacaggttacgaaccgaacaggcatacgctacttgcattacaggttacgaaccgaacaggcatacgctacttgcattacaggttacgaaccgaacaggcatacgctacttgcattacaggttacaggaaccaggcatacgctacttgcattacaggttacaggaaccaggaacaggcatacgctacttgcattacaggttacaggaaccaggaacaggcatacgcatacgctacttgcattacaggttacaggaaccagaaccaggaacaggcatacgcatacgcatacgctacttgcattacaggttacaggaaccaggaaccaggaacaggcatacgcatacgctacttgcattacaggttacaggaaccaggaaccaggaacaggcatacgcatacgcatacaggattatgtcaactgggttcgtgccttcatccgtttccacggtgtgcgtcacccggcaaccttgggcagcagcga agtcgccataacttcgtatagcatacattatacgaagttatctgtaactataacggtcctaaggtagcgagt $\verb|ttaaacactag| tatcgattcgcgacctactccggaatattaatagatcattggagataattaaaatgataacc| \\$ $\verb|atctcgcaaataaataagtattttactgttttcgtaacagttttgtaataaaaaaacctataaatattccgg|$ attattcataccgtcccaccatcgggcgcggatcccggtccgaagcgcgcggaattcaaaggcctacgtcga $\verb|cgagctcacttgtcgcggccgctttcgaatctagagcctgcagtctcgacaagcttgtcgagaagtactaga|\\$ ggatcataatcagccataccacatttgtagaggttttacttgctttaaaaaaacctcccacacctcccctga acctgaaacataaaatgaatgcaattgttgttgttaacttgtttattgcagcttataatggttacaaataaa tcaatgtatcttatcatgtctggatctgatcactgcttgagcctagaagatccggctgctaacaaagcccga a agga agct gagt tggctgctgccaccgct gagca at aactat cata acccct agggt at acccat cta at the same of thggaaccagataagtgaaatctagttccaaactattttgtcatttttaattttcgtattagcttacgacgcta cacccagttcccatctattttgtcactcttccctaaataatccttaaaaaactccatttccacccctcccagt tcccaactattttgtccgcccaca-3'



E.4.1.2 pACEBac2: 2761 bp

Multiple Cloning Site

KpnI PaeI NheI Acc65I SphI NsiI PvuII

NcoI XhoI XmaI BclI BbsI(-)

C C A T G G C T C G A G A T C C C G G G T G A T C A A G T C T T C G T C G A G T G A T

Enzymes that cut pACEBac2 once

2386	Acc65I	1	Age I	425	AlwNI	2496	Asel, Asnl
1629	Aspl	2146	Banll	2444	Bbsl (-)	2437	Bcll
2223	Bsal	666	Bsil	1582	BsmBI	2591	BstXI
1169	BstZI, Eagl KspI	2075	Bsu36I	1226	EcoRV	2390	Kpnl
1111	MluNI, MscI	2417	Ncol	2410	Nhel	2403	Nsil
2090	Pmel	2407	Pvull	114	Rcal	1169	SacII
2433	Smal, Xmal	2396	SphI	2423	Xhol		

Bold type: restriction enzymes cutting in the MCS (-) cutting on the negative strand

Enzymes that cut pACEBac2 twice

2569, 2579 Accl 949, 1418 Bglll 312, 1077 Eco57l

594, 1471 Haell 464, 1546 Hinfl

Enzymes that do not cut pACEBac2

Aatil Afiii Apal Asci Avrii BamHi Bcli Bgli

BssHII	ClaI	Drall	EcoNI	EcoRI	Hpal	Mfel	Nael
Narl	Ndel	Notl	Nrul	PstI	Pvul	RsrII	SacI
Sall	Sbfl	Scal	Sfil	SnaBl	Spel	Stul	Xbal
XcmI	XmnI						

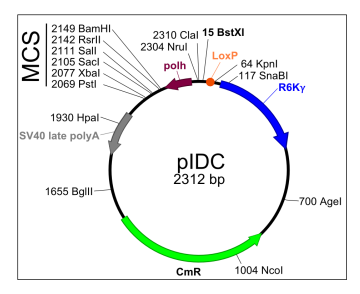
Sequence

5**′** -

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E.4.2 Donor vectors

E.4.2.1 pIDC: 2312 bp



Multiple Cloning Site

PstI XbaI SacI

 $\verb|TTCTCGACAAGCTT|| \underline{GTCGAGACTGCAGGCTCTAGATTCGAAAGCGGCCGCGACTAGTGAGCTC| \\$

AccI AatI

SalI StuI BssHII RsrII BamHI

Enzymes that cut pIDC once

2121	Aatl	64	Acc65I	700	Age I	760	Aspl
1696	AvrII	2149	BamHI	2109	BanII	1655	BgIII
2134	BssHII	2149	Bstl (BamHl)	15	BstXI	650	Bsu36I
2310	Clal	1930	Hpal	64	KpnI	1941	Mfel
1040	MscI, MluNI	1004	Ncol	2304	Nrul	2069	PstI
2142	RsrII	2105	SacI	68	SacII	2111	Sall
117	SnaBl	2119	SseBI, Stul	2109	SstI	2077	Xbal

Bold type: restriction enzymes cutting in the MCS

Enzymes that cut pIDC twice

281, 2285	Asel, Asnl	766, 2108	BstBI	70, 2090	BstZI, EagI
1305, 2127	EcoRI	359, 2056	HindIII	70, 2090	Notl
580, 1407	Pvull	890, 2043	Scal	4, 2099	Spel
1004, 1696	Styl				

Enzymes that do not cut pIDC

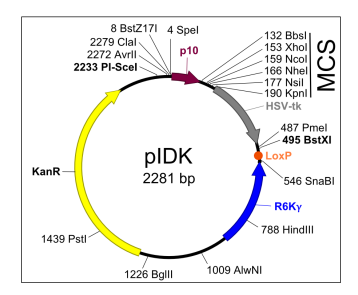
AatII	AfIII	AlwNI	Apal	Ascl	Bcll	Bgll	BstZ17I
Drall	EcoNI	EcoRV	Mbol	MluI	Nael	Narl	Ndel
Nhel	Nsil	Pmel	Pvul	SbfI	Sfil	Smal	SphI
XcmI	XhoI	Xmal	XmnI				

Sequence

5**′** –

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E.4.2.2 pIDK: 2281 bp



Multiple Cloning Site

BbsI XhoI NcoI NheI

ACAATCACTCGACGAAGACTTGATCACCCGGGATCTCGAGCCATGGTGCTAGCAGCTG

Acc65I

NsiI KpnI

 $\underline{\texttt{A} \texttt{ T} \texttt{ G} \texttt{ C} \texttt{ A} \texttt{ T} \texttt{ A} \texttt{ G} \texttt{ C} \texttt{ A} \texttt{ T} \texttt{ G} \texttt{ C} \texttt{ G} \texttt{ G} \texttt{ T} \texttt{ A} \texttt{ C}} \texttt{ C} \texttt{ G} \texttt{ G} \texttt{ G}$

Enzymes that cut pIDK once

190	Acc65I, KpnI	1014	AlwNI	1508	Aspl	2272	AvrII, BlnI
132	BbsI	1226	BgIII	363	Bsal	1787	BssHII
2072	BstBI	2744	BstXI	1296	BstZI, EagI	8	BstZ17I
1492	FspI	788	HindIII	1472	MscI, MluNI	159	Ncol
166	Nhel	181	Nsil	490	Pmel	1443	PstI
1906	RsrII	2072	Sful	548	SnaBI	4	Spel
1206	Xcml	153	Xhol				

Bold type: restriction enzymes cutting in the MCS

Enzymes that cut pIDK twice

82, 711	Asel, Asnl	1145, 1393	Bbel	139, 1231	Bcll
1744, 1954	Earl	1145, 1393	Haell	1892, 2175	Nael
188, 1795	Pael	147, 2229	Smal, Xmal	188, 1795	SphI

Enzymes that do not cut pIDK

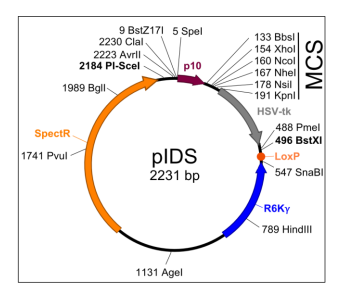
AatII	AfIII	Agel	Apal	ApaLl	Ascl	BamHI	Bgll
Drall	EcoNI	EcoRI	EcoRV	Drall	Mbol	Mlul	Ndel
Pvul	SbfI	Sfil	Stul	Xbal	XmnI		

Sequence

5**′** –

gatactagtatacggacctttaattcaacccaacacaatatattatagttaaaataagaattattatcaa at catttg tatatta atta aatactatactg ta aatta cattttattta caatcactcg acgaa gacttgatcacccgggatctcgagccatggtgctagcagctgatgcatagcatgcggtaccgggagatggggg $\verb| aataaaacgcacgggtgttgggtcgtttgttcataaacgcggggttcggtcccagggcttggcactctgt| \\$ $\verb|cgataccccaccgagaccccattgggaccaatacgcccgcgtttcttccttttccccaccccaaccccc|\\$ aagttcgggtgaaggcccagggctcgcagccaacgtcggggcggcaagccctgccatagccactacggg taagctctcatgtttcacgtactaagctctcatgtttaacgtactaagctctcatgtttaacgaactaa $\verb"accctcatggctaacgtactaagctctcatggctaacgtactaagctctcatgtttcacgtactaagct"$ $\verb|ctcatgtttgaaca| ataaaattaatataaatcagcaacttaaatagcctctaaggttttaagttttata| \\$ agaa aa aa aa aa gaata ta ta aa ggctttta aa gctttta aa ggttta ac ggttg tg gac aa caa gcc ag ggaar ac gaata ag gaata aa gaata ag ${\tt tgtaacgcactgagaagcccttagagcctctcaaagcaattttcagtgacacaggaacacttaacggct}$ $\tt gacagaattagcttcacgctgccgcaagcactcagggcgcaagggctgctaaaggaagcggaacacgta$ gaaagccagtccgcagaaacggtgctgaccccggatgaatgtcagctactgggctatctggacaaggga aaacgcaagcgcaaagagaaagcaggtagcttgcagtgggcttacatggcgatagctagactgggcggt tttatggacagcaagcgaaccggaattgccagctggggcgccctctggtaaggttgggaagccctgcaa ${\tt agtaaactggatggctttcttgccgccaaggatctgatggcgcaggggatcaagatctgatcaagagac}$ aggatgaggatcgtttcgcatgattgaacaagatggattgcacgcaggttctccggccgcttgggtgga $\tt gaggctattcggctattgactgggcacaacagacaatcggctgctctgatgccgccgtgttccggctgtc$ $\verb|ggcagcggctatcgtggctagcgacgggggttccttgcgcagctgtgctcgacgttgtcactga|\\$ agcgggaagggactggctattgggcgaagtgccggggcaggatctcctgtcatctcaccttgctcctgccgagaaagtatccatcatggctgatgcaatgcggcggctgcatacgcttgatccggctacctgccc attcgaccaccaagcgaaacatcgcatcgagcgagcacgtactcggatggaagccggtcttgtcgatca $\verb|ggatgatctggacgaagagcatcaggggctcgcgccagccgaactgttcgccaggctcaaggcgcgcat|$ gcccgacggcgaggatctcgtcgtgacacatggcgatgcctgcttgccgaatatcatggtggaaaatgg $\verb|ccgcttttctggattcatcgactgtggccggctgggtgtggcggaccgctatcaggacatagcgttggc|\\$ $\verb|tacccgtgatattgctgaagagcttggcggaatgggctgaccgcttcctcgtgctttacggtatcgc| \\$ $\verb|cgctcccgattcgcatcgccttctatcgccttcttgacgagttcttctgagcgggactctgggg|$ $\verb|ttcgaaattgaccgaccaagcgacgcccaacctgccatcacgagatttcgattccaccgccgccttctat|$ gaaaggttgggcttcggaatcgttttccgggacgccggctggatgatcctccagcgcggggatctcatg ctggagttcttcgcccaccccgggatctatgtcgggtgcggagaaagaggtaatgaaatggcacctagg tatc-3'

E.4.2.3 pIDS: 2231 bp



Multiple Cloning Site

SmaI

BbsI XmaI XhoI NcoI NheI NsiI

 $\texttt{CAATCA} \underline{\texttt{CTCGACGAAGACTTGATCACCCGGGATCTCGAGCCATGGTGCTAGCAGCTGATGCAT}$

Acc65I SphI KpnI AGCATGCGGTACC

Enzymes that cut pIDS once

191	Acc65I, KpnI	1131	Agel	1638	ApaLI	2223	AvrII, BlnI
133	BbsI	1989	BgII	2087	BssHII	496	BstXI
9	BstZ17I	1081	Bsu36I	2230	ClaI	789	HindIII
2012	Nael	160	Ncol	167	Nhel	178	Nsil
488	Pmel	1741	Pvul	146	Smal, Xmal	547	SnaBl
5	Spel	185	SphI	154	Xhol		

Bold type: restriction enzymes cutting in the MCS

Enzymes that cut pIDS twice

82, 711	Asel, Asnl	489, 785	Dral	172, 1011	Pvull
82, 711	VspI				

Enzymes that do not cut pIDS

AatII	Afel	AlwNI	Apal	Ascl	BamHI	BgIII	BstBI
Drall	EcoNI	EcoRI	EcoRV	HincII	Hpal	Mbol	Mlul

Mscl Narl Ndel Notl Nrul Pstl Rsrll Sacl
Sall Sbfl Sfil Stul Xbal Xcml Xmnl

Sequence

5**′** –

cqatactagtatacqqacctttaattcaacccaacacaatatattataqttaaataaqaattattatca ttgatcacccgggatctcgagccatggtgctagcagctgatgcatagcatgcggtaccgggagatgggg gaataaaacgcacgggtgttgggtcgtttgttcataaacgcggggttcggtcccagggctggcactctg $\verb|tcgataccccaccgagaccccattgggaccaatacgcccgcgtttcttccttttccccaccccaacccc|$ caagttcgggtgaaggcccagggctcgcagccaacgtcggggcggcaagccctgccatagccactacgg gtacgtttaaacccatgtgcctggcagataacttcgtataatgtatgctatacgaagttatggtacgta ctaagctctcatgtttcacgtactaagctctcatgtttaacgtactaagctctcatgtttaacgaacta aaccctcatggctaacgtactaagctctcatggctaacgtactaagctctcatgtttcacgtactaagc tctcatgtttgaacaataaaattaatataaatcagcaacttaaatagcctctaaggttttaagttttat aaqaaaaaaaaqaatatataaqqctttttaaaqctttttaaqqttttaacqqttgtqqacaacaaqccaqqq atgtaacgcactgagaagcccttagagcctctcaaagcaattttgagtgacacaggaacacttaacggc tgacataattcagcttcacgctgccgcaagcactcagggcgcaagggctgctaaaaggaagcggaacacg tagaaagccagtccgcagaaacggtgctgaccccggatgaatgtcagctgggaggcagaataaatgatc atatcqtcaattattacctccacqqqqaqaqcctqaqcaaactqqcctcaqqcatttqaqaaqcacacq gtcacactgcttccggtagtcaataaaccggtaagtagcgtatgcgctcacgcaactggtccagaacct tgaccgaacgcagcggtggtaacggcgcagtggcggttttcatggcttgttatgactgtttttttgggg tacagtctatgcctcgggcatccaagcagcaagcgcgttacgccgtgggtcgatgtttgatgttatgga gcagcaacgatgttacgcagcagggcagtcgccctaaaacaaagttaaacatcatgagggaagcggtga tcgccgaagtatcgactcaactatcagaggtagttggcgtcatcgagcgccatctcgaaccgacgttgc tggccgtacatttgtacggctccgcagtggatggcggcctgaagccacacagtgatattgatttgctggttacggtgaccgtaaggcttgatgaaacaacgcggcgagctttgatcaacgaccttttggaaacttcgg $\verb|cttcccctggagagagagattctccgcgctgtagaagtcaccattgttgtgcacgacgacatcattc|\\$ cgtggcgttatccagctaagcgcgaactgcaatttggagaatggcagcgcaatgacattcttgcaggta tcttcgagccagccacgatcgacattgatctggctatcttgctgacaaaagcaagagaacatagcgttg ccttggtaggtccagcggcggaggaactctttgatccggttcctgaacaggatctatttgaggcgctaa $\verb|atgaaaccttaacgctatggaactcgccgactgggctggcgatgagcgaaatgtagtgcttacgt|$ tgtcccgcatttggtacagcgcagtaaccggcaaaatcgcgccgaaggatgtcgctgccgactgggcaa tggagcgcctgccggcccagtatcagcccgtcatacttgaagctagacaggcttatcttggacaagaagaagatcgcttggcctcgcgcgcagatcagttggaagaatttgtccactacgtgaaaggcgagatcacca aggtagtcggcaaataatgtctaacaattcgttcaagccgacggatctatgtcgggtgcggagaaagag gtaatgaaatggcacctaggtat-3'

E.4.3. Analytical restriction digest patterns

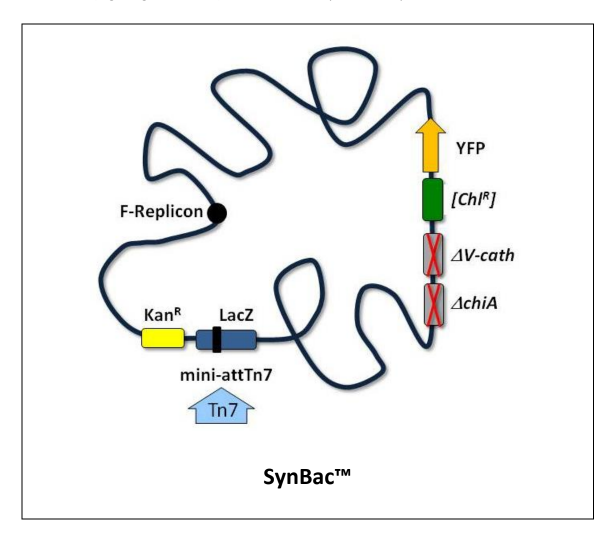
All SynBac™ acceptor and donor vectors yield characteristic restriction digest patterns when plasmid DNA is restricted with HindIII and/or Hinfl or a combination of both.

Vector	Length [bp]	Restriction	Fragment(s) [bp]
pACEBac1	2904	HindIII	2904
		Hinfl	1082, 1037, 558, 227
pACEBac2	2761	HindIII	2761
		Hinfl	1679, 1082
pIDC	2312	HindIII	1697, 615
		Hinfl	1125, 960, 227
		HindIII/Hinfl	1100, 597, 363, 227, 25
pIDK	2281	HindIII	2281
		Hinfl	2000, 134, 58, 52, 37
		HindIII/Hinfl	1087, 913, 134, 58, 52, 37
pIDS	2231	HindIII	2231
		Hinfl	2018, 213
		HindIII/Hinfl	1413, 605, 213

E.4.4 DH10SynBac™ baculoviral genome

Genome (size approx. 100 kb) is a derivative of the *Autographa californica* nucleopolyhedrovirus (AcMNPV) genome. It has been genetically engineered for improved protein production and reduced protein degradation (details on p. 18). In addition, it contains an element for accepting donor DNA into its transposition acceptor site (mini-attTn7) that concomitantly allows bluewhite selection to identify successful transposition events. The bacmid possesses the F replicon from the F plasmid that keeps the plasmid copy number at 1 (single copy).

It is hosted in DH10SynBac[™] *E.coli* cells and can be isolated from the bacteria using commercial "large construct" kits (e.g. Qiagen) or appropriate protocols for preparation of large DNA molecules (e.g. King et al., 2007). Also refer to Chapter D.5 on p. 30.



NOTES

F. References

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