

## Champion<sup>™</sup> pET Directional TOPO<sup>®</sup> Expression Kits

Five-minute, directional TOPO<sup>®</sup> Cloning of blunt-end PCR products into vectors for high-level, inducible expression in *E. coli* Catalog nos. K100-01, K101-01, K102-01, K151-01, K200-01

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**User Manual** 

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### **TOPO<sup>®</sup> Cloning Procedure for Experienced Users**

Introduction	This quick reference sheet is provided for experienced users of the TOPO <sup>®</sup>
	Cloning procedure. If you are performing the TOPO® Cloning procedure for the
	first time, we recommend that you follow the detailed protocols provided in the
	manual.

Step		Action	
Design PCR Primers	•	Include the 4 base pair sequences (CACC) necessary for directional cloning on the 5' end of the forward primer.	
	•	Design the primers such that your gene of interest will be optimally expressed and fused in frame with any epitope tags, if desired.	
Amplify Your Gene of Interest	1.	Use a thermostable, proofreading DNA polymerase and the PCR primers above to produce your blunt-end PCR product.	
	2.	Use agarose gel electrophoresis to check the integrity and yield of your PCR product.	
Perform the TOPO <sup>®</sup> Cloning Reaction	1.	Set up the following TOPO <sup>®</sup> Cloning reaction. <b>For optimal results, use a 0.5:1 to 2:1 molar ratio of PCR product:TOPO<sup>®</sup> vector</b> .	
		<b>Note:</b> If you plan to transform electrocompetent <i>E. coli</i> , use Dilute Salt Solution in the TOPO <sup>®</sup> Cloning reaction.	
		Fresh PCR product 0.5 to 4 µl	
		Salt Solution 1 µl	
		Sterile water add to a final volume of $5 \mu l$	
		<u>TOPO<sup>®</sup> vector 1 µl</u>	
		Total volume 6 µl	
	2.	Mix gently and incubate for 5 minutes at room temperature.	
	3.	Place on ice and proceed to transform One Shot <sup>®</sup> TOP10 chemically competent <i>E. coli</i> , below.	
Transform TOP10 Chemically Competent	1.	Add 3 μl of the TOPO <sup>®</sup> Cloning reaction into a vial of One Shot <sup>®</sup> TOP10 chemically competent <i>E. coli</i> and mix gently.	
E. coli	2.	Incubate on ice for 5 to 30 minutes.	
	3.	Heat-shock the cells for 30 seconds at 42°C without shaking. Immediately transfer the tube to ice.	
	4.	Add 250 µl of room temperature S.O.C. medium.	
	5.	Incubate at 37°C for 1 hour with shaking.	
	6.	Spread 100-200 $\mu$ l of bacterial culture on a prewarmed selective plate and incubate overnight at 37°C.	
-			

## **Control Reaction** We recommend using the Control PCR Template and the Control PCR Primers included with the kit to perform the control reaction. See the protocol on pages 37-39 for instructions.

#### **Kit Contents and Storage**

### **Types of Kits** This manual is supplied with the following pET Directional TOPO<sup>®</sup> Expression kits listed below.

Kit	Quantity	Catalog no.
Champion <sup>™</sup> pET100 Directional TOPO <sup>®</sup> Expression Kit	20 reactions	K100-01
Champion <sup>™</sup> pET101 Directional TOPO <sup>®</sup> Expression Kit	20 reactions	K101-01
Champion <sup>™</sup> pET102 Directional TOPO <sup>®</sup> Expression Kit	20 reactions	K102-01
Champion <sup>™</sup> pET151 Directional TOPO <sup>®</sup> Expression Kit	20 reactions	K151-01
Champion <sup>™</sup> pET200 Directional TOPO <sup>®</sup> Expression Kit	20 reactions	K200-01

## **Shipping/Storage** The Champion<sup>™</sup> pET Directional TOPO<sup>®</sup> Expression Kits are shipped on dry ice. Each kit contains three boxes. Upon receipt, store the boxes as detailed below.

Box	Item	Storage
1	pET TOPO <sup>®</sup> Reagents	-20°C
2	One Shot® TOP10 Chemically Competent E. coli	-80°C
3	BL21 Star <sup>™</sup> (DE3) One Shot <sup>®</sup> Chemically Competent <i>E. coli</i>	-80°C

# Vectors and<br/>PrimersEach Champion™ pET Directional TOPO® Expression Kit contains a pET-TOPO®<br/>vector, an expression control, and primers for sequencing. The pET-TOPO®<br/>vector, expression control, and primers differ depending on the kit. The table<br/>below lists the vectors and primers supplied with each kit. For details on the<br/>amount of each component provided, see the next page.

pET TOPO® Kit	Catalog no.	TOPO <sup>®</sup> Vector	Expression Control	Primers
pET100	K100-01	pET100/D-TOPO®	pET100/D/lacZ	Τ7
				T7 Reverse
pET101	K101-01	pET101/D-TOPO®	pET101/D/lacZ	Τ7
				T7 Reverse
pET102	K102-01	pET102/D-TOPO®	pET102/D/lacZ	TrxFus Forward
				T7 Reverse
pET151	K151-01	pET151/D-TOPO®	pET151/D/lacZ	Τ7
				T7 Reverse
pET200	K200-01	pET200/D-TOPO®	pET200/D/lacZ	Τ7
				T7 Reverse

#### Kit Contents and Storage, continued

#### pET TOPO<sup>®</sup> Reagents

pET TOPO<sup>®</sup> Reagents (Box 1) are listed below. Each box includes PCR reagents and the appropriate vectors and primers. **Note that the user must supply a thermostable, proofreading polymerase and the appropriate PCR buffer. Store Box 1 at -20°C.** 

Item	Concentration	Amount
pET TOPO <sup>®</sup> vector, TOPO <sup>®</sup> -adapted	15-20 ng/μl linearized plasmid DNA in:	20 µl
	50% glycerol	
	50 mM Tris-HCl, pH 7.4 (at 25°C)	
	1 mM EDTA	
	2 mM DTT	
	0.1% Triton X-100	
	100 μg/ml BSA	
	30 µM bromophenol blue	
dNTP Mix	12.5 mM dATP	10 µl
	12.5 mM dCTP	
	12.5 mM dGTP	
	12.5 mM dTTP	
	in water, pH 8	
Salt Solution	1.2 M NaCl	50 µl
	0.06 M MgCl <sub>2</sub>	
Sterile Water		1 ml
Forward Sequencing Primer	$0.1 \ \mu g/\mu l$ in TE Buffer, pH 8	20 µl
Reverse Sequencing Primer	0.1 μg/μl in TE Buffer, pH 8	20 µl
Control PCR Primers	0.1 μg/μl <b>each</b> in TE Buffer, pH 8	10 µl
Control PCR Template	0.1 μg/μl in TE Buffer, pH 8	10 µl
Expression Control Plasmid	0.01 μg/μl in TE buffer, pH 8	10 µl

#### Kit Contents and Storage, continued

#### Sequences of the Primers

Each Champion<sup>™</sup> pET Directional TOPO<sup>®</sup> Expression Kit provides a forward and reverse sequencing primer to facilitate sequence analysis of your expression constructs (see the table on page vi for the specific primers included with each kit). The sequences of the forward and reverse primers are listed below. Two micrograms of each primer are supplied.

Primer	Sequence	pMoles Supplied
Τ7	5'-TAATACGACTCACTATAGGG-3'	327
TrxFus Forward	5'-TTCCTCGACGCTAACCTG-3'	371
T7 Reverse	5'-TAGTTATTGCTCAGCGGTGG-3'	325

#### One Shot<sup>®</sup> TOP10 Reagents

The table below lists the items included in the One Shot<sup>®</sup> TOP10 Chemically Competent *E. coli* kit (Box 2). Transformation efficiency is at least  $1 \times 10^9$  cfu/µg DNA. **Store Box 2 at -80°C.** 

Item	Composition	Amount
S.O.C. Medium	2% Tryptone	6 ml
(may be stored at room	0.5% Yeast Extract	
temperature or +4°C)	10 mM NaCl	
	2.5 mM KCl	
	10 mM MgCl <sub>2</sub>	
	10 mM MgSO <sub>4</sub>	
	20 mM glucose	
TOP10 cells		21 x 50 μl
pUC19 Control DNA	10 pg/μl in 5 mM Tris-HCl, 0.5 mM EDTA, pH 8	50 µl

#### Kit Contents and Storage, continued

#### BL21 Star<sup>™</sup>(DE3) One Shot<sup>®</sup> Reagents

The table below describes the items included in the BL21 Star<sup>T</sup>(DE3) One Shot<sup>®</sup> Chemically Competent *E. coli* kit (Box 3). Transformation efficiency is at least 1 x 10<sup>8</sup> cfu/µg DNA. **Store Box 3 at -80°C.** 

Item	Composition	Amount
S.O.C. Medium	2% Tryptone	6 ml
(may be stored at room	0.5% Yeast Extract	
temperature or $+4^{\circ}C$ )	10 mM NaCl	
	2.5 mM KCl	
	10 mM MgCl <sub>2</sub>	
	10 mM MgSO <sub>4</sub>	
	20 mM glucose	
BL21 Star <sup>™</sup> (DE3)		21 x 50 μl
pUC19 Control DNA	10 pg/µl in 5 mM Tris-HCl, 0.5 mM EDTA, pH 8	50 µl

## Genotype of<br/>TOP10Use this *E. coli* strain for general cloning of blunt-end PCR products into the pET<br/>TOPO® vectors.Complement Former A (unrum hed BMC unruPC) @80/ac7AM15 AlacY74 useA1 areD120

**Genotype:**  $F^{-}mcrA \Delta(mrr-hsdRMS-mcrBC) \Phi 80lacZ\Delta M15 \Delta lacX74 recA1 araD139 \Delta(ara-leu)7697 galU galK rpsL (Str<sup>R</sup>) endA1 nupG$ 

## Genotype of BL21 Use this *E. coli* strain for expression only. Do not use these cells to propagate or maintain your construct.

**Genotype:** F<sup>-</sup> *ompT hsdS*<sub>B</sub> (r<sub>B</sub>-m<sub>B</sub>-) *gal dcm rne131* (DE3)

The DE3 designation means this strain contains the lambda DE3 lysogen which carries the gene for T7 RNA polymerase under the control of the *lac*UV5 promoter. IPTG is required to induce expression of the T7 RNA polymerase.

The strain is an *E. coli* B/r strain and does not contain the *lon* protease. It also has a mutation in the outer membrane protease, OmpT. The lack of these two key proteases reduces degradation of heterologous proteins expressed in the strain.

The strain carries a mutated *rne* gene (*rne131*) which encodes a truncated RNase E enzyme that lacks the ability to degrade mRNA, resulting in an increase in mRNA stability (see page 6).

### **Accessory Products**

Introduction	The products listed in this section are intended for use with the Champion <sup>™</sup> pET Directional TOPO <sup>®</sup> Expression Kits. For more information, refer to our Web site (www.invitrogen.com) or call Technical Service (see page 56).			
Additional Products	Many of the reagents supplied in the Champion <sup>™</sup> pET Directional TOPO <sup>®</sup> Expression Kits and other reagents suitable for use with the kits are available separately from Invitrogen. Ordering information for these reagents is provided below.			
	Item	Quantity	Catalog no.	
	One Shot <sup>®</sup> TOP10 Chemically Competent	10 x 50 μl	C4040-10	
	E. coli	20 x 50 μl	C4040-03	
	BL21 Star <sup>™</sup> (DE3) One Shot <sup>®</sup> Chemically Competent <i>E. coli</i>	20 x 50 μl	C6010-03	
	BL21 Star <sup>™</sup> (DE3)pLysS One Shot <sup>®</sup> Chemically Competent <i>E. coli</i>	20 x 50 μl	C6020-03	
	BL21-AI <sup>™</sup> One Shot <sup>®</sup> Chemically Competent <i>E. coli</i>	20 x 50 μl	C6070-03	
	PureLink <sup>™</sup> HQ Mini Plasmid Purification Kit	100 reactions	K2100-01	
	PureLink <sup>™</sup> Quick Gel Extraction Kit	50 reactions	K2100-12	
	Ampicillin	200 mg	11593-027	
	Kanamycin Sulfate	5 g	11815-024	
		25 g	11815-032	
	Carbenicillin	5 g	10177-012	
	Isopropylthio-β-galactoside (IPTG)	1 g	15529-019	
	EKMax™	250 units	E180-01	
		1000 units	E180-02	
	AcTEV Protease	1000 units	12575-015	

#### **Accessory Products, continued**

Products to Detect	Expression of your recombinant fusion protein can be detected using an
Recombinant	antibody to the appropriate epitope. The table below describes the products
Proteins	available from Invitrogen for detection of fusion proteins expressed using the
	appropriate pET TOPO <sup>®</sup> vector (see pages 11-15 for details about the N- and/or
	C-terminal tags present on each pET TOPO <sup>®</sup> vector).

Product	Mechanism of Detection	Catalog no.
Anti-Xpress <sup>™</sup> Antibody	Detects 8 amino acid Xpress™	R910-25
Anti-Xpress <sup>™</sup> -HRP Antibody	epitope:	R911-25
	DLYDDDDK	
Anti-HisG Antibody	Detects the N-terminal polyhistidine (6xHis) tag	R940-25
Anti-HisG-HRP Antibody		R941-25
Anti-HisG-AP Antibody	НННННG	R942-25
Anti-V5 Antibody	Detects 14 amino acid epitope	R960-25
Anti-V5-HRP Antibody	derived from the P and V proteins of the paramyxovirus, SV5 (Southern <i>et al.</i> , 1991):	R961-25
Anti-V5-AP Antibody		R962-25
	GKPIPNPLLGLDST	
Anti-His (C-term) Antibody	Detects the C-terminal	R930-25
Anti-His(C-term)-HRP Antibody	polyhistidine (6xHis) tag (requires the free carboxyl group	R931-25
Anti-His(C-term)-AP Antibody	1997): НННННН-СООН	R932-25
Anti-Thio <sup>™</sup> Antibody	Detects His-Patch thioredoxin fusion proteins	R920-25
	<b>Note:</b> The exact epitope detected by this antibody has not been mapped	

The amount of antibody supplied is sufficient for 25 western blots.

#### **Accessory Products, continued**

#### Products to Purify Recombinant Fusion Proteins

If you clone your gene of interest in frame with a C-terminal or N-terminal peptide containing a polyhistidine (6xHis) tag, you may use Invitrogen's ProBond<sup>™</sup> or Ni-NTA resins to purify your recombinant fusion protein. See the table below for ordering information.

Product	Quantity	Catalog no.
ProBond <sup>™</sup> Nickel-Chelating Resin	50 ml	R801-01
	150 ml	R801-15
ProBond <sup>™</sup> Purification System	6 purifications	K850-01
ProBond <sup>™</sup> Purification System with	1 kit	K851-01
Anti-Xpress <sup>™</sup> -HRP Antibody		
ProBond <sup>™</sup> Purification System with	1 kit	K853-01
Anti-His(C-term)-HRP Antibody		
ProBond <sup>™</sup> Purification System with Anti-V5-HRP Antibody	1 kit	K854-01
Ni-NTA Agarose	10 ml	R901-01
	25 ml	R901-15
	100 ml	R901-10
Ni-NTA Purification System	6 purifications	K950-01
Ni-NTA Purification System with Anti-Xpress <sup>™</sup> -HRP Antibody	1 kit	K951-01
Ni-NTA Purification System with	1 kit	K953-01
Anti-His(C-term)-HRP Antibody		
Ni-NTA Purification System with Anti-V5-HRP Antibody	1 kit	K954-01
Polypropylene Columns (empty)	50	R640-50

#### Introduction

#### **Overview**

## **Introduction** The Champion<sup>™</sup> pET Directional TOPO<sup>®</sup> Expression Kits utilize a highly efficient, 5-minute cloning strategy ("TOPO<sup>®</sup> Cloning") to directionally clone a blunt-end PCR product into a vector for high-level, T7-regulated expression in *E. coli*. Blunt-

ligase, post-PCR procedures, or restriction enzymes required. Depending on the vector chosen, the pET TOPO<sup>®</sup> vectors are available with:

end PCR products clone directionally at greater than 90% efficiency, with no

- N-terminal or C-terminal peptide tags for production of recombinant fusion proteins that may be easily detected or purified
- Protease recognition site for removal of the N-terminal peptide tag from your recombinant fusion protein
- Antibiotic resistance marker for selection of transformants

See the table below for a list of the available pET TOPO<sup>®</sup> vectors and the fusion tag, cleavage site, and selection marker for each vector.

pET TOPO® vector	Fusion Peptide	Fusion Tag	Cleavage Site	Selection Marker
pET100/D-TOPO®	N-terminal	Xpress <sup>™</sup> , 6xHis	EK	Ampicillin
pET200/D-TOPO®				Kanamycin
pET101/D-TOPO®	C-terminal	V5, 6xHis		Ampicillin
pET102/D-TOPO®	N-terminal	His-Patch thioredoxin	EK	Ampicillin
	C-terminal	V5, 6xHis		
pET151/D-TOPO®	N-terminal	V5, 6xHis	TEV protease	Ampicillin

EK = enterokinase; TEV = tobacco etch virus

#### The Champion<sup>™</sup> pET Expression System

The Champion<sup>™</sup> pET Expression System is based on expression vectors originally developed by Studier and colleagues, and takes advantage of the high activity and specificity of the bacteriophage T7 RNA polymerase to allow regulated expression of heterologous genes in *E. coli* from the T7 promoter (Rosenberg *et al.*, 1987; Studier & Moffatt, 1986; Studier *et al.*, 1990). For more information about the Champion<sup>™</sup> pET Expression System, see page 4.

#### **Overview**, continued

Features of the Champion<sup>™</sup> pET Directional TOPO<sup>®</sup> Vectors The pET TOPO<sup>®</sup> vectors are designed to facilitate rapid, directional TOPO<sup>®</sup> Cloning of blunt-end PCR products for regulated expression in *E. coli*. Features of the vectors include:

- T7*lac* promoter for high-level, IPTG-inducible expression of the gene of interest in *E. coli* (Dubendorff & Studier, 1991; Studier *et al.*, 1990)
- Directional TOPO<sup>®</sup> Cloning site for rapid and efficient directional cloning of blunt-end PCR products (see page 3 for more information)
- N- or C-terminal fusion tags for detection and purification of recombinant fusion proteins (choice of tag depends on the particular vector; see the previous page)
- Protease recognition site for cleavage of the fusion tag from the recombinant protein of interest (present on N-terminal fusion vectors)
- N-terminal His-Patch thioredoxin for increased translation efficiency and solubility of heterologous proteins (pET102/D-TOPO<sup>®</sup> only)
- *lacI* gene encoding the lac repressor to reduce basal transcription from the T7*lac* promoter in the pET TOPO<sup>®</sup> vector and from the *lacUV5* promoter in the *E. coli* host chromosome (see page 4 for more information)
- Antibiotic resistance marker for selection in E. coli
- pBR322 origin for low-copy replication and maintenance in E. coli

## How Directional TOPO<sup>®</sup> Cloning Works

How Topoisomerase I Works	Topoisomerase I from <i>Vaccinia</i> virus binds to duplex DNA at specific sites and cleaves the phosphodiester backbone after 5'-CCCTT in one strand (Shuman, 1991). The energy from the broken phosphodiester backbone is conserved by formation of a covalent bond between the 3' phosphate of the cleaved strand and a tyrosyl residue (Tyr-274) of topoisomerase I. The phospho-tyrosyl bond between the DNA and enzyme can subsequently be attacked by the 5' hydroxyl of the original cleaved strand, reversing the reaction and releasing topoisomerase (Shuman, 1994). TOPO <sup>®</sup> Cloning exploits this reaction to efficiently clone PCR products.		
Directional TOPO <sup>®</sup> Cloning	Directional joining of double-strand DNA using TOPO <sup>®</sup> -charged oligonucleotides occurs by adding a 3' single-stranded end (overhang) to the incoming DNA (Cheng & Shuman, 2000). This single-stranded overhang is identical to the 5' end of the TOPO <sup>®</sup> -charged DNA fragment. At Invitrogen, this idea has been modified by adding a 4 nucleotide overhang sequence to the TOPO <sup>®</sup> -charged DNA and adapting it to a 'whole vector' format.		
	In this system, PCR products are directionally cloned by adding four bases to the forward primer (CACC). The overhang in the cloning vector (GTGG) invades the 5' end of the PCR product, anneals to the added bases, and stabilizes the PCR product in the correct orientation. Inserts can be cloned in the correct orientation with efficiencies equal to or greater than 90%.		
	Topoisomerase		
	Tyr-274 CACC ATG NNN NNN AAG GG GTGG TAC NNN NNN TTC CC		
	Overhang		
	Overhang invades double-stranded DNA, displacing the bottom strand.		
	Topoisomerase		
	CCCTTCACC ATG NNN NNN AAG GG GGGAAGTGG TAC NNN NNN TTC CC		

### **T7-Regulated Expression**

The Basis of T7- Regulated Expression	The Champion <sup>™</sup> pET Expression System uses elements from bacteriophage T7 to control expression of heterologous genes in <i>E. coli</i> . In the pET TOPO <sup>®</sup> vectors, expression of the gene of interest is controlled by a strong bacteriophage T7 promoter that has been modified to contain a <i>lac</i> operator sequence (see below). In bacteriophage T7, the T7 promoter drives expression of gene 10 ( $\phi$ 10). T7 RNA polymerase specifically recognizes this promoter. To express the gene of interest, it is necessary to deliver T7 RNA polymerase to the cells by inducing expression of the polymerase or infecting the cell with phage expressing the polymerase. In the Champion <sup>™</sup> pET Directional TOPO <sup>®</sup> Expression System, T7 RNA polymerase is supplied by the BL21 Star <sup>™</sup> (DE3) host <i>E. coli</i> strain in a regulated manner (see below). When sufficient T7 RNA polymerase is produced, it binds to the T7 promoter and transcribes the gene of interest.
Regulating Expression of T7 RNA Polymerase	The BL21 Star <sup>™</sup> (DE3) <i>E. coli</i> strain is specifically included in each Champion <sup>™</sup> pET Directional TOPO <sup>®</sup> Expression kit for expression of T7-regulated genes. This strain carries the DE3 bacteriophage lambda lysogen. This λDE3 lysogen contains a <i>lac</i> construct consisting of the following elements:
	the <i>lacI</i> gene encoding the lac repressor
	• the T7 RNA polymerase gene under control of the <i>lacUV5</i> promoter
	• a small portion of the <i>lacZ</i> gene.
	This <i>lac</i> construct is inserted into the <i>int</i> gene such that it inactivates the <i>int</i> gene. Disruption of the <i>int</i> gene prevents excision of the phage ( <i>i.e.</i> lysis) in the absence of helper phage. The <i>lac</i> repressor (encoded by <i>lac1</i> ) represses expression of T7 RNA polymerase. Addition of the gratuitous inducer, isopropyl β-D-thiogalacto- side (IPTG) allows expression of T7 RNA polymerase from the <i>lacUV5</i> promoter.
	The BL21 Star <sup>™</sup> (DE3) strain also contains other features which facilitate high-level expression of heterologous genes. For more information, see page 6.
T7 <i>lac</i> Promoter	Studies have shown that there is always some basal expression of T7 RNA polymerase from the <i>lacUV5</i> promoter in $\lambda$ DE3 lysogens even in the absence of inducer (Studier & Moffatt, 1986). In general, this is not a problem, but if the gene of interest is toxic to the <i>E. coli</i> host, basal expression of the gene of interest may lead to plasmid instability and/or cell death.
	To address this problem, the pET TOPO <sup>®</sup> vectors have been designed to contain a T7 <i>lac</i> promoter to drive expression of the gene of interest. The T7 <i>lac</i> promoter consists of a <i>lac</i> operator sequence placed downstream of the T7 promoter. The <i>lac</i> operator serves as a binding site for the lac repressor (encoded by the <i>lacI</i> gene) and functions to further repress T7 RNA polymerase-induced basal transcription of the gene of interest in BL21 Star <sup>™</sup> (DE3) cells.

## T7-Regulated Expression, continued

Expressing Toxic Genes	In some cases, the gene of interest is so toxic to BL21 Star <sup><math>TM</math></sup> (DE3) cells that other <i>E. coli</i> host strains may be required for expression. For a discussion of other alternative strains that may be used, see page 6.
Using TOP10 Cells	One Shot <sup>®</sup> TOP10 competent <i>E. coli</i> , which do not contain T7 RNA polymerase, are included in each Champion <sup>™</sup> pET Directional TOPO <sup>®</sup> Expression kit to provide a host for stable propagation and maintenance of recombinant plasmids. As mentioned on the previous page, the presence of T7 RNA polymerase, even at basal levels, can lead to expression of the desired gene even in the absence of inducer. If the gene of interest is toxic to the <i>E. coli</i> host, plasmid instability and/or cell death may result. We recommend that you transform your TOPO <sup>®</sup> Cloning reaction into TOP10 cells for characterization of the construct, propagation, and maintenance. When you are ready to perform an expression experiment, transform your construct into BL21 Star <sup>™</sup> (DE3) <i>E. coli</i> .

## BL21 Star<sup>™</sup> *E. coli* Strains

BL21 Star <sup>™</sup> Strains	The BL21 Star <sup>TM</sup> (DE3) <i>E. coli</i> strain is included in each Champion <sup>TM</sup> pET Directional TOPO <sup>®</sup> Expression Kit for use as a host for expression. Other BL21 Star <sup>TM</sup> strains are also available from Invitrogen (see below). In addition to the $\lambda$ DE3 lysogen which allows high-level expression of T7-regulated genes (see page 3), the BL21 Star <sup>TM</sup> strains also contain the <i>rne131</i> mutation. This particular mutation further enhances the expression capabilities of BL21 Star <sup>TM</sup> .
<i>rne131</i> Mutation	The <i>rne</i> gene encodes the RNase E enzyme, an essential, 1061 amino acid <i>E. coli</i> endonuclease which is involved in rRNA maturation and mRNA degradation as a component of a protein complex known as a "degradosome" (Grunberg-Manago, 1999; Lopez <i>et al.</i> , 1999). Various studies have shown that the N-terminal portion of RNase E (approximately 584 amino acids) is required for rRNA processing and cell growth while the C-terminal portion of the enzyme (approximately 477 amino acids) is required for mRNA degradation (Kido <i>et al.</i> , 1996; Lopez <i>et al.</i> , 1999). The <i>rne131</i> mutation (present in the BL21 Star <sup>™</sup> strains) encodes a truncated RNase E which lacks the C-terminal 477 amino acids of the enzyme required for mRNA degradation (Kido <i>et al.</i> , 1996; Lopez <i>et al.</i> , 1999). Thus, mRNAs expressed in the RNase E-defective BL21 Star <sup>™</sup> strains exhibit increased stability when compared to other BL21 strains. When heterologous genes are expressed in the BL21 Star <sup>™</sup> strains from T7-based expression vectors, the yields of recombinant proteins generally increase.
BL21 Star <sup>™</sup> (DE3)pLysS Strain	If you discover that your gene is toxic to BL21 Star <sup>™</sup> (DE3) cells, you may want to perform your expression experiments in the BL21 Star <sup>™</sup> (DE3)pLysS strain (see page x for ordering information). The BL21 Star <sup>™</sup> (DE3)pLysS strain contains the pLysS plasmid, which produces T7 lysozyme. T7 lysozyme binds to T7 RNA polymerase and inhibits transcription. This activity results in reduced basal levels of T7 RNA polymerase, leading to reduced basal expression of T7-driven heterologous genes. For more information about BL21 Star <sup>™</sup> (DE3)pLysS, refer to our Web site (www.invitrogen.com) or call Technical Service (see page 56).
Note	Note that while BL21 Star <sup>™</sup> (DE3)pLysS reduces basal expression from the gene of interest when compared to BL21 Star <sup>™</sup> (DE3), it also generally reduces the overall induced level of expression of recombinant protein.

#### Thioredoxin

Introduction	The pET102/D-TOPO <sup>®</sup> vector allows you to clone your gene of interest as a fusion to a mutated thioredoxin protein (His-Patch thioredoxin). For more information about thioredoxin and His-Patch thioredoxin, see below. The 11.7 kDa thioredoxin protein is found in yeast, plants, and mammals, as well as in bacteria. It was originally isolated from <i>E. coli</i> as a hydrogen donor for ribonuclease reductase (see Holmgren, 1985 for a review). The gene has been completely sequenced (Wallace & Kushner, 1984). The protein has been crystallized and its three-dimensional structure determined (Katti <i>et al.</i> , 1990). When overexpressed in <i>E. coli</i> , thioredoxin is able to accumulate to approximately 40% of the total cellular protein and still remain soluble. When used as a fusion partner, thioredoxin can increase translation efficiency, and in some cases, solubility, of eukaryotic proteins expressed in <i>E. coli</i> .			
Thioredoxin				
	Examples of eukaryotic proteins that have been produced as soluble C-terminal fusions to the thioredoxin protein in <i>E. coli</i> (LaVallie <i>et al.</i> , 1993) include:			
	• Murine interleukins 2, 4, and 5 and human interleukin-3			
	Human macrophage colony stimulating factor			
	Murine steel factor			
	Murine leukemia inhibitory factor			
	Human bone morphogenetic protein-2			
His-Patch Thioredoxin	The thioredoxin protein in pET102/D-TOPO <sup>®</sup> has been mutated to contain a metal binding domain, and is termed "His-Patch thioredoxin". To create a metal binding domain in the thioredoxin protein, the glutamate residues at position 32 and the glutamine residue at position 64 were mutated to histidine residues. When His-Patch thioredoxin folds, the histidines at positions 32 and 64 interact with a native histidine at position 8 to form a "patch". This histidine patch has been shown to have high affinity for divalent cations (Lu <i>et al.</i> , 1996). His-Patch thioredoxin) proteins can therefore be purified on metal chelating resins ( <i>e.g.</i> ProBond <sup>™</sup> or Ni-NTA).			

#### **Experimental Outline**



### **Designing PCR Primers**

#### **Basic Requirements**

Designing Your PCR Primers	The design of the PCR primers to amplify your gene of interest is critical for expression. Depending on the pET TOPO <sup>®</sup> vector you are using, consider the following when designing your PCR primers:		
	Sequences required to facilitate directional cloning (see below)		
	• Whether or not you wish to clone your PCR product in frame with the appropriate N-terminal and/or C-terminal peptide tag (see pages 11-15 for information about each pET TOPO <sup>®</sup> vector)		
General Requirements for the Forward	To enable directional cloning, the forward PCR primer <b>must</b> contain the sequence, CACC, at the 5' end of the primer. The 4 nucleotides, CACC, base pair with the overhang sequence, GTGG, in each pET TOPO <sup>®</sup> vector.		
Primer	For example, below is the DNA sequence of the N-terminus of a theoretical protein and the proposed sequence for your forward PCR primer:		
	DNA sequence: 5'-ATG GGA TCT GAT AAA Proposed Forward PCR primer: 5'-C ACC ATG GGA TCT GAT AAA		
	See pages 11-15 for other factors to consider when designing the forward primer.		
General Requirements for the Reverse Primer	In general, design the reverse PCR primer to allow you to clone your PCR product in frame with any C-terminal tag, if desired. To ensure that your PCR product clones directionally with high efficiency, the reverse PCR primer MUST NOT be complementary to the overhang sequence GTGG at the 5' end. A one base pair mismatch can reduce the directional cloning efficiency from 90% to 75%, and may increase the chances of your ORF cloning in the opposite orientation. We have not observed evidence of PCR products cloning in the opposite orientation from a two base pair mismatch, but this has not been tested thoroughly.		
	<b>Example:</b> Below is the sequence of the C-terminus of a theoretical protein. You want to clone in frame with the C-terminal tag. The stop codon is underlined.		
	DNA sequence: AAG TCG GAG CAC TCG ACG ACG GTG <u>TAG</u> -3'		
	One solution is to design the reverse PCR primer to start with the codon just up- stream of the stop codon, but the last two codons contain GTGG (underlined below), which is identical to the overhang sequence. As a result, the reverse primer will be complementary to the overhang sequence, increasing the probability that the PCR product will clone in the opposite orientation. You want to avoid this situation.		
	DNA sequence: AAG TCG GAG CAC TCG ACG AC <u>G GTG</u> TAG-3' Proposed Reverse PCR primer sequence: TG AGC TGC TGC CAC-5'		
	Another solution is to design the reverse primer so that it hybridizes just down- stream of the stop codon, but still includes the C-terminus of the ORF. Note that you will need to replace the stop codon with a codon for an innocuous amino acid such as glycine or alanine.		

### **Basic Requirements, continued**

Important	•	Remember that the pET TOPO <sup>®</sup> vectors accept blunt-end PCR products. Refer to pages 11-15 for a discussion of specific factors to consider when designing PCR primers for cloning into each pET TOPO <sup>®</sup> vector. Do not add 5 <sup>′</sup> phosphates to your primers for PCR. This will prevent ligation into the pET TOPO <sup>®</sup> vectors.
	•	We recommend gel-purifying your oligonucleotides, especially if they are long (> 30 nucleotides).
Example of Primer Design	Th on int end	e example below uses a theoretical protein and is for illustration purposes ly. In this case, PCR primers are designed to allow cloning of the PCR product o pET101/D-TOPO <sup>®</sup> . In this example, the N-terminus of the protein is coded by:
		5'-ATGGCCCCCCGACCGATGTCAGCCTGGGGGACGAA
	1.	Design the forward PCR primer to be:
		5'-CACC <u>ATG</u> GCCCCCCGACCGAT-3'
	2.	For the reverse primer, analyze the C-terminus of the protein. The stop codon is underlined (see the top strand below).
		GCG GTT AAG TCG GAG CAC TCG ACG ACT GCA <u>TAG</u> -3' CGC CAA <u>TTC AGC CTC GTG AGC TGC TGA CGT</u> ATC-5'
	3.	To fuse the ORF in frame with the V5 epitope and 6xHis tag, remove the stop codon by starting with nucleotides homologous to the last codon (TGC) and continue upstream (underlined sequence in the bottom strand above). The reverse primer will be:
		5'-TGC AGT CGT CGA GTG CTC CGA CTT-3'
	4.	This will amplify the C-terminus without the stop codon and allow you to clone the ORF in frame with the V5 epitope and 6xHis tag.
		If you don't want the V5 epitope and 6xHis tag, simply begin with the stop codon:

5'-<u>CTA</u> TGC AGT CGT CGA GTG CTC CGA CTT-3'

## Specific Requirements for Cloning into pET100/D-TOPO<sup>®</sup> and pET200/D-TOPO<sup>®</sup>

Introduction	pET100/D-TOPO <sup>®</sup> and pET200/D-TOPO <sup>®</sup> allow expression of recombinant protein with an N-terminal tag containing the Xpress <sup>™</sup> epitope and a 6xHis tag. The N-terminal tag also includes an enterokinase (EK) recognition site to enable removal of the tag after protein purification using enterokinase ( <i>e.g.</i> EKMax <sup>™</sup> ).
Additional Cloning Considerations	In addition to the guidelines on page 9, consider the following when designing PCR primers to clone your DNA into pET100/D-TOPO <sup>®</sup> or pET200/D-TOPO <sup>®</sup> .
	Be sure to include a stop codon in the reverse primer or design the reverse primer to hybridize downstream of the native stop codon.

If you wish to	Then
include the Xpress <sup>™</sup> epitope and 6xHis tag	design the forward PCR primer to place the gene of interest in frame with the N-terminal tag. Note that:
	• a ribosome binding site (RBS) is included upstream of the initiation ATG in the N-terminal tag to ensure optimal spacing for proper translation
	• at least five nonnative amino acids will be present between the EK cleavage site and the start of your gene
express your protein with	design the forward PCR primer to include the following:
a native N-terminus, <i>i.e.</i>	a stop codon to terminate the N-terminal peptide
without the N-terminal peptide	• a second ribosome binding site (AGGAGG) 9-10 base pairs 5' of the initiation ATG codon of your protein

Note: The first three base pairs of the PCR product following the 5' CACC overhang will constitute a complete codon.

#### TOPO<sup>®</sup> Cloning Site of pET100/D-TOPO<sup>®</sup> and pET200/D-TOPO<sup>®</sup>

Use the diagram below to help you design suitable PCR primers to clone your PCR product into pET100/D-TOPO<sup>®</sup> or pET200/D-TOPO<sup>®</sup>. Restriction sites are labeled to indicate the actual cleavage site. **The sequence of each vector is available for downloading from our Web site or from Technical Service (see page 56).** For more information about pET100/D-TOPO<sup>®</sup> or pET200/D-TOPO<sup>®</sup>, see pages 42-43.

121	ATAGGCGCCA GCAACCGCAC CTGTGG	CGCC GGTGATGCCG GCCACGATGC	GTCCGGCGTA GAGGATCGAG ATCTCGATCC
	T7 promoter/priming si	te	
	T7 promoter	lac operator	
201	CGCGAAATTA ATACGACTCA CTATAG	GGGA ATTGTGAGCG GATAACAATT	CCCCTCTAGA AATAATTTTG TTTAACTTTA
	RBS Nde I	Polyhistidine region	Nhe I
281	AGAAGGAGAT ATACAT <b>ATG</b> CGG GG	T TCT CAT CAT CAT CAT CAT C	CAT GGT ATG GCT AGC ATG ACT GGT GGA
	Met Arg Gl	y Ser His His His His His H	His Gly Met Ala Ser Met Thr Gly Gly
		Xpress <sup>™</sup> epitope	
351	CAG CAA ATG GGT CGG GAT CTG	TAC GAC GAT GAC GAT AAG GAT	CAT CCC TT <u>C ACC</u> AAGGGC
	Gin Gin Met Giv Arg Asp Leu	Tyr Asn Asn Asn Asn Lys Asr	GGG AAG TGG
	Sin Sin net Siy nig hop lea		
		EK recognition site	avage site
			17 reverse priming site
411	GAGCTCAACG ATCCGGCTGC TAACAA	AGCC CGAAAGGAAG CTGAGTTGGC	TGCTGCCACC GCTGAGCAAT AACTAGCATA

#### Specific Requirements for Cloning into pET101/D-TOPO®

## Additional Cloning Considerations

pET101/D-TOPO<sup>®</sup> allows expression of recombinant protein with a native N-terminus and a C-terminal fusion tag. In addition to the guidelines on page 9, consider the following when designing PCR primers to clone your DNA into pET101/D-TOPO<sup>®</sup>.

For maximal expression of native protein, the forward PCR primer should be designed to place the initial ATG codon of the desired protein approximately 9 to 10 base pairs from the ribosome binding site (Gold, 1988; Miller, 1992). This will ensure the optimal spacing for proper translation.

If you wish to	Then
express your protein with a native N-terminus using the vector encoded ribosome binding site	design the forward PCR primer such that the initial ATG codon of your protein directly follows the 5' CACC overhang.
include the C-terminal V5 epitope and 6xHis tag	design the reverse PCR primer to remove the native stop codon in the gene of interest and preserve the reading frame through the C-terminal tag.
<b>not</b> include the C-terminal V5 epitope and 6xHis tag	include the native sequence containing the stop codon in the reverse primer or make sure the stop codon is upstream from the reverse PCR primer binding site.

**Note:** The first three base pairs of the PCR product following the 5' CACC overhang will constitute a complete codon.

#### TOPO<sup>®</sup> Cloning Site of pET101/D-TOPO<sup>®</sup>

O<sup>®</sup> Use the diagram below to help you design suitable PCR primers to clone your
 PO<sup>®</sup> PCR product into pET101/D-TOPO<sup>®</sup>. Restriction sites are labeled to indicate the actual cleavage site. The sequence of the vector is available for downloading from our Web site or from Technical Service (see page 56). For more information about pET101/D-TOPO<sup>®</sup>, see pages 44-45.

								Τ7	promoter/	priming sit	te
							T7	promoter			
151	GGTGATGCCG GCCACO	GATGC GTCCG	GCGTA GA	GGATCGAG	ATCTCGAT	CC CGCG	AAATTA	ATACGA	CTCA C	TATAG	GGGA
	lac operator					R	BS	RBS	1		
231	ATTGTGAGCG GATAAC	CAATT CCCCT	CTAGA AA'	TAATTTTG	TTTAACTT	TA AGAA	GGAATT	CAGGAG	CCCT I GGGA A	C ACC G TGC	ATG TAC
										C F	7
	Sac I	BstB I				V5 e	epitope				
306	Sacl AAG GGC GAG CTC A Lys Gly Glu Leu A	BstBI I AAT TCG AAG Asn Ser Lys	CTT GAA Leu Glu	GGT AAG Gly Lys	CCT ATC Pro Ile	V5 e CCT AAC Pro Asn	CCT CT Pro Le	C CTC u Leu	GGT CI Gly Le	C GAT u Asp	TCT Ser
306	Saci AAG GGC GAG CTC A Lys Gly Glu Leu A Agel	BstB     AAT TCG AAG Asn Ser Lys Polyhisti	CTT GAA Leu Glu dine region	GGT AAG Gly Lys	CCT ATC Pro Ile	V5 CCT AAC Pro Asn	CCT CT Pro Le	C CTC u Leu	GGT CI Gly Le	C GAT u Asp	TCT Ser
306 372	Saci AAG GGC GAG CTC A Lys Gly Glu Leu A Agel ACG CGT ACC GGT C Thr Arg Thr Gly F	BstB I I TCG AAG ASN Ser Lys Polyhisti CAT CAT CAC His His His	CTT GAA Leu Glu dine region CAT CAC His His	GGT AAG Gly Lys CAT TGA His ***	CCT ATC Pro Ile GTTTGA T	V5 CCT AAC Pro Asn	ppitope CCT CT Pro Le CT AACA	C CTC u Leu AAGCCC	GGT CI Gly Le GAAAG	C GAT u Asp GAAGC	TCT Ser
306 372	Saci AAG GGC GAG CTC A Lys Gly Glu Leu A Agel ACG CGT ACC GGT C Thr Arg Thr Gly F	BstB I AAT TCG AAG Asn Ser Lys Polyhisti CAT CAT CAC His His His T7 reverse p	CTT GAA Leu Glu dine region CAT CAC His His riming site	GGT AAG Gly Lys CAT TGA His ***	CCT ATC Pro Ile GTTTGA T	V5 CCT AAC Pro Asn	CCT CT Pro Le CT AACA	C CTC u Leu AAGCCC	GGT CI Gly Le	C GAT u Asp GAAGC	TCT Ser

#### Specific Requirements for Cloning into pET102/D-TOPO®

Specific Features for Expression in pET102/D-TOPO<sup>®</sup> pET102/D-TOPO<sup>®</sup> is designed with some specific features to facilitate expression. They are:

- The initiation ATG is correctly spaced from the optimized ribosome binding site (RBS) to ensure optimal translation
- HP-thioredoxin acts to increase translation efficiency and in some cases, solubility
- HP-thioredoxin can be removed after protein purification using enterokinase (*e.g.* EKMax<sup>™</sup>, Catalog no. E180-01).

Additional Cloning Considerations

In addition to the guidelines on page 9, you should consider the following when designing PCR primers to clone your DNA into pET102/D-TOPO<sup>®</sup>. Use the diagram on the next page to help you design your PCR primers.

If you wish to	Then
clone in frame with thioredoxin	design the forward PCR primer to ensure that your protein is in frame with the N-terminal leader peptide.
include the C-terminal V5 epitope and 6xHis tag	design the reverse PCR primer to remove the native stop codon in the gene of interest and preserve the reading frame through the C-terminal tag.
<b>not</b> include the C-terminal V5 epitope and 6xHis tag	include the native sequence containing the stop codon in the reverse primer or make sure the stop codon is upstream from the reverse PCR primer binding site.

**Note:** The first three base pairs of the PCR product following the 5' CACC overhang will constitute a complete codon.

## Specific Requirements for Cloning into pET102/D-TOPO<sup>®</sup>, continued

#### **TOPO<sup>®</sup> Cloning Site** of pET102/D-TOPO<sup>®</sup> Use the diagram below to help you design appropriate PCR primers to clone your PCR product into pET102/D-TOPO<sup>®</sup>. Restriction sites are labeled to indicate the actual cleavage site. The complete sequence of the vector is available for downloading from our Web site or from Technical Service (see page 56). For more information about pET102/D-TOPO<sup>®</sup>, refer to the Appendix, pages 46-47.

																-		Т	7 prom	oter/prii	ming sit	9
		~~ ~~ ~			~~~~	~~ ~					~~~~~	3 00 01					T7 pr	romoter				
151	GGT	5A'I'G(	CG (	JCCA	CGAT	GC G	rccg	GCG17	A GA	3GA'I'(	CGAG	ATC	I'CGA'	rcc (	CGCG.	AAA'I''	l'A A'	TACG	ACTC	A CTA	ATAGO	JGGA
			<i>lac</i> ope	erator			_								R	BS						
231	ATTO	GTGA	GCG (	GATA	ACAA	TT C	CCCT	CTAGA	A AA'	FAAT	FTTG	TTT	AACT	TTA J	AGAA	GGAG	AT A	TACA	TA A	<b>FG</b> G	GA TO	CT GAT
										His-pat	ch (HP)	thiored	loxin						1416	9C G.	LY SE	er Asp
310	AAA	ATT	ATT	CAT	CTG	ACT	GAT	GAT	TCT	TTT	GAT	ACT	GAT	GTA	CTT	AAG	GCA	GAT	GGT	GCA	ATC	CTG
	Lys	Ile	Ile	His	Leu	Thr	Asp	Asp	Ser	Phe	Asp	Thr	Asp	Val	Leu	Lys	Ala	Asp	Gly	Ala	Ile	Leu
276		<b>C N H</b>	mma	mag	007	010	mag	mag	COT	000	mag		3.00	3.00	000	000	200	OT C	C 3 0	C 7 7	380	
370	Val	Asp	Phe	Trp	Ala	His	Trp	Cys	Gly	Pro	Cys	Lys	Met	Ile	Ala	Pro	Ile	Leu	Asp	GAA Glu	Ile	Ala
442	GAC	GAA	TAT	CAG	GGC	AAA	CTG	ACC	GTT Val	GCA	AAA	CTG	AAC	ATC	GAT	CAC	AAC	CCG	GGC	ACT	GCG	CCG
	лэр	Gru	тут	GTH	GTĀ	цүз	цец	TIIT	vai	лıа	цүр	цец	ASII	TTC	лэр	1113	ASII	FIO	GTĂ	TIIT	ліа	FIO
508	AAA	TAT	GGC	ATC	CGT	GGT	ATC	CCG	ACT	CTG	CTG	CTG	TTC	AAA	AAC	GGT	GAA	GTG	GCG	GCA	ACC	AAA
	Lys	Tyr	Gly	Ile	Arg	Gly	Ile	Pro	Thr	Leu	Leu	Leu	Phe TrxF	Lys us forw	Asn ard prin	Gly ning site	Glu	Val	Ala	Ala	Thr	Lys
574	GTG	GGT	GCA	CTG	TCT		GGT	CAG	TTG	ΔΔΔ	GAG	TTC	СТС	GAC	GCT	AAC	СТС	GCC	I GGC	TCT	GGA	TCC
0/1	Val	Gly	Ala	Leu	Ser	Lys	Gly	Gln	Leu	Lys	Glu	Phe	Leu	Asp	Ala	Asn	Leu	Ala	Gly	Ser	Gly	Ser
		Enter	okinase	(EK) re	cognitio	on site	EK cle	eavage	site										Sac I	<i>Hin</i> d	111	
640	GGT	GAT	GAC	GAT	GAC	AAG	CTG	GGA	ATT	GAT	CCC	TTC	ACC			AAG	GGC	GAG	CTC	AAG	CTT	GAA
	Gly	Asp	Asp	Asp	Asp	Lys	Leu	Gly	Ile	Asp	Pro	Phe	Thr	\$ \$	•••	Lys	Gly	Glu	Leu	Lys	Leu	Glu
						V	5 epitop	e					G			Aae I		Po	olyhistid	ine (6xl	His) regi	on
700	GGT	AAG	CCT	ATC	CCT	AAC	CCT	CTC	CTC	GGT	CTC	GAT	TCT	ACG	1 CGT	ACC	GGT	CAT	CAT	CAC	CAT	CAC
	Gly	Lys	Pro	Ile	Pro	Asn	Pro	Leu	Leu	Gly	Leu	Asp	Ser	Thr	Arg	Thr	Gly	His	His	His	His	His
		1														г		T7 rev	verse p	riming s	ite	_
766	CAT His	TGA ***	GTT	rgat(	CC G	GCTG	CTAA	C AAA	AGCC	CGAA	AGG	AAGC	IGA (	GTTG	GCTG	CT G	CCAC	CGCT	g Ag	CAAT	AACT	AGCA

#### Specific Requirements for Cloning into pET151/D-TOPO®

Introduction	pET151/D-TOPO <sup>®</sup> allows expression of recombinant protein with an N-terminal tag containing the V5 epitope and a 6xHis tag. The N-terminal tag also includes a TEV protease cleavage site to enable removal of the tag after protein purification using TEV protease.

Additional Cloning<br/>ConsiderationsIn addition to the guidelines on page 9, consider the following when designing<br/>PCR primers to clone your DNA into pET151/D-TOPO®.

Be sure to include a stop codon in the reverse primer or design the reverse primer to hybridize downstream of the native stop codon.

If you wish to	Then
include the V5 epitope and 6xHis tag	design the forward PCR primer to place the gene of interest in frame with the N-terminal tag. Note that:
	a ribosome binding site (RBS) is included upstream of the initiation ATG in the N-terminal tag to ensure optimal spacing for proper translation
	at least six nonnative amino acids will be present between the TEV cleavage site and the start of your gene
express your protein with a native N-terminus, <i>i.e.</i> without the	design the forward PCR primer to include the following: a stop codon to terminate the N-terminal peptide
N-terminal peptide	a second ribosome binding site (AGGAGG) 9-10 base pairs 5' of the initiation ATG codon of your protein

**Note:** The first three base pairs of the PCR product following the 5' CACC overhang will constitute a complete codon.

#### TOPO<sup>®</sup> Cloning Site of pET151/D-TOPO<sup>®</sup>

Use the diagram below to help you design suitable PCR primers to clone your PCR product into pET151/D-TOPO<sup>®</sup>. Restriction sites are labeled to indicate the actual cleavage site. **The sequence of pET151/D-TOPO<sup>®</sup> is available for downloading from our Web site or from Technical Service (see page 56).** For more information about pET151/D-TOPO<sup>®</sup>, see pages 48-49.

GATCC
CTTTA
GGT CTC
Gly Leu
GA
£

## **Producing Blunt-End PCR Products**

Introduction	Once you have decided on a PCR strategy and have synthesized the primers, you are ready to produce your blunt-end PCR product using any thermostable, proofreading polymerase. Follow the guidelines below to produce your blunt-end PCR product.							
Materials Needed	You should have the following materials on hand before beginning.							
	Thermocycler and thermostable, proofreading polymerase							
	<ul> <li>10X PCR buffer appropriate for your polymerase</li> </ul>							
	<ul> <li>DNA template and primers for PCR product</li> </ul>							
Producing PCR	Set up a 25 $\mu$ l or 50 $\mu$ l PCR reaction using the guidelines below:							
Products	<ul> <li>Follow the instructions and recommendations provided by the manufacture of your thermostable, proofreading polymerase to produce blunt-end PCR products.</li> </ul>							
	• Use the cycling parameters suitable for your primers and template. Make sure to optimize PCR conditions to produce a single, discrete PCR product.							
	• Use a 7 to 30 minute final extension to ensure that all PCR products are completely extended.							
	• After cycling, place the tube on ice or store at -20°C for up to 2 weeks. Proceed to <b>Checking the PCR Product</b> , below.							
Checking the PCR Product	After you have produced your blunt-end PCR product, use agarose gel electrophoresis to verify the quality and quantity of your PCR product. Check for the following outcomes below.							
	• Be sure you have a single, discrete band of the correct size. If you do not have a single, discrete band, follow the manufacturer's recommendations for optimizing your PCR with the polymerase of your choice. Alternatively, you may gel-purify the desired product (see page 40s).							
	• Estimate the concentration of your PCR product. You will use this information when setting up your TOPO <sup>®</sup> Cloning reaction (see <b>Amount of PCR Product to Use in the TOPO<sup>®</sup> Cloning Reaction</b> , next page for details).							

#### **TOPO<sup>®</sup> Cloning Reaction and Transformation**

#### Setting Up the TOPO<sup>®</sup> Cloning Reaction

#### Introduction

Once you have produced the desired PCR product, you are ready to TOPO<sup>®</sup> Clone it into the pET TOPO<sup>®</sup> vector and transform the recombinant vector into One Shot<sup>®</sup> TOP10 *E. coli*. You should have everything you need set up and ready to use to ensure that you obtain the best possible results. We suggest that you read the this section and the section entitled **Transforming Competent Cells** before beginning. If this is the first time you have TOPO<sup>®</sup> Cloned, perform the control reactions on pages 37-39 in parallel with your samples.

#### Amount of PCR Product to Use in the TOPO<sup>®</sup> Cloning Reaction

When performing directional TOPO<sup>®</sup> Cloning, we have found that the molar ratio of PCR product:TOPO<sup>®</sup> vector used in the reaction is critical to its success. **To obtain the highest TOPO<sup>®</sup> Cloning efficiency, use a 0.5:1 to 2:1 molar ratio of PCR product:TOPO<sup>®</sup> vector (see figure below).** Note that the TOPO<sup>®</sup> Cloning efficiency decreases significantly if the ratio of PCR product: TOPO<sup>®</sup> vector is <0.1:1 or >5:1 (see figure below). These results are generally obtained if too little PCR product is used (*i.e.* PCR product is too dilute) or if too much PCR product is used in the TOPO<sup>®</sup> Cloning reaction. If you have quantitated the yield of your PCR product, you may need to adjust the concentration of your PCR product before proceeding to TOPO<sup>®</sup> Cloning.

**Tip:** For the pET TOPO<sup>®</sup> vectors, using 1-5 ng of a 1 kb PCR product or 5-10 ng of a 2 kb PCR product in a TOPO<sup>®</sup> Cloning reaction generally results in a suitable number of colonies.



#### Setting Up the TOPO<sup>®</sup> Cloning Reaction, continued

**Using Salt** You will perform TOPO<sup>®</sup> Cloning in a reaction buffer containing salt (*i.e.* using the Solution in the stock salt solution provided in the kit). Note that the amount of salt added to the **TOPO<sup>®</sup> Cloning** TOPO<sup>®</sup> Cloning reaction varies depending on whether you plan to transform chemically competent cells (provided) or electrocompetent cells (see page x for Reaction ordering information). If you are transforming chemically competent E. coli, use the stock Salt Solution as supplied and set up the TOPO<sup>®</sup> Cloning reaction as directed below. If you are transforming electrocompetent E. coli, the amount of salt in the TOPO<sup>®</sup> Cloning reaction **must be reduced** to 50 mM NaCl, 2.5 mM MgCl<sub>2</sub> to prevent arcing during electroporation. Dilute the stock Salt Solution 4-fold with water to prepare a 300 mM NaCl, 15 mM MgCl<sub>2</sub> Dilute Salt Solution. Use the Dilute Salt Solution to set up the TOPO® Cloning reaction as directed below.

## Performing the TOPO<sup>®</sup> Cloning Reaction

Use the procedure below to perform the TOPO<sup>®</sup> Cloning reaction. Set up the TOPO<sup>®</sup> Cloning reaction depending on whether you plan to transform chemically competent *E. coli* or electrocompetent *E. coli*. **Reminder:** For optimal results, be sure to use a 0.5:1 to 2:1 molar ratio of PCR product:TOPO<sup>®</sup> vector in your TOPO<sup>®</sup> Cloning reaction.

Note: The blue color of the  $\textsc{TOPO}^{\circledast}$  vector solution is normal and is used to visualize the solution.

Reagents*	Chemically Competent E. coli	Electrocompetent E. coli
Fresh PCR product	0.5 to 4 μl	0.5 to 4 µl
Salt Solution	1 µl	
Dilute Salt Solution (1:4)		1 μl
Sterile Water	add to a final volume of 5 $\mu$ l	add to a final volume of 5 $\mu$ l
TOPO <sup>®</sup> vector	1 μl	1 µl
Total Volume	6 µl	6 µl

\*Store all reagents at -20°C when finished. Salt solution and water can be stored at room temperature or +4°C.

1. Mix reaction gently and incubate for 5 minutes at room temperature (22-23°C).

**Note:** For most applications, 5 minutes will yield plenty of colonies for analysis. Depending on your needs, the length of the TOPO<sup>®</sup> Cloning reaction can be varied from 30 seconds to 30 minutes. For routine subcloning of PCR products, 30 seconds may be sufficient. For large PCR products (> 1 kb) or if you are TOPO<sup>®</sup> Cloning a pool of PCR products, increasing the reaction time may yield more colonies.

2. Place the reaction on ice and proceed to **Transforming One Shot**<sup>®</sup> **TOP10 Competent Cells**, next page.

Note: You may store the TOPO<sup>®</sup> Cloning reaction at -20°C overnight.

## Transforming One Shot<sup>®</sup> TOP10 Competent Cells

Introduction	Once you have performed the TOPO <sup>®</sup> Cloning reaction, you will transform your pET TOPO <sup>®</sup> construct into competent <i>E. coli</i> . One Shot <sup>®</sup> TOP10 Chemically Competent <i>E. coli</i> (Box 2) are included with the kit to facilitate transformation, however, you may also transform electrocompetent cells. Protocols to transform chemically competent or electrocompetent <i>E. coli</i> are provided in this section.
	To maintain the stability of your construct, we recommend that you transform your TOPO <sup>®</sup> Cloning reaction into TOP10 cells and characterize transformants in TOP10 before proceeding to expression studies using BL21 Star <sup>™</sup> (DE3). Expression of T7 RNA polymerase in BL21 Star <sup>™</sup> (DE3) may be leaky and may lead to rearrangement or loss of your plasmid.
Materials Supplied by the User	In addition to general microbiological supplies ( <i>i.e.</i> plates, spreaders), you will need the following reagents and equipment.
	• 42°C water bath (or electroporator with cuvettes, optional)
	• LB plates containing the appropriate antibiotic for selection (two for each transformation)
	• 37°C shaking and non-shaking incubator
Note	There is no blue-white screening for the presence of inserts. Most transformants will contain recombinant plasmids with the PCR product of interest cloned in the correct orientation, reducing the number of colonies to be analyzed. Sequencing primers are included in the kit to sequence across an insert in the multiple cloning site to confirm orientation and reading frame.
Preparing for Transformation	For each transformation, you will need one vial of competent cells and two selective plates.
	• Equilibrate a water bath to 42°C (for chemical transformation) or set up your electroporator if you are using electrocompetent <i>E. coli</i> .
	• Warm the vial of S.O.C. medium from Box 2 to room temperature.
	<ul> <li>Warm LB plates containing the appropriate antibiotic (<i>i.e.</i> 50-100 μg/ml ampicillin or 50-100 μg/ml kanamycin, as appropriate) at 37°C for 30 minutes</li> </ul>
	• Thaw <u>on ice</u> 1 vial of One Shot <sup>®</sup> TOP10 cells from Box 2 for each transformation.
	continued on next page

## Transforming One Shot<sup>®</sup> TOP10 Competent Cells, continued

<b>Q</b> Important	Th Or col	The number of colonies obtained after transforming the pET TOPO <sup>®</sup> vectors into One Shot <sup>®</sup> TOP10 cells is generally lower when compared to the number of colonies obtained after transforming other prokaryotic TOPO <sup>®</sup> vectors ( <i>e.g.</i> pCR <sup>®</sup> T7 TOPO <sup>®</sup> , pBAD/Thio-TOPO <sup>®</sup> ). This is due to the following:				
	•	Directional TOPO <sup>®</sup> Cloning generally yields 2 to 5-fold fewer colonies than traditional bidirectional TOPO TA Cloning <sup>®</sup>				
	•	Transforming low-copy number TOPO <sup>®</sup> plasmids generally yields 2 to 5- fold fewer colonies than transforming high-copy number TOPO <sup>®</sup> plasmids				
	If y Or ad	you have TOPO <sup>®</sup> Cloned previously, note that we have slightly modified the ne Shot <sup>®</sup> TOP10 transformation protocols (see below and the next page) to dress this issue. Briefly, we recommend the following:				
	•	Increase the amount of TOPO $^{\otimes}$ Cloning reaction that you transform into TOP10 cells (use 3 $\mu l)$ and				
	•	Increase the amount of transformed cells that you plate (use 100-200 $\mu$ l for chemically competent cells and 50-100 $\mu$ l for electrocompetent cells)				
	<b>Example:</b> When directionally TOPO <sup>®</sup> Cloning a 750 bp test insert into any of the pET TOPO <sup>®</sup> vectors, we generally obtain 500-1500 total colonies. Although fewer total colonies are obtained, greater than 90% of the colonies will contain plasmid with your PCR insert in the correct orientation.					
One Shot <sup>®</sup> TOP10 Chemical Transformation	1.	Add 3 μl of the TOPO <sup>®</sup> Cloning reaction from <b>Performing the TOPO<sup>®</sup></b> <b>Cloning Reaction</b> , Step 2, page 18 into a vial of One Shot <sup>®</sup> TOP10 Chemically Competent <i>E. coli</i> and mix gently. <b>Do not mix by pipetting up and down</b> .				
Protocol	2.	Incubate on ice for 5 to 30 minutes.				
		<b>Note:</b> Longer incubations on ice seem to have a minimal effect on transformation efficiency. The length of the incubation is at the user's discretion.				
	3.	Heat-shock the cells for 30 seconds at 42°C without shaking.				
	4.	Immediately transfer the tubes to ice.				
	5.	Add 250 1 of room temperature S.O.C. medium.				
	6.	Cap the tube tightly and shake the tube horizontally (200 rpm) at 37°C for 1 hour.				
	7.	Spread 100-200 l from each transformation on a prewarmed selective plate and incubate overnight at 37°C. We recommend plating two different volumes to ensure that at least one plate will have well-spaced colonies.				
	8.	An efficient TOPO <sup>®</sup> Cloning reaction may produce several hundred colonies. Pick ~5 colonies for analysis (see <b>Analyzing Positive Clones</b> , page 22).				
		<b>Note:</b> If you see few transformants, refer to the <b>Troubleshooting</b> section, page 34 for tips to optimize your TOPO <sup>®</sup> Cloning and transformation reactions.				

#### **Transforming Competent Cells, continued**

Transformation by Electroporation	Use ONLY electrocompetent cells for electroporation to avoid arcing. Do not use the One Shot <sup>®</sup> TOP10 chemically competent cells for electroporation.			
	1.	. Add 3 μl of the TOPO <sup>®</sup> Cloning reaction from Performing the TOPO <sup>®</sup> Cloning Reaction, Step 2, page 18 into 50 μl of electrocompetent <i>E. coli a</i> mix gently. Do not mix by pipetting up and down. Avoid formation of bubbles. Transfer the electrocompetent cells to a 0.1 cm cuvette.		
	2.	Electroporate your samples using your own protocol and an electroporator.		
		Note: If you have problems with arcing, see below.		
	3.	Immediately add 250 µl of room temperature S.O.C. medium.		
	4.	Transfer the solution to a 15 ml snap-cap tube (i.e. Falcon) and shake for at least 1 hour at 37°C to allow expression of the antibiotic resistance marker.		
	5.	Spread 50-100 $\mu$ l from each transformation on a prewarmed selective plate and incubate overnight at 37°C. We recommend plating two different volumes to ensure that at least one plate will have well-spaced colonies.		
	6.	An efficient TOPO <sup>®</sup> Cloning reaction may produce several hundred colonies. Pick ~5 colonies for analysis (see <b>Analyzing Positive Clones</b> , page 22).		
		<b>Note:</b> If you see few transformants, refer to the <b>Troubleshooting</b> section, page 34 for tips to optimize your TOPO <sup>®</sup> Cloning and transformation reactions.		
	To prevent arcing of your samples during electroporation, the volume of cells should be between 50 and 80 $\mu$ l (0.1 cm cuvettes) or 100 to 200 $\mu$ l (0.2 cm cuvettes).			
	If y su	you experience arcing during transformation, try one of the following ggestions:		

- Reduce the voltage normally used to charge your electroporator by 10%
- Reduce the pulse length by reducing the load resistance to 100 ohms
- Ethanol precipitate the TOPO<sup>®</sup> Cloning reaction and resuspend in water prior to electroporation

#### **Analyzing Transformants**

Analyzing Positive Clones	nalyzing Positive1.Pick 5 colonies and culture them overnight in LB or S.O.B. medialonescontaining the appropriate antibiotic.					
	<ol> <li>Isolate plasmid DNA using your method of choice. We recomm PureLink<sup>™</sup> HQ Mini Plasmid Purification Kit (Catalog no. K210</li> </ol>					
		<b>Note:</b> Since the pET TOPO <sup>®</sup> vectors are low-copy number plasmids, you may need t increase the amount of bacterial culture to obtain enough plasmid DNA for sequenc or analysis purposes. Use extra care during purification to obtain plasmid DNA of sufficiently pure quality for sequencing (see below).				
	3.	Analyze the plasmids by restriction analysis to confirm the presence and correct orientation of the insert. Use a restriction enzyme or a combination of enzymes that cut once in the vector and once in the insert.				
Sequencing	We recommend sequencing your construct to confirm that your gene is in frame with the appropriate N-terminal or C-terminal fusion tag, if desired. The table below lists the primers included in each kit to help you sequence your insert.					
		Vector	Forward Primer	Reverse Primer		
	pł	ET100/D-TOPO®	Τ7	T7 Reverse		
	pł	ET101/D-TOPO®	T7	T7 Reverse		
	pł	ET102/D-TOPO®	TrxFus Forward	T7 Reverse		

T7

T7

pET151/D-TOPO®

pET200/D-TOPO®



If you download the sequence from our Web site, note that the overhang sequence (GTGG) will be shown already hybridized to CACC. No DNA sequence analysis program allows us to show the overhang without the complementary sequence.

T7 Reverse

T7 Reverse

## Analyzing Transformants, continued

Analyzing Transformants by PCR	You may analyze positive transformants using PCR. For PCR primers, use a combination of the Forward sequencing primer or the Reverse sequencing primer and a primer that hybridizes within your insert. You will have to determine the amplification conditions. If you are using this technique for the first time, we recommend performing restriction analysis in parallel. Artifacts may be obtained because of mispriming or contaminating template. The protocol below is provided for your convenience. Other protocols are suitable.				
	Materials Needed PCR SuperMix High Fidelity (Invitrogen, Catalog no. 10790-020)				
	Appropriate forward and reverse PCR primers (20 $\mu$ M each)				
	Procedure				
	1. For each sample, aliquot 48 $\mu$ l of PCR SuperMix High Fidelity into a 0.5 ml microcentrifuge tube. Add 1 $\mu$ l each of the forward and reverse PCR primer.				
	2. Pick 5 colonies and resuspend them individually in 50 $\mu$ l of the PCR cocktail from Step 1, above.				
	3. Incubate reaction for 10 minutes at 94°C to lyse cells and inactivate nucleases.				
	4. Amplify for 20 to 30 cycles.				
	5. For the final extension, incubate at $72^{\circ}$ C for 10 minutes. Store at $+4^{\circ}$ C.				
	6. Visualize by agarose gel electrophoresis.				
<b>Q</b> Important	If you have problems obtaining transformants or the correct insert, perform the control reactions described on page 37-39. These reactions will help you troubleshoot your experiment. Refer to the <b>Troubleshooting</b> section, page 34 for additional tips.				
Long-Term Storage	Once you have identified the correct clone, be sure to purify the colony and make a glycerol stock for long-term storage. We recommend that you store a stock of plasmid DNA at -20°C.				
	1. Streak the original colony out for single colony on LB plates containing the appropriate antibiotic.				
	2. Isolate a single colony and inoculate into 1-2 ml of LB containing the appropriate antibiotic.				
	3. Grow until culture reaches stationary phase.				
	4. Mix 0.85 ml of culture with 0.15 ml of sterile glycerol and transfer to a cryovial.				
	5. Store at -80°C.				

### **Expression and Purification**

### **General Guidelines for Expression**

Introduction	BL21 Star <sup>™</sup> (DE3) One Shot <sup>®</sup> <i>E. coli</i> (Box 3) are included with Directional TOPO <sup>®</sup> Expression Kit for use as the host for en- pure plasmid DNA of your pET TOPO <sup>®</sup> construct to trans Star <sup>™</sup> (DE3) for expression studies. Since each recombinant characteristics that may affect optimal expression, we reco- time course of expression to determine the best conditions protein. Each Champion <sup>™</sup> pET Directional TOPO <sup>®</sup> Express appropriate pET TOPO <sup>®</sup> vector containing the <i>lacZ</i> gene for expression control (see below).	th each Champion <sup>™</sup> pET xpression. You will need form into BL21 t protein has different ommend performing a 6 for expression of your sion Kit also includes the or use as a positive			
BL21 Star <sup>™</sup> Strains	The BL21 Star <sup>TM</sup> (DE3) <i>E. coli</i> strain is specifically designed for expression of geness regulated by the T7 promoter. Each time you perform an expression experiment, you will transform your plasmid into BL21 Star <sup>TM</sup> (DE3). <b>Do not use this strain fo</b> <b>propagation and maintenance of your plasmid. Use TOP10 instead.</b> Basal level expression of T7 polymerase, particularly in BL21 Star <sup>TM</sup> (DE3) cells, may lead to plasmid instability if your gene of interest is toxic to <i>E. coli</i> .				
	<b>Note:</b> If you are expressing a highly toxic gene, the BL21 Star <sup>™</sup> (DE3)pLysS strain is also available from Invitrogen for expression purposes. The BL21 Star <sup>™</sup> (DE3)pLysS strain contains the pLysS plasmid to further reduce basal level expression of the gene of interest. For more information, see page 6.				
Positive Controls	<b>Te Controls</b> Each Champion <sup>™</sup> pET Directional TOPO <sup>®</sup> Expression Kit includes a positive control vector for use as an expression control (see the table below). In each case the gene encoding β-galactosidase is directionally TOPO <sup>®</sup> Cloned into the appropriate pET TOPO <sup>®</sup> vector (see pages 50-53 for details). Transform 10 ng of each plasmid into BL21 Star <sup>™</sup> (DE3) cells using the procedure on page 26.				
	Kit	Positive Control			
	Champion <sup>™</sup> pET100 Directional TOPO <sup>®</sup> Expression Kit	pET100/D/lacZ			
	Champion <sup>™</sup> pET101 Directional TOPO <sup>®</sup> Expression Kit	pET101/D/lacZ			
	Champion <sup>™</sup> pET102 Directional TOPO <sup>®</sup> Expression Kit	pET102/D/lacZ			
	Champion <sup>™</sup> pET151 Directional TOPO <sup>®</sup> Expression Kit	pET151/D/lacZ			
	Champion <sup>™</sup> pET200 Directional TOPO <sup>®</sup> Expression Kit	pET200/D/lacZ			
## General Guidelines for Expression, continued

Basic Strategy	The basic steps needed to induce expression of your gene in BL21 Star <sup><math>TM</math></sup> (DE3) <i>E. coli</i> are outlined below.
	<ol> <li>Isolate plasmid DNA using standard procedures and transform your construct and the positive control separately into BL21 Star<sup>™</sup>(DE3) One Shot<sup>®</sup> cells.</li> </ol>
	2. Grow the transformants and induce expression with IPTG over several hours. Take several time points to determine the optimal time of expression.
	3. Optimize expression to maximize the yield of protein.
Plasmid Preparation	You may prepare plasmid DNA using your method of choice. We recommend using the PureLink <sup>™</sup> HQ Mini Plasmid Purification Kit (Catalog no. K2100-01) for isolation of pure plasmid DNA. Note that since you are purifying a low-copy number plasmid, you may need to increase the amount of bacterial culture that you use to prepare your plasmid construct.
Ampicillin Selection	For pET TOPO <sup>®</sup> vectors containing the ampicillin resistance gene, ampicillin generally works well for selection of transformants and expression experiments. However, if you find that your expression levels are low, you may want to use carbenicillin instead. The resistance gene for ampicillin encodes the protein, $\beta$ -lactamase. $\beta$ -lactamase is secreted into the medium where it hydrolyzes ampicillin, inactivating the antibiotic. Since $\beta$ -lactamase is catalytic, ampicillin is rapidly removed from the medium, resulting in non-selective conditions. If your plasmid is unstable, this may result in the loss of plasmid and low expression levels.
Using Carbenicillin	Carbenicillin is generally more stable than ampicillin, and studies have shown that using carbenicillin in place of ampicillin may help to increase expression levels by preventing loss of the pET TOPO <sup>®</sup> plasmid. If you wish to use carbenicillin, perform your transformation and expression experiments in LB containing 50 $\mu$ g/ml carbenicillin.
	<b>Note:</b> If your gene of interest is highly toxic, increasing the concentration of carbenicillin used from 50 $\mu$ g/ml to 200 $\mu$ g/ml may help to increase expression levels.
Note	Note that cyclic AMP-mediated derepression of the <i>lacUV5</i> promoter in $\lambda$ DE3 lysogens can result in an increase in basal expression of T7 RNA polymerase. If you are expressing an extremely toxic gene, the pET construct may be unstable in BL21 Star <sup>™</sup> (DE3) cells. Adding 1% glucose to the bacterial culture medium may help to repress basal expression of T7 RNA polymerase and stabilize your pET construct.

# Expressing the PCR Product

Materials to Have on Hand	Be sure to have the following solutions and equipment on hand before starting your expression experiment:		
	•	Your pET TOPO <sup>®</sup> expression construct (>10 µg/ml)	
	•	The appropriate pET/D/ <i>lacZ</i> positive control plasmid	
	•	BL21 Star <sup>™</sup> (DE3) One Shot <sup>®</sup> cells (Box 3 supplied with the kit)	
<ul> <li>S.O.B. or LB containing the appropriate antibiotic for selection 1% glucose, if desired)</li> </ul>		S.O.B. or LB containing the appropriate antibiotic for selection (plus 1% glucose, if desired)	
	•	37°C incubator (shaking and nonshaking)	
	•	42°C water bath	
	•	1 M isopropyl β-D-thiogalactoside (IPTG; Invitrogen, Catalog no. 15529-019)	
	•	Liquid nitrogen	
Transforming BL21 Star™(DE3)To tr Star™One Shot® Cellsof ce of ce Note grow		transform your construct or the positive control (10 ng each) into BL21 r <sup>™</sup> (DE3) One Shot <sup>®</sup> cells, follow the instructions below. You will need one vial cells per transformation. te: You will not plate the transformation reaction, but inoculate it into medium for wth and subsequent expression.	
	1.	Thaw on ice, one vial of BL21 Star <sup>™</sup> (DE3) One Shot <sup>®</sup> cells per transformation.	
	2.	Add 5-10 ng plasmid DNA in a 1 to 5 µl volume into each vial of BL21 Star <sup>™</sup> (DE3) One Shot <sup>®</sup> cells and mix by stirring gently with the pipette tip. <b>Do not mix by pipetting up and down.</b>	
	3.	Incubate on ice for 30 minutes.	
	4.	Heat-shock the cells for 30 seconds at 42°C without shaking.	
	5.	Immediately transfer the tubes to ice.	
	6.	Add 250 µl of room temperature S.O.C. medium.	
	7.	Cap the tube tightly, tape the tube on its side (for better aeration), and incubate at 37°C for 30 minutes with shaking (200 rpm).	
	8.	Add the <b>entire</b> transformation reaction to 10 ml of LB containing the appropriate antibiotic (and 1% glucose, if desired).	
	9.	Grow overnight at 37°C with shaking. Proceed to <b>Pilot Expression</b> , next page.	

# Expressing the PCR Product, continued

Pilot Expression	1.	Inoculate 10 ml of LB containing the appropriate antibiotic (and 1% glucose, if desired) with 500 $\mu$ l of the overnight culture from Step 8, previous page.
	2.	Grow two hours at 37°C with shaking. $OD_{600}$ should be about 0.5-0.8 (midlog).
	3.	Split the culture into two 5 ml cultures. Add IPTG to a final concentration of 0.5-1 mM to one of the cultures. You will now have two cultures: one induced, one uninduced.
	4.	Remove a 500 $\mu$ l aliquot from <b>each</b> culture, centrifuge at maximum speed in a microcentrifuge for 30 seconds, and aspirate the supernatant.
	5.	Freeze the cell pellets at -20°C. These are the zero time point samples.
	6.	Continue to incubate the cultures at 37°C with shaking. Take time points for each culture every hour for 4 to 6 hours.
	7.	For each time point, remove 500 $\mu$ l from the induced and uninduced cultures and process as described in Steps 4 and 5. Proceed to <b>Analyzing Samples</b> , next page.

## **Analyzing Samples**

Materials to Have on Hand	<ul> <li>Have the following materials on hand before starting:</li> <li>Lysis Buffer (see page 55 for recipe)</li> <li>1X and 2X SDS-PAGE sample buffer (see page 55 for recipes)</li> <li>Reagents and apparatus to perform SDS-PAGE electrophoresis</li> <li>Boiling water bath</li> </ul>			
Preparing Samples	Once you have finished your pilot expression, you are ready to analyze the samples you have collected. Before starting, prepare SDS-PAGE gels or use one of the pre-cast polyacrylamide gels available from Invitrogen (see below).			
	Note: If you wish to analyze your samples for soluble protein, see below.			
	1. Thaw the samples (from Pilot Expression, Steps 5 and 7, previous page) and resuspend each cell pellet in 80 $\mu$ l of 1X SDS-PAGE sample buffer.			
	2. Boil 5 minutes and centrifuge briefly.			
	<ol> <li>Load 5-10 μl of each sample on an SDS-PAGE gel and electrophorese. Save your samples by storing them at -20°C.</li> </ol>			
Preparing Samples for	<ol> <li>Thaw and resuspend each cell pellet in 500 μl of Lysis Buffer (see Recipes, page 55).</li> </ol>			
Soluble/Insoluble Protein	2. Freeze sample in dry ice or liquid nitrogen and then thaw at 42°C. Repeat 2 to 3 times.			
	Note: To facilitate lysis, you may need to add lysozyme or sonicate the cells.			
	3. Centrifuge samples at maximum speed in a microcentrifuge for 1 minute at +4°C to pellet insoluble proteins. Transfer supernatant to a fresh tube and store on ice.			
	4. Mix together equivalent amounts of supernatant and 2X SDS-PAGE sample buffer and boil for 5 minutes.			
	5. Add 500 μl of 1X SDS-PAGE sample buffer to the pellets from Step 3 and boil 5 minutes.			
	<ol> <li>Load 10 μl of the supernatant sample and 5 μl of the pellet sample onto an SDS-PAGE gel and electrophorese.</li> </ol>			
Polyacrylamide Gel Electrophoresis	To facilitate separation and visualization of your recombinant fusion protein by polyacrylamide gel electrophoresis, a wide range of pre-cast NuPAGE <sup>®</sup> and Novex <sup>®</sup> Tris-Glycine polyacrylamide gels and electrophoresis apparatus are available from Invitrogen. In addition, Invitrogen also carries a large selection of molecular weight protein standards and staining kits. For more information about the appropriate gels, standards, and stains to use to visualize your recombinant protein, refer to our Web site (www.invitrogen.com) or call Technical Service (see page 56).			

#### **Analyzing Samples, continued**

Analyzing Samples	To c perf	To determine the success of your expression experiment, you may want to perform the following types of analyses:		
	1.	Stain the polyacrylamide gel with Coomassie blue and look for a band of increasing intensity in the expected size range for the recombinant protein. Use the uninduced culture as a negative control.		
	2.	Perform a western blot to confirm that the overexpressed band is your desired protein (see below).		
	3.	<ol> <li>Use the positive control to confirm that growth and induction were performed properly. The table below lists the size of the β-galactosidase fusion protein expressed from each positive control vector.</li> <li>Note: β-galactosidase Antiserum is available from Invitrogen (Catalog no. R901-25) to detect β-galactosidase fusion proteins by western blot.</li> </ol>		
		Vector	Size of the Control Protein	
		pET100/D/lacZ	121 kDa	
		pET101/D/lacZ	120 kDa	
		pET102/D/lacZ	133 kDa	
		pET151/D/lacZ	120 kDa	
		pET200/D/lacZ	121 kDa	

#### Detecting Recombinant Fusion Proteins

To detect expression of your recombinant fusion protein by western blot analysis, you may use:

- Antibodies against the appropriate epitope available from Invitrogen (see page xi for ordering information)
- An antibody to your protein of interest

In addition, the Positope<sup>™</sup> Control Protein (Catalog no. R900-50) is available from Invitrogen for use as a positive control for detection of fusion proteins containing an Xpress<sup>™</sup>, HisG, V5, or C-terminal 6xHis epitope. The WesternBreeze<sup>®</sup> Chromogenic Kits and WesternBreeze<sup>®</sup> Chemiluminescent Kits are available from Invitrogen to facilitate detection of antibodies by colorimetric or chemiluminescence methods. For more information, refer to our Web site (www.invitrogen.com) or call Technical Service (see page 56).

### **Analyzing Samples, continued**



Expression of your protein with the N- and/or C-terminal tags will increase the size of your recombinant protein. The table below lists the increase in the molecular weight of your recombinant fusion protein that you should expect from the particular N- or C-terminal fusion tag in each pET TOPO® vector. Be sure to account for any additional amino acids between the fusion tag and the start of your protein.

Vector	Fusion Tag	Expected Size Increase (kDa)
pET100/D-TOPO®	N-terminal	3 kDa
pET101/D-TOPO®	C-terminal	3 kDa
pET102/D-TOPO®	N-terminal	13 kDa
	C-terminal	3 kDa
pET151/D-TOPO®	N-terminal	4 kDa
pET200/D-TOPO®	N-terminal	3 kDa

The Next StepIf you are satisfied with expression of your gene of interest, proceed to Purifying<br/>the Recombinant Fusion Protein, page 31.

If you have trouble expressing your protein or wish to optimize expression, refer to the **Troubleshooting** section, page 37.

## Purifying the Recombinant Fusion Protein

Introduction	The presence of the N-terminal or C-terminal polyhistidine (6xHis) tag in all pET TOPO <sup>®</sup> vectors allows purification of your recombinant fusion protein with a metal-chelating resin such as ProBond <sup>™</sup> or Ni-NTA.			
ProBond <sup>™</sup> and Ni-NTA	ProBond <sup>™</sup> and Ni-NTA are nickel-charged agarose resins that can be used for affinity purification of fusion proteins containing the 6xHis tag and/or the HP-thioredoxin protein. Proteins bound to the resin may be eluted with either low pH buffer or competition with imidazole or histidine.			
	• To scale up your pilot expression for purification, see below.			
	• To purify your fusion protein using ProBond <sup>™</sup> or Ni-NTA, refer to the manual included with each product. You may download the manuals from our Web site (www.invitrogen.com).			
	• To purify your fusion protein using another metal-chelating resin, refer to the manufacturer's instructions.			
Important	Note that under denaturing conditions, the Ni <sup>2+</sup> binding site encoded by the histidine "patch" in pET102/D-TOPO <sup>®</sup> will be destroyed because the HP-thioredoxin protein will be denatured. The binding of nickel ion to the 6xHis tag is not affected by denaturing conditions.			
Scaling-up Expression for Purification	We generally scale-up expression to a 50 ml bacterial culture for purification using a 2 ml ProBond <sup>™</sup> or Ni-NTA column. Depending on the expression level of your recombinant fusion protein, you may need to adjust the culture volume to bind the maximum amount of recombinant fusion protein to your column.			
	To grow and induce a 50 ml bacterial culture:			
	<ol> <li>Inoculate 10 ml of S.O.B. or LB containing the appropriate antibiotic with 500 μl of the culture of the transformation reaction (see page 26, step 9).</li> </ol>			
	2. Grow overnight at 37°C with shaking (225-250 rpm) to $OD_{600} = 1-2$ .			
	3. The next day, inoculate 50 ml of S.O.B. or LB containing the appropriate antibiotic with 1 ml of the overnight culture.			
	<b>Note</b> : You can scale up further and inoculate all of the 10 ml overnight culture into 500 ml of medium, but you will need to adjust the bed volume of your ProBond <sup>™</sup> or Ni-NTA column accordingly.			
	4. Grow the culture at $37^{\circ}$ C with shaking (225-250 rpm) to an OD <sub>600</sub> = ~0.5 (2-3 hours). The cells should be in mid-log phase.			
	5. Add 0.5-1 mM IPTG to induce expression.			
	6. Grow at 37°C with shaking until the optimal time point determined by the pilot expression is reached. Harvest the cells by centrifugation (3000 x g for 10 minutes at +4°C).			
	7. Proceed to purification or store the cells at -80°C for future use.			

## Purifying the Recombinant Fusion Protein, continued

# Additional There may be cases when your specific fusion protein may not be completely purification Steps There may be utilized in conjunction with ProBond<sup>™</sup> or Ni-NTA to purify the fusion protein (see Deutscher, 1990 for more information).

### **Removing N-terminal Fusion Tags**

pET200/D-TOPO®

Introduction	If you have expressed your recombinant fusion protein from any N-terminal fusion pET TOPO <sup>®</sup> vector, the presence of either the enterokinase (EK) or TEV recognition site in the vector allows removal of the N-terminal fusion tag from the recombinant fusion protein after purification using enterokinase or TEV protease, as appropriate. General guidelines are provided below.		
Removal of the N-terminal Fusion Tag Using Enterokinase	If you have expressed your recombinant fusion protein from pET TOPO <sup>®</sup> vectors containing the EK recognition site (see below), you may use a recombinant preparation of the catalytic subunit of bovine enterokinase (EKMax <sup>™</sup> ) available from Invitrogen (Catalog no. E180-01) to remove the N-terminal fusion tag. Instructions for digestion are included with the product. For more information, contact Technical Service (see page 56). To remove EKMax <sup>™</sup> from the digest, EK-Away <sup>™</sup> (Catalog no. R180-01) is also available		
	<b>Note:</b> After digestion with enterokinase, a number of vector-encoded amino acids will remain at the N-terminus of your protein (see table below).		
	Vector	Number of Amino Acids Remaining After EK Cleavage	
	pET100/D-TOPO®	5	
	pET102/D-TOPO®	7	

#### Removal of the N-terminal Fusion Tag Using TEV Protease

If you have expressed your recombinant fusion protein from pET151/D-TOPO<sup>®</sup>, you may use recombinant TEV Protease available from Invitrogen (Catalog no. 12575-015) to remove the N-terminal fusion tag. Instructions for digestion are included with the product. For more information, contact Technical Service (see page 56).

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**Note:** After digestion with enterokinase, six vector-encoded amino acids will remain at the N-terminus of your protein.

## Troubleshooting

TOPO <sup>®</sup> Cloning	The table below lists some potential problems and possible solutions that may
Reaction and	help you troubleshoot the TOPO® Cloning and transformation reactions. To help
Transformation	evaluate your results, we recommend that you perform the control reactions (see pages 37-39) in parallel with your samples.

Problem	Reason	Solution
Few or no colonies obtained from sample reaction <b>and</b> the transformation control	Suboptimal ratio of PCR product:TOPO <sup>®</sup> vector used in the TOPO <sup>®</sup> Cloning reaction	Use a 0.5:1 to 2:1 molar ratio of PCR product:TOPO <sup>®</sup> vector.
gave colonies	Too much PCR product used in	Dilute the PCR product.
	the TOPO <sup>®</sup> Cloning reaction	Use a 0.5:1 to 2:1 molar ratio of PCR product:TOPO <sup>®</sup> vector.
	PCR product too dilute	• Concentrate the PCR product.
		• Use a 0.5:1 to 2:1 molar ratio of PCR product:TOPO <sup>®</sup> vector.
	PCR primers contain 5' phosphates	Do not add 5' phosphates to your PCR primers.
	Incorrect PCR primer design	Make sure that the forward PCR primer contains the sequence, CACC, at the 5' end.
		Make sure that the reverse PCR primer <b>does not</b> contain the sequence, CACC, at the 5' end.
	Used <i>Taq</i> polymerase or a <i>Taq</i> /proofreading polymerase mixture for PCR	Use a proofreading polymerase for PCR.
	Large PCR product	Increase the incubation time of the TOPO <sup>®</sup> reaction from 5 minutes to 30 minutes.
		Gel-purify the PCR product to remove primer-dimers and other artifacts.
	PCR reaction contains artifacts ( <i>i.e.</i> does not run as a single, discrete band on an agarose gel)	• Optimize your PCR using the proofreading polymerase of your choice.
		• Gel-purify your PCR product.
	Cloning large pool of PCR products or a toxic gene	• Increase the incubation time of the TOPO <sup>®</sup> reaction from 5 minutes to 30 minutes.
		• Use a 0.5:1 to 2:1 molar ratio of PCR product:TOPO <sup>®</sup> vector.

## Troubleshooting, continued

Problem	Reason	Solution
Large percentage of inserts cloned in the incorrect orientation	Incorrect PCR primer design	Make sure that the forward PCR primer contains the sequence, CACC, at the 5' end.
	Reverse PCR primer is complementary to the GTGG overhang at the 5' end	Make sure that the reverse PCR primer <b>does not</b> contain the sequence, CACC, at the 5' end.
Large number of incorrect inserts cloned	PCR cloning artifacts	Gel-purify your PCR product to remove primer-dimers and smaller PCR products.
		Optimize your PCR.
		Include a final extension step of 7 to 30 minutes during PCR. Longer PCR products will need a longer extension time.
	Incorrect PCR primer design	Make sure that the forward and reverse PCR primers are designed correctly.
Few or no colonies obtained from sample reaction <b>and</b>	One Shot <sup>®</sup> competent <i>E. coli</i> stored incorrectly	Store One Shot <sup>®</sup> competent <i>E. coli</i> at -80°C.
the transformation control gave <b>no</b> colonies		If you are using another <i>E. coli</i> strain, follow the manufacturer's instructions.
	One Shot <sup>®</sup> transformation protocol not followed correctly	Follow the One Shot <sup>®</sup> transformation protocol provided on page 20.
	Insufficient amount of <i>E. coli</i> plated	Increase the amount of <i>E. coli</i> plated.
	Transformants plated on selective plates containing the wrong antibiotic	Use the appropriate antibiotic for selection.

#### TOPO<sup>®</sup> Cloning Reaction and Transformation, continued

## Troubleshooting, continued

#### Expression

The table below lists some potential problems and possible solutions that may help you troubleshoot your expression experiment. To help evaluate your results, we recommend including the expression control supplied with kit in your experiment.

Problem	Reason	Solution
No expression of recombinant protein	Gene of interest not in frame with the epitope tag	Sequence your construct to verify if the insert is in frame with the epitope tag. If not in frame, redesign your PCR primers.
	Incorrect antibody used for detection	Use an antibody to your protein or one of the antibodies listed on page xi, as appropriate
Low expression	Plasmid instability observed when using ampicillin for selection	Substitute carbenicillin for ampicillin in your transformation and expression experiments (see page 25).
	Toxic gene <b>Note:</b> Evidence of toxicity includes loss of plasmid or slow growth relative to the control.	• Add 1% glucose to the bacterial culture medium during transformation and expression.
		• Transform BL21 Star <sup>™</sup> (DE3) cells using the protocol on page 26, then perform the expression by growing cells at room temperature rather than 37°C for 24-48 hours.
		• Transform your expression construct into a pLysS-containing strain ( <i>e.g.</i> BL21 Star <sup>™</sup> (DE3)pLysS).
		<ul> <li>Transform your expression construct into an <i>E. coli</i> strain in which expression of T7 RNA polymerase is tightly regulated (<i>e.g.</i> BL21-AI<sup>™</sup> available from Invitrogen; see our Web site for more information).</li> </ul>
		• Infect TOP10F' (or other suitable F' episome-containing host strain) with M13 or lambda phage expressing T7 RNA polymerase.

# Appendix

## **Performing the Control Reactions**

Introduction	We recommend performing the following control TOPO <sup>®</sup> Cloning reactions the first time you use the kit to help you evaluate your results. Performing the control reactions involves producing a control PCR product using the reagents included in the kit and using this product directly in a TOPO <sup>®</sup> Cloning reaction.				
Before Starting	For each transformation, prepare two LB plates containing the appropriate antibiotic (see page 54 for a recipe).				
Producing the Control PCR Product	Use your thermostable, proofreading polymerase and the appropriate buffer to amplify the control PCR product. Follow the manufacturer's recommendations for the polymerase you are using.				
	1.	To produce the 750 bp o	control PCR prod	uct, set up the follo	owing 50 µl PCR:
		Control DNA Template	e (100 ng)	1 µl	
		10X PCR Buffer (approj	priate for enzyme	e) 5 μl	
		dNTP Mix 0.5 μl			
		Control PCR Primers (0.1 $\mu$ g/ $\mu$ l each) 1 $\mu$		1 µl	
		Sterile Water		41.5 μl	
Thermostable polymerase (1-2.5 units/ $\mu$ l) 1 $\mu$ l					
	<ul> <li>Total Volume 50 μl</li> <li>Overlay with 70 μl (1 drop) of mineral oil, if required.</li> <li>Amplify using the following cycling parameters:</li> </ul>				
		Step	Time	Temperature	Cycles
		Initial Denaturation	2 minutes	94°C	1X
		Denaturation	1 minute	94°C	
		Annealing	1 minute	55°C	25X
		Extension	1 minute	72°C	-
		Final Extension	7 minutes	72°C	1X
	4.	Remove 10 μl from the discrete 750 bp band sh	reaction and anal ould be visible.	yze by agarose gel	electrophoresis. A
	<ol> <li>Estimate the concentration of the PCR product, and adjust as necessary s that the amount of PCR produce used in the control TOPO<sup>®</sup> Cloning rea results in an optimal molar ratio of PCR product:TOPO<sup>®</sup> vector (<i>i.e.</i> 0.5:1 2:1). Proceed to Control TOPO<sup>®</sup> Cloning Reactions, next page.</li> </ol>			as necessary such <sup>®</sup> Cloning reaction ector ( <i>i.e.</i> 0.5:1 to page.	

#### Performing the Control Reactions, continued

#### Control TOPO<sup>®</sup> Cloning Reactions

1. Using the control PCR product produced on the previous page and the pET TOPO<sup>®</sup> vector, set up two 6 µl TOPO<sup>®</sup> Cloning reactions as described below. If you plan to transform electrocompetent *E. coli*, use Dilute Salt Solution in place of the Salt Solution.

2. Set up control TOPO<sup>®</sup> Cloning reactions:

Reagent	"Vector Only"	"Vector + PCR Insert"
Sterile Water	4 μl	3 μl
Salt Solution	1 μl	1 µl
Control PCR Product		1 µl
pET TOPO <sup>®</sup> vector	1 µl	1 μl
Final volume	6 µl	6 µl

3. Incubate at room temperature for 5 minutes and place on ice.

- 4. Transform 3 1 of each reaction into separate vials of One Shot<sup>®</sup> TOP10 cells (page 20).
- 5. Spread 100-200 l of each transformation mix onto LB plates containing the appropriate antibiotic. Be sure to plate two different volumes to ensure that at least one plate has well-spaced colonies.
- 6. Incubate overnight at 37°C.

#### Analysis of Results

Hundreds of colonies from the vector + PCR insert reaction should be produced. To analyze the transformations, isolate plasmid DNA and digest with the appropriate restriction enzyme as listed below. The table below lists the digestion patterns that you should see for inserts that are cloned in the correct orientation or in the reverse orientation.

Vector	Restriction Enzyme	Expected Digestion Patterns (bp)
pET100/D-TOPO®	Hind III	Correct orientation: 838, 5674
		Reverse orientation: 500, 6012
		Empty vector: 5764
pET101/D-TOPO®	Cla I	Correct orientation: 599, 5896
		Reverse orientation: 929, 5566
		Empty vector: 5753
pET102/D-TOPO®	Pst I	Correct orientation: 1990, 5065
		Reverse orientation: 1484, 5571
		Empty vector: 6315

### Performing the Control Reactions, continued

#### Analysis of Results, continued

Vector	Restriction Enzyme	Expected Digestion Patterns (bp)
pET151/D-TOPO®	EcoR I and Sst I	Correct orientation: 653, 5861
		Reverse orientation: 101, 6413
		Empty vector: 5760
pET200/D-TOPO®	Hind III	Correct orientation: 845, 5646
		Reverse orientation: 511, 5980
		Empty vector: 5741

Greater than 90% of the colonies should contain the 750 bp insert in the correct orientation.

Relatively few colonies should be produced in the vector-only reaction.

# Transformation<br/>ControlpUC19 plasmid is included to check the transformation efficiency of the One<br/>Shot® TOP10 competent cells. Transform one vial of One Shot® TOP10 cells with<br/>10 pg of pUC19 using the protocol on page 20. Plate 10 µl of the transformation<br/>mixture plus 20 µl of S.O.C. on LB plates containing 100 µg/ml ampicillin.<br/>Transformation efficiency should be ~1 x 10° cfu/µg DNA.

## Gel Purifying PCR Products

Introduction	Smearing, multiple banding, primer-dimer artifacts, or large PCR products (>3 kb) may necessitate gel purification. If you intend to purify your PCR product, be extremely careful to remove all sources of nuclease contamination. There are many protocols to isolate DNA fragments or remove oligonucleotides. Refer to <i>Current Protocols in Molecular Biology</i> , Unit 2.6 (Ausubel <i>et al.</i> , 1994) for the most common protocols. Two simple protocols are provided below.	
Using the PureLink <sup>™</sup> Quick	The PureLink™ Quick Gel Extraction Kit (page x) allows you to rapidly purify PCR products from regular agarose gels.	
Gel Extraction Kit	1. Equilibrate a water bath or heat block to 50°C.	
	2. Cut the area of the gel containing the desired DNA fragment using a clean, sharp blade. Minimize the amount of surrounding agarose excised with the fragment.	
	3. Weigh the gel slice.	
	4. Add Gel Solubilization Buffer (GS1) supplied in the kit as follows:	
	<ul> <li>For ≤2% agarose gels, place up to 400 mg gel into a sterile, 1.5-ml polypropylene tube. Divide gel slices exceeding 400 mg among additional tubes. Add 30 µl Gel Solubilization Buffer (GS1) for every 10 mg of gel.</li> </ul>	
	<ul> <li>For &gt;2% agarose gels, use sterile 5-ml polypropylene tubes and add 60 µl Gel Solubilization Buffer (GS1) for every 10 mg of gel.</li> </ul>	
	<ol> <li>Incubate the tube at 50°C for 15 minutes. Mix every 3 minutes to ensure gel dissolution. After gel slice appears dissolved, incubate for an additional 5 minutes.</li> </ol>	
	6. Preheat an aliquot of TE Buffer (TE) to 65-70°C	
	7. Place a Quick Gel Extraction Column into a Wash Tube. Pipette the mixture from Step 5, above onto the column. Use 1 column per 400 mg agarose.	
	8. Centrifuge at >12,000 x g for 1 minute. Discard the flow-through. Place the column back into the Wash Tube.	
	9. <b>Optional:</b> Add 500 µl Gel Solubilization Buffer (GS1) to the column. Incubate at room temperature for 1 minute. Centrifuge at >12,000 x g for 1 minute. Discard the flow-through. Place the column back into the Wash Tube.	
	10. Add 700 μl Wash Buffer (W9) with ethanol (add 96–100% ethanol to the Wash Buffer according to instructions on the label of the bottle) to the column and incubate at room temperature for 5 minutes. Centrifuge at >12,000 x g for 1 minute. Discard flow-through.	
	<ol> <li>Centrifuge the column at &gt;12,000 x g for 1 minute to remove any residual buffer. Place the column into a 1.5 ml Recovery Tube.</li> </ol>	
	<ol> <li>Add 50 μl warm (65-70°C) TE Buffer (TE) to the center of the cartridge. Incubate at room temperature for 1 minute.</li> </ol>	
	13. Centrifuge at >12,000 x g for 2 minutes. <i>The Recovery Tube contains the purified DNA</i> . Store DNA at –20°C. Discard the column.	
	14. Use 4 $\mu$ l of the purified DNA for the TOPO <sup>®</sup> Cloning reaction.	

## Gel Purifying PCR Products, continued

Low-Melt Agarose Method	If y pur cloi	ou prefer to use low-melt agarose, use the procedure below. Note that gel rification will result in dilution of your PCR product and a potential loss of ning efficiency. Use only chemically competent cells for transformation.
	1.	Electrophorese as much as possible of your PCR reaction on a low-melt agarose gel (0.8 to 1.2%) in TAE buffer.
	2.	Visualize the band of interest and excise the band.
	3.	Place the gel slice in a microcentrifuge tube and incubate the tube at 65°C until the gel slice melts.
	4.	Place the tube at 37°C to keep the agarose melted.
	5.	Add 4 $\mu$ l of the melted agarose containing your PCR product to the TOPO <sup>®</sup> Cloning reaction as described on page 18.
	6.	Incubate the TOPO <sup>®</sup> Cloning reaction <b>at 37°C for 5 to 10 minutes</b> . This is to keep the agarose melted.
	7.	Transform 2 to 4 $\mu l$ directly into One Shot® TOP10 cells using the method on page 20.
Note	The ma	e cloning efficiency may decrease with purification of the PCR product. You y wish to optimize your PCR to produce a single band.

## Map and Features of pET100/D-TOPO<sup>®</sup> and pET200/D-TOPO<sup>®</sup>

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The figure below shows the features of the pET100/D-TOPO<sup>®</sup> (5764 bp) and pET200/D-TOPO<sup>®</sup> (5741 bp) vectors. The complete sequence of each vector is available for downloading from our Web site (www.invitrogen.com) or by contacting Technical Service (see page 56).



	<u>pET100/D-TOPO®</u>	<u>pET200/D-TOPO®</u>
T7 promoter	209-225	209-225
T7 promoter/priming site	209-228	209-228
<i>lac</i> operator (lacO)	228-252	228-252
Ribosome binding site (RBS)	282-288	282-288
Initiation ATG	297-299	297-299
Polyhistidine (6xHis) region	309-326	309-326
Xpress <sup>™</sup> epitope	366-389	366-389
EK recognition site	375-389	375-389
TOPO <sup>®</sup> Cloning site (directional)	396-409	396-409
T7 reverse priming site	466-485	466-485
T7 transcription termination region	427-555	427-555
<i>bla</i> promoter	856-954	856-954
Ampicillin (bla) resistance gene	955-1815	
Kanamycin resistance gene		955-1749
pBR322 origin	2022-2757	1969-2610
ROP ORF (complementary strand)	3001-3192	2978-3169
lacI ORF (complementary strand)	4507-5595	4481-5572

# Map and Features of pET100/D-TOPO<sup>®</sup> and pET200/D-TOPO<sup>®</sup>, continued

Features of pET100/D-TOPO<sup>®</sup> and pET200/D-TOPO<sup>®</sup> The pET100/D-TOPO<sup>®</sup> (5764 bp) and pET200/D-TOPO<sup>®</sup> (5741 bp) vectors contain the following elements. All features have been functionally tested.

Feature	Benefit
T7 promoter	Permits high-level, IPTG-inducible expression of your recombinant protein in <i>E. coli</i> strains expressing the T7 RNA polymerase.
T7 forward priming site	Allows sequencing in the sense orientation.
<i>lac</i> operator (lacO)	Binding site for lac repressor that serves to reduce basal expression of your recombinant protein.
Ribosome binding site	Optimally spaced from the TOPO <sup>®</sup> Cloning site for efficient translation of PCR product.
N-terminal 6xHis tag	Permits purification of recombinant fusion protein on metal-chelating resin ( <i>i.e.</i> ProBond <sup>™</sup> or Ni-NTA).
	In addition, allows detection of recombinant protein with the Anti-HisG Antibodies.
Xpress <sup>™</sup> epitope	Allows detection of the fusion protein by the Anti-
(Asp-Leu-Tyr-Asp-Asp-Asp-Asp-Lys)	Xpress <sup>™</sup> Antibodies.
Enterokinase (EK) recognition site	Allows removal of the N-terminal tag from your
(Asp-Asp-Asp-Asp-Lys)	recombinant protein using an enterokinase such as EKMax <sup>™</sup> .
TOPO <sup>®</sup> Cloning site (directional)	Permits rapid cloning of your PCR product for expression in <i>E. coli</i> .
T7 Reverse priming site	Allows sequencing of the insert.
T7 transcription termination region	Sequence from bacteriophage T7 which permits efficient transcription termination.
bla promoter	Allows expression of the ampicillin resistance gene.
Ampicillin resistance gene (β-lactamase) (pET100/D-TOPO <sup>®</sup> only)	Allows selection of the plasmid in <i>E. coli</i> .
Kanamycin resistance gene	Allows selection of the plasmid in <i>E. coli</i> .
(pET200/D-TOPO <sup>®</sup> only)	
pBR322 origin of replication ( <i>ori</i> )	Permits replication and maintenance in <i>E. coli</i> .
ROP ORF	Interacts with the pBR322 origin to facilitate low- copy replication in <i>E. coli</i> .
lacI ORF	Encodes lac repressor which binds to the T7 <i>lac</i> promoter to block basal transcription of the gene of interest and to the <i>lacUV5</i> promoter in the host chromosome to repress transcription of T7 RNA polymerase.

## Map and Features of pET101/D-TOPO®

pET101/D-TOPO<sup>®</sup> Map The figure below shows the features of the pET101/D-TOPO<sup>®</sup> (5753 bp) vector. The complete sequence of the vector is available for downloading from our Web site (www.invitrogen.com) or by contacting Technical Service (see page 56).



# Map and Features of pET101/D-TOPO<sup>®</sup>, continued

# Features of pET101/D-TOPO<sup>®</sup>

The pET101/D-TOPO<sup>®</sup> (5753 bp) vector contains the following elements. All features have been functionally tested.

Feature	Benefit	
T7 promoter	Permits high-level, IPTG-inducible expression of your recombinant protein in <i>E. coli</i> strains expressing the T7 RNA polymerase.	
T7 forward priming site	Allows sequencing in the sense orientation.	
<i>lac</i> operator (lacO)	Binding site for lac repressor that serves to reduce basal expression of your recombinant protein.	
Ribosome binding site	Optimally spaced from the TOPO <sup>®</sup> Cloning site for efficient translation of PCR product.	
TOPO <sup>®</sup> Cloning site (directional)	Permits rapid cloning of your PCR product for expression in <i>E. coli</i> .	
C-terminal V5 epitope tag (Gly-Lys-Pro-Ile-Pro-Asn-Pro-Leu-Leu- Gly-Leu-Asp-Ser-Thr)	Allows detection of the fusion protein by the Anti- V5 Antibodies (Southern <i>et al.</i> , 1991).	
C-terminal 6xHis tag	Permits purification of recombinant fusion protein on metal-chelating resins ( <i>i.e.</i> ProBond <sup>™</sup> or Ni-NTA).	
	In addition, allows detection of recombinant protein with the Anti-His(C-term) Antibodies (Lindner <i>et al.</i> , 1997).	
T7 Reverse priming site	Allows sequencing of the insert.	
T7 transcription termination region	Sequence from T7 bacteriophage which permits efficient transcription termination.	
<i>bla</i> promoter	Allows expression of the ampicillin resistance gene.	
Ampicillin resistance gene (β-lactamase)	Allows selection of the plasmid in <i>E. coli</i> .	
pBR322 origin of replication (ori)	Permits replication and maintenance in <i>E. coli</i> .	
ROP ORF	Interacts with the pBR322 origin to facilitate low-copy replication in <i>E. coli</i> .	
lacI ORF	Encodes lac repressor which binds to the T7 <i>lac</i> promoter to block basal transcription of the gene of interest and to the <i>lacUV5</i> promoter in the host chromosome to repress transcription of T7 RNA polymerase.	

# Map and Features of pET102/D-TOPO®

pET102/D-TOPO<sup>®</sup> Map The figure below shows the features of the pET102/D-TOPO<sup>®</sup> (6315 bp) vector. The complete sequence of the vector is available for downloading from our Web site (www.invitrogen.com) or by contacting Technical Service (see page 56).



	pET102/D-TOPO®
T7 promoter	209-225
T7 promoter/priming site	209-228
lac operator (lacO)	228-252
Ribosome binding site (RBS)	282-288
His-patch (HP) thioredoxin ORF	298-627
TrxFus forward priming site	607-624
EK recognition site	643-657
TOPO <sup>®</sup> Cloning site (directional)	670-683
V5 epitope	700-741
Polyhistidine (6xHis) region	751-768
T7 reverse priming site	822-841
T7 transcription termination region	783-911
<i>bla</i> promoter	1407-1505
Ampicillin (bla) resistance gene	1506-2366
pBR322 origin	2511-3184
ROP ORF (complementary strand)	3552-3743
<i>lacI</i> ORF (complementary strand)	5055-6146

# Map and Features of pET102/D-TOPO<sup>®</sup>, continued

# Features of pET102/D-TOPO<sup>®</sup>

The pET102/D-TOPO<sup>®</sup> (6315 bp) vector contains the following elements. All features have been functionally tested.

Feature	Benefit	
T7 promoter	Permits high-level, IPTG-inducible expression of your protein in <i>E. coli</i> strains expressing the T7 RNA polymerase.	
T7 forward priming site	Allows sequencing in the sense orientation.	
<i>lac</i> operator (lacO)	Binding site for lac repressor that serves to reduce basal expression of your recombinant protein.	
Ribosome binding site	Optimally spaced from the TOPO <sup>®</sup> Cloning site for efficient translation of PCR product.	
His-Patch (HP)-thioredoxin	Provides an efficient fusion partner for translation of the fusion protein.	
TrxFus forward priming site	Permits sequencing of the insert in the sense orientation.	
Enterokinase (EK) recognition site	Allows removal of the N-terminal tag from your	
(Asp-Asp-Asp-Asp-Lys)	protein using an enterokinase such as EKMax <sup>™</sup> .	
TOPO <sup>®</sup> Cloning site (directional)	Permits rapid cloning of your PCR product.	
C-terminal V5 epitope tag	Allows detection of the fusion protein by the Anti- V5 Antibodies (Southern <i>et al.</i> , 1991).	
C-terminal 6xHis tag	Permits purification of recombinant fusion protein on metal-chelating resins ( <i>i.e.</i> ProBond <sup>™</sup> or Ni-NTA).	
	In addition, allows detection of the recombinant protein with the Anti-His(C-term) Antibodies (Lindner <i>et al.</i> , 1997).	
T7 Reverse priming site	Allows sequencing of the insert.	
T7 transcription termination region	Sequence from T7 bacteriophage which permits efficient transcription termination.	
<i>bla</i> promoter	Allows expression of the ampicillin resistance gene.	
Ampicillin resistance gene ( $\beta$ -lactamase) (pET102/D-TOPO <sup>®</sup> only)	Allows selection of the plasmid in <i>E. coli</i> .	
pBR322 origin of replication ( <i>ori</i> )	Permits replication and maintenance in <i>E. coli</i> .	
ROP ORF	Interacts with the pBR322 origin to facilitate low- copy replication in <i>E. coli</i> .	
lacI ORF	Encodes lac repressor which binds to the T7 <i>lac</i> promoter to block basal transcription of the gene of interest and to the <i>lacUV5</i> promoter in the host chromosome to repress transcription of T7 RNA polymerase.	

## Map and Features of pET151/D-TOPO<sup>®</sup>

The figure below shows the features of the pET151/D-TOPO<sup>®</sup> (5760 bp) vector. pET151/D-TOPO® The complete sequence of the vector is available for downloading from our Web site (www.invitrogen.com) or by contacting Technical Service (see page 56).



continued on next page

Map of

# Map and Features of pET151/D-TOPO<sup>®</sup>, continued

# Features of pET151/D-TOPO<sup>®</sup>

The pET151/D-TOPO<sup>®</sup> (5760 bp) vector contains the following elements. All features have been functionally tested.

Feature	Benefit
T7 promoter	Permits high-level, IPTG-inducible expression of your recombinant protein in <i>E. coli</i> strains expressing the T7 RNA polymerase.
T7 forward priming site	Allows sequencing in the sense orientation.
<i>lac</i> operator (lacO)	Binding site for lac repressor that serves to reduce basal expression of your recombinant protein.
Ribosome binding site	Optimally spaced from the TOPO <sup>®</sup> Cloning site for efficient translation of PCR product.
N-terminal 6xHis tag	Permits purification of recombinant fusion protein on metal-chelating resin ( <i>i.e.</i> ProBond <sup>™</sup> or Ni-NTA).
	In addition, allows detection of recombinant protein with the Anti-HisG Antibodies.
V5 epitope	Allows detection of the fusion protein by the Anti- V5 Antibodies (Southern <i>et al.</i> , 1991).
TEV recognition site	Allows removal of the N-terminal tag from your recombinant protein using TEV protease (Carrington & Dougherty, 1988; Dougherty <i>et al.</i> , 1988).
TOPO <sup>®</sup> Cloning site (directional)	Permits rapid cloning of your PCR product for expression in <i>E. coli</i> .
T7 Reverse priming site	Allows sequencing of the insert.
T7 transcription termination region	Sequence from bacteriophage T7 which permits efficient transcription termination.
<i>bla</i> promoter	Allows expression of the ampicillin resistance gene.
Ampicillin resistance gene (β-lactamase)	Allows selection of the plasmid in <i>E. coli</i> .
pBR322 origin of replication (ori)	Permits replication and maintenance in E. coli.
<i>ROP</i> ORF	Interacts with the pBR322 origin to facilitate low- copy replication in <i>E. coli</i> .
lacI ORF	Encodes lac repressor which binds to the T7 <i>lac</i> promoter to block basal transcription of the gene of interest and to the <i>lacUV5</i> promoter in the host chromosome to repress transcription of T7 RNA polymerase.

#### Map of pET100/D/lacZ and pET200/D/lacZ

#### Description

pET100/D/lacZ (8836 bp) and pET200/D/lacZ (8813 bp) are vectors containing a lacZ gene that has been directionally TOPO<sup>®</sup> Cloned into pET100/D-TOPO<sup>®</sup> and pET200/D-TOPO<sup>®</sup>, respectively. In each vector, the *lacZ* gene is cloned in frame with the N-terminal peptide containing the Xpress<sup>™</sup> epitope and the 6xHis tag. The size of the β-galactosidase fusion protein is approximately 121 kDa. The **nucleotide sequence of each vector is available for downloading from Web site (www.invitrogen.com) or by contacting Technical Service (see page 56).** 



	<u>pET100/D/lacZ</u>	<u>pET200/D/lacZ</u>
T7 promoter	209-225	209-225
T7 promoter/priming site	209-228	209-228
<i>lac</i> operator (lacO)	228-252	228-252
Ribosome binding site (RBS)	282-288	282-288
Initiation ATG	297-299	297-299
Polyhistidine (6xHis) region	309-326	309-326
Xpress <sup>™</sup> epitope	366-389	366-389
EK recognition site	375-389	375-389
lacZ ORF	405-3476	405-3476
T7 reverse priming site	3538-3557	3538-3557
T7 transcription termination region	3499-3627	3499-3627
<i>bla</i> promoter	3928-4026	3928-4026
Ampicillin (bla) resistance gene	4027-4887	
Kanamycin resistance gene		4027-4821
pBR322 origin	5032-5705	5009-5682
ROP ORF (complementary strand)	6073-6264	6050-6241
<i>lacI</i> ORF (complementary strand)	7576-8667	7553-8644

#### Map of pET101/D/lacZ

#### Description

The pET101/D/lacZ (8825 bp) vector contains a lacZ gene that has been directionally TOPO<sup>®</sup> Cloned into pET101/D-TOPO<sup>®</sup> in frame with the C-terminal peptide containing the V5 epitope and the 6xHis tag. The size of the  $\beta$ -galactosidase fusion protein is approximately 120 kDa. The nucleotide sequence of the vector is available for downloading from Web site (www.invitrogen.com) or by contacting Technical Service (see page 56).



## Map of pET102/D/lacZ

#### Description

pET102/D/lacZ (9384 bp) is a vector containing a lacZ gene that has been directionally TOPO<sup>®</sup> Cloned into pET102/D-TOPO<sup>®</sup>. The lacZ gene is cloned in frame with the N-terminal HP thioredoxin and the C-terminal peptide tag. The size of the  $\beta$ -galactosidase fusion protein is approximately 133 kDa. The nucleotide sequence of the vector is available for downloading from Web site (www.invitrogen.com) or by contacting Technical Service (see page 56).



	<u>perioz/D/mcz</u>
T7 promoter	209-225
T7 promoter/priming site	209-228
<i>lac</i> operator (lacO)	228-252
Ribosome binding site (RBS)	282-288
His-patch (HP) thioredoxin ORF	298-627
TrxFus forward priming site	607-624
EK recognition site	643-657
lacZ ORF	679-3732
V5 epitope	3769-3810
Polyhistidine (6xHis) region	3820-3837
T7 reverse priming site	3891-3910
T7 transcription termination region	3852-3980
<i>bla</i> promoter	4476-4574
Ampicillin (bla) resistance gene	4575-5435
pBR322 origin	5580-6253
ROP ORF (complementary strand)	6621-6812
<i>lacI</i> ORF (complementary strand)	8124-9215

#### Map of pET151/D/lacZ

#### Description

The pET151/D/*lacZ* (8832 bp) vector contains a *lacZ* gene that has been directionally TOPO<sup>®</sup> Cloned into pET151/D-TOPO<sup>®</sup> in frame with the N-terminal peptide containing the V5 epitope and the 6xHis tag. The size of the  $\beta$ -galactosidase fusion protein is approximately 120 kDa. The nucleotide sequence of the vector is available for downloading from Web site (www.invitrogen.com) or by contacting Technical Service (see page 56).



## Recipes

LB (Luria-Bertani) Medium and Plates	1.0% Tryptone 0.5% Yeast Extract 1.0% NaCl pH 7.0
	<ol> <li>For 1 liter, dissolve 10 g tryptone, 5 g yeast extract, and 10 g NaCl in 950 ml deionized water.</li> </ol>
	2. Adjust the pH of the solution to 7.0 with NaOH and bring the volume up to 1 liter.
	3. Autoclave on liquid cycle for 20 minutes. Allow solution to cool to ~55°C and add antibiotic if needed.
	4. Store at room temperature or at +4°C.
	LB agar plates
	1. Prepare LB medium as above, but add 15 g/L agar before autoclaving.
	2. Autoclave on liquid cycle for 20 minutes.
	3. After autoclaving, cool to ~55°C, add antibiotic and pour into 10 cm plates.
	4. Let harden, then invert and store at $+4^{\circ}$ C, in the dark.
S.O.B. Medium (with Antibiotic)	2% Tryptone 0.5% Yeast Extract 0.05% NaCl 2.5 mM KCl 10 mM MgCl <sub>2</sub>
	1. Dissolve 20 g tryptone, 5 g yeast extract, and 0.5 g NaCl in 950 ml deionized water.
	2. Make a 250 mM KCl solution by dissolving 1.86 g of KCl in 100 ml of deionized water. Add 10 ml of this stock KCl solution to the solution in Step 1.
	3. Adjust pH to 7.5 with 5 M NaOH and add deionized water to 1 liter.
	<ol> <li>Autoclave this solution, cool to ~55°C, and add 10 ml of sterile 1 M MgCl<sub>2</sub>. You may also add antibiotic, if needed.</li> </ol>
	5. Store at +4°C. <b>Medium is stable for only 1-2 weeks.</b>

# **Recipes**, continued

Lysis Buffer	50 400 100 109 0.5 10	50 mM potassium phosphate, pH 7.8 400 mM NaCl 100 mM KCl 10% glycerol 0.5% Triton X-100 10 mM imidazole			
	1.	Prepare 1 M stock solutions of KH <sub>2</sub> PO <sub>4</sub> and K <sub>2</sub> HPO <sub>4</sub> .			
	2.	For 100 ml, dissolve the following reagents in 90 ml of deionized water:			
		0.3 ml KH <sub>2</sub> PO <sub>4</sub> 4.7 ml K <sub>2</sub> HPO <sub>4</sub> 2.3 g NaCl 0.75 g KCl 10 ml glycerol 0.5 ml Triton X-100 68 mg imidazole			
	3.	Mix thoroughly and adjust pH to 7.8 with HCl. Bring the volume to 100 ml.			
	4.	Store at +4°C.			
2X SDS-PAGE Sample Buffer	1.	Combine the following reagents:			
		0.5 M Tris-HCl, pH 6.8 Glycerol (100%) β-mercaptoethanol Bromophenol Blue SDS	2.5 ml 2.0 ml 0.4 ml 0.02 g 0.4 g		
	2.	Bring the volume to 10 ml with sterile water.			
	3.	Aliquot and freeze at -20°C until needed.			
1X SDS-PAGE	1.	Combine the following reag	gents:		
Sample Buffer		0.5 M Tris-HCl, pH 6.8 Glycerol (100%) β-mercaptoethanol Bromophenol Blue SDS	1.25 ml 1.0 ml 0.2 ml 0.01 g 0.2 g		
	2.	Bring the volume to 10 ml v	vith sterile water.		
	3.	Aliquot and freeze at -20°C until needed.			

## **Technical Service**

#### Web Resources



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#### References

- Ausubel, F. M., Brent, R., Kingston, R. E., Moore, D. D., Seidman, J. G., Smith, J. A., and Struhl, K. (1994) *Current Protocols in Molecular Biology*, Greene Publishing Associates and Wiley-Interscience, New York
- Carrington, J. C., and Dougherty, W. G. (1988) A Viral Cleavage Site Cassette: Identification of Amino Acid Sequences Required for Tobacco Etch Virus Polyprotein Processing. Proc. Natl. Acad. Sci. USA *85*, 3391-3395
- Cheng, C., and Shuman, S. (2000) Recombinogenic Flap Ligation Pathway for Intrinsic Repair of Topoisomerase IB-Induced Double-Strand Breaks. Mol. Cell. Biol. 20, 8059-8068
- Deutscher, M. P. (ed)) (1990) *Guide to Protein Purification* Vol. 182. Methods in Enzymology. Edited by Abelson, J. N., and Simon, M. I., Academic Press, San Diego, CA.
- Dougherty, W. G., Carrington, J. C., Cary, S. M., and Parks, T. D. (1988) Biochemical and Mutational Analysis of a Plant Virus Polyprotein Cleavage Site. EMBO J. 7, 1281-1287
- Dubendorff, J. W., and Studier, F. W. (1991) Controlling Basal Expression in an Inducible T7 Expression System by Blocking the Target T7 Promoter with lac Repressor. J. Mol. Biol. 219, 45-59
- Gold, L. (1988) Posttranscriptional Regulatory Mechanisms in Escherichia coli. Ann. Rev. Biochem. 57, 199-233
- Grunberg-Manago, M. (1999) Messenger RNA Stability and its Role in Control of Gene Expression in Bacteria and Phages. Annu. Rev. Genet. 33, 193-227
- Holmgren, A. (1985) Thioredoxin. Ann. Rev. Biochem. 54, 237-271
- Katti, S. K., LeMaster, D. M., and Eklund, H. (1990) Crystal Structure of Thioredoxin from *E. coli* at 1.68 Angstroms Resolution. J. Mol. Biol. 212, 167-184
- Kido, M., Yamanaka, K., Mitani, T., Niki, H., Ogura, T., and Hiraga, S. (1996) RNase E Polypeptides Lacking a Carboxyl-terminal Half Suppress a *mukB* mutation in *Escherichia coli*. J. Bacteriol. 178, 3917-3925
- LaVallie, E. R., DiBlasio, E. A., Kovacic, S., Grant, K. L., Schendel, P. F., and McCoy, J. M. (1993) A Thioredoxin Gene Fusion Expression System That Circumvents Inclusion Body Formation in the *E. coli* Cytoplasm. Bio/Technology *11*, 187-193
- Lindner, P., Bauer, K., Krebber, A., Nieba, L., Kremmer, E., Krebber, C., Honegger, A., Klinger, B., Mocikat, R., and Pluckthun, A. (1997) Specific Detection of His-tagged Proteins With Recombinant Anti-His Tag scFv-Phosphatase or scFv-Phage Fusions. BioTechniques 22, 140-149
- Lopez, P. J., Marchand, I., Joyce, S. A., and Dreyfus, M. (1999) The C-terminal Half of RNase E, Which Organizes the *Escherichia coli* Degradosome, Participates in mRNA Degradation but not rRNA Processing *in vivo*. Mol. Microbiol. *33*, 188-199
- Lu, Z., DiBlasio-Smith, E. A., Grant, K. L., Warne, N. W., LaVallie, E. R., Collins-Racie, L. A., Follettie, M. T., Williamson, M. J., and McCoy, J. M. (1996) Histidine Patch Thioredoxins. J. Biol. Chem. 271, 5059-5065
- Miller, J. H. (1992) A Short Course in Bacterial Genetics: A Laboratory Manual and Handbook for Escherichia coli and Related Bacteria, Cold Spring Harbor Laboratory Press, Plainview, New York
- Rosenberg, A. H., Lade, B. N., Chui, D.-S., Lin, S.-W., Dunn, J. J., and Studier, F. W. (1987) Vectors for Selective Expression of Cloned DNAs by T7 RNA Polymerase. Gene *56*, 125-135
- Shuman, S. (1991) Recombination Mediated by Vaccinia Virus DNA Topoisomerase I in *Escherichia coli* is Sequence Specific. Proc. Natl. Acad. Sci. USA *88*, 10104-10108
- Shuman, S. (1994) Novel Approach to Molecular Cloning and Polynucleotide Synthesis Using Vaccinia DNA Topoisomerase. J. Biol. Chem. 269, 32678-32684
## **References**, continued

- Southern, J. A., Young, D. F., Heaney, F., Baumgartner, W., and Randall, R. E. (1991) Identification of an Epitope on the P and V Proteins of Simian Virus 5 That Distinguishes Between Two Isolates with Different Biological Characteristics. J. Gen. Virol. 72, 1551-1557
- Studier, F. W., and Moffatt, B. A. (1986) Use of Bacteriophage T7 RNA Polymerase to Direct Selective High-Level Expression of Cloned Genes. J. Mol. Biol. 189, 113-130
- Studier, F. W., Rosenberg, A. H., Dunn, J. J., and Dubendorff, J. W. (1990) Use of T7 RNA Polymerase to Direct Expression of Cloned Genes. Meth. Enzymol. 185, 60-89
- Wallace, B. J., and Kushner, S. R. (1984) Genetic and Physical Analysis of the Thioredoxin (*trx*A) Gene of *Escherichia coli* K-12. Gene 32, 399-408

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