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6 x 0.2 ml/tube

Lot: 18

Store at -80°C

**CAUTION:** This product contains DMSO, a hazardous material. Review the MSDS before handling.

**Description:** Chemically competent *E. coli* cells suitable for high efficiency transformation and protein expression.

#### Features:

- Transformation efficiency: 0.6–1 x 10<sup>9</sup> cfu/µg pUC19
- Enhanced BL21 derivative for T7 expression
- T7 RNA Polymerase in the *lac* operon no lambda prophage
- Deficient in proteases Lon and OmpT
- Resistant to phage T1 (fhuA2)
- Does not restrict methylated DNA (McrA-, McrBC-, EcoBr-m-, Mrr-)
- B Strain
- Free of animal products

#### **Reagents Supplied:**

6 x 0.2 ml/tube of chemically competent T7 Express Competent *E. coli* cells (Store at -80°C)

25 ml of SOC Outgrowth Medium (Store at room temperature) 0.025 ml of 50 pg/µl pUC19 Control DNA (Store at -20°C)

### **Quality Control Assays**

**Transformation Efficiency:** 100 pg of pUC19 plasmid DNA was used to transform one tube of T7 Express Competent *E. coli* following the high efficiency protocol provided.  $0.6-1 \times 10^9$  colonies formed/µg after an overnight incubation on LB-ampicillin plates at 37°C.

Untransformed cells were also tested for resistance to phage  $\phi 80$ , a standard test for resistance to phage T1, and sensitivity to ampicillin, chloramphenicol, kanamycin, spectinomycin, streptomycin and tetracycline.

# **High Efficiency Transformation Protocol**

- Thaw a tube of T7 Express Competent E. coli cells on ice until the last ice crystals disappear. Mix gently and carefully pipette 50 µl of cells into a transformation tube on ice.
- Add 1–5 µl containing 1 pg–100 ng of plasmid DNA to the cell mixture. Carefully flick the tube 4–5 times to mix cells and DNA. Do not vortex.
- 3. Place the mixture on ice for 30 minutes. Do not mix.
- 4. Heat shock at exactly 42°C for exactly 10 seconds. Do not mix.
- 5. Place on ice for 5 minutes. Do not mix.
- 6. Pipette 950 µl of room temperature SOC into the mixture.
- 7. Place at 37°C for 60 minutes. Shake vigorously (250 rpm) or rotate.
- 8. Warm selection plates to 37°C.
- Mix the cells thoroughly by flicking the tube and inverting, then perform several 10-fold serial dilutions in SOC.

STORAGE AND HANDLING: Competent cells should be stored at -80°C. Storage at -20°C will result in a significant decrease in transformation efficiency. Cells lose efficiency whenever they are warmed above -80°C, even if they do not thaw.

 Spread 50–100 μl of each dilution onto a selection plate and incubate overnight at 37°C. Alternatively, incubate at 30°C for 24–36 hours or at 25°C for 48 hours.

#### **5 Minute Transformation Protocol**

A shortened transformation protocol resulting in approximately 10% efficiency compared to the standard protocol may be suitable for applications where a reduced total number of transformants is acceptable. Follow the High Efficiency Transformation Protocol above with the following changes:

- 1. Steps 3 and 5 are reduced to 2 minutes.
- Omit outgrowth (step 7) completely for ampicillin-resistant plasmids or reduce the outgrowth time for other selective media as appropriate.

## **Protocol for Expression Using T7 Express**

- Transform expression plasmid into T7 Express. Plate on antibiotic selection plates and incubate overnight at 37°C.
- 2. Resuspend a single colony in 10 ml liquid culture with antibiotic.
- 3. Incubate at 37°C until OD<sub>600</sub> reaches 0.4–0.8.
- 4. Induce with 40  $\mu$ I of a 100 mM stock of IPTG (final concentration of 0.4 mM) and induce for 2 hours at 37°C.
- Check for expression either by Coomassie stained protein gel, Western Blot or activity assay. Check expression in both the total cell extract (soluble + insoluble) and the soluble fraction alone.
- 6. For large scale, inoculate 1 L of liquid medium (with antibiotic) with a freshly grown colony or 10 ml of freshly grown culture. Incubate at  $37^{\circ}\text{C}$  until OD $_{600}$  reaches 0.4–0.8. Add IPTG to 0.4 mM. Induce 2 hours at  $37^{\circ}\text{C}$  or  $15^{\circ}\text{C}$  overnight.

## **Transformation Protocol Variables**

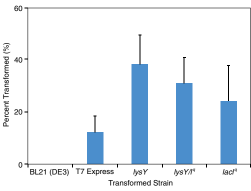
**Thawing:** Cells are best thawed on ice and DNA added as soon as the last bit of ice in the tube disappears. Cells can also be thawed by hand, but warming above 0°C will decrease the transformation efficiency.

**Incubation of DNA with Cells on Ice:** For maximum transformation efficiency, cells and DNA should be incubated together on ice for 30 minutes. Expect a 2-fold loss in transformation efficiency for every 10 minutes this step is shortened.

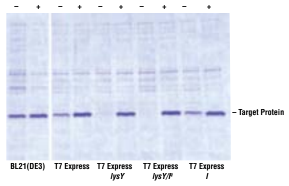
**Heat Shock:** Both the temperature and the timing of the heat shock step are important and specific to the transformation volume and vessel. Using the transformation tube provided, 10 seconds at 42°C is optimal.

**Outgrowth:** Outgrowth at 37°C for 1 hour is best for cell recovery and for expression of antibiotic resistance. Expect a 2-fold loss in transformation efficiency for every 15 minutes this step is shortened. SOC gives 2-fold higher transformation efficiency than LB medium; and incubation with shaking or rotating the tube gives 2-fold higher transformation efficiency than incubation without shaking.

**Plating:** Selection plates can be used warm or cold, wet or dry without significantly affecting the transformation efficiency. However, warm, dry plates are easier to spread and allow for the most rapid colony formation.



**Transformation of a toxic mammalian clone into E. coli hosts.** A T7 expression plasmid and the same plasmid containing a gene encoding a toxic mammalian protein were transformed into each host. Comparison of the relative transformation efficiencies demonstrates that the T7 Express hosts provide the levels of control necessary for transformation of potentially toxic clones. BL21(DE3) could not be transformed with the toxic clone.



**T7-controlled expression of a non-toxic protein in E. coli hosts.** A T7 expression plasmid containing a gene encoding an E. coli protein was transformed into each host, grown to 0.6 OD and induced for 3 hours. Comparison of soluble extracts from uninduced (-) and induced (+) cells shows superior control of basal expression in the T7 Express hosts while maintaining high levels of induced expression.

## **DNA Contaminants to Avoid**

Contaminant	Removal Method
Detergents	Ethanol precipitate
Phenol	Extract with chloroform and ethanol precipitate
Ethanol or Isopropanol	Dry pellet before resuspending
PEG*	Column purify or phenol/chloroform extract and ethanol precipitate
DNA binding proteins* (e.g. Ligase)	Column purify or phenol/chloroform extract and ethanol precipitate

<sup>\*</sup>Ideally, DNA for transformation should be purified and resuspended in water or TE. However, up to 10 µl of DNA directly from a ligation mix can be used with only a twofold loss of transformation efficiency. Where it is necessary to maximize the number of transformants (e.g. a library), a purification step, either a spin column or phenol/ chloroform extraction and ethanol precipitation should be added.

### **Troubleshooting T7 Protein Expression**

**No colonies or no growth in liquid culture:** Even though T7 expression is tightly regulated, there may be a low level of basal expression in the T7 Express host. If toxicity of the expressed protein is likely, transformation of the expression plasmid should be carried out in one of the following strains:

- T7 Express I<sup>q</sup>: over-expression of the LacI repressor reduces basal expression of the T7 RNA polymerase
- T7 Express IysY: IysY produces mutant T7 Iysozyme which binds to T7 RNA polymerase, reducing basal expression of the target protein. Upon induction, newly made T7 RNA polymerase titrates out the Iysozyme and results in expression of the target protein
- T7 Express *lysY/lq* combines both above effects.

Incubation at 30°C or room temperature may also alleviate toxicity issues. In addition, check antibiotic concentration (test with control plasmid).

**No protein visible on gel or no activity:** Check for toxicity - no protein may mean the cells have eliminated or deleted elements in the expression plasmid.

- Culture cells for protein induction. Just before induction, plate a sample on duplicate plates with and without antibiotic selection. If toxicity is an issue, there will be a significant difference between the number of colonies on the plates. Fewer colonies will be seen on plates containing antibiotic (indicating that the plasmid has been lost) compared to plates without antibiotic.
- If toxicity is the problem test the above I<sup>n</sup> and IysY hosts to reduce basal level expression.

**Induced protein is insoluble:** Check for insolubility - this is important because T7 expression often leads to very high production of protein that can result in the target protein becoming insoluble. Potential solutions for this are:

- Induce at lower temperatures (as low as 15°C overnight)
- Reduce IPTG concentration to 0.1 mM and 0.01 mM
- Induce for less time (as little as 15 minutes)
- Induce earlier in growth  $(OD_{600} = 0.3 \text{ or } 0.4)$

#### Solutions/Recipes

SOB:		SOC:	
2% Ve	getable peptone (or Tryptone)	SOB + 20	) mM Glucose
0.5% Ye	ast Extract	LB agar:	
2.5 mM KC			Tryptone
10 mM M	gCl <sub>a</sub>	0.5%	Yeast Extract
10 mM M	gSÓ,	0.17 M	NaCl
	4	1.5%	Aga

#### **Antibiotics for Plasmid Selection**

Antibiotic	Working Concentration
Ampicillin	100 μg/ml
Carbenicillin	100 μg/ml
Chloramphenicol	33 μg/ml
Kanamycin	30 μg/ml
Streptomycin	25 μg/ml
Tetracycline	15 μg/ml

**Genotype:** fhuA2 lacZ::T7 gene1 [lon] ompT gal sulA11  $R(mcr-73::miniTn10--Tet^s)2$  [dcm]  $R(zgb-210::Tn10--Tet^s)$  endA1  $\Delta(mcrC-mrr)114::IS10$ 

## **Strain Properties**

The properties of this strain that contribute to its usefulness as a protein expression strain are described below. The genotypes underlying these properties appear in parentheses.

T7 RNA Polymerase (*lacZ::T7 gene1*): T7-Express has the T7 RNA polymerase gene inserted into the lac operon on the *E. coli* chromosome and is expressed under the control of the *lac* promoter. This configuration provides controlled induction of the polymerase and consequently, inducible control of transcription of genes downstream of the T7 promoter. This system provides potential advantages over strains such as BL21(DE3), that carry the T7 RNA polymerase on a lysogenic prophage. Although λDE3 is normally dormant in the host chromosome, the induction of the SOS cascade can occur as the result of expressing proteins that damage the *E. coli* chromosome, either directly or indirectly. This may lead to cell lysis.

Protease Deficient ([lon] ompT): E. coli B strains are "naturally" deficient in the lon protease which in K-12 strains serves to degrade misfolded proteins and to prevent some cell cycle-specific proteins from accumulating. The OmpT protease resides at the surface of wild type E. coli in both K-12 and B strains, presumably helping the cells to derive amino acids from their external environment. Cells deficient in both these proteases are much more amenable to the production of proteins from cloned genes. Mutations of other genes can help to ameliorate the sometimes-deleterious effects of these protease defects (e.g. sulA, below).

Recovery from DNA Damage (sulA11): E. coli cells can tolerate a substantial amount of chronic DNA damage as long as repair is allowed to proceed. This capacity is compromised if the cells are unable to divide following repair. In lon cells, SulA, a cell division inhibitor, accumulates and causes cells to become hypersensitive to DNA damage. The sulA mutation introduced into the T7 Express strain allows cells to divide more normally in the absence of Lon protease.

Endonuclease I Deficient (endA1): The periplasmic space of wild type E. coli cells contains a nonspecific endonuclease. Extreme care must be taken to avoid degradation of plasmids prepared from these cells. The endA mutation deletes this endonuclease and can significantly improve the quality of plasmid preparations.

Restriction Deficient ( $\Delta(mcrC-mrr)114::IS10$ ): Wild type *E. coli* B strains carry a Type I restriction endonuclease which cleaves DNA with the site TGA(N8)TGCT. While *E. coli* DNA is protected from degradation by a cognate methyl-transferase, foreign DNA will be cut at these sites. The deletion described above eliminates both the methylase and the endonuclease.

Methyl Restriction Deficient ( $\Delta(mcrC-mrr)114::IS10$  and  $R(mcr-73::miniTn10-Tet^s)2$ ): E. coli has a system of enzymes encoded by mcrA, mcrBC and mrr which will cleave DNA with methylation patterns found in higher eukaryotes, as well as some plant and bacterial strains. All three Mcr enzymes and Mrr have been inactivated in T7 Express allowing the introduction of eukaryotic DNA of genomic origin (e.g. primary libraries) if desired.

T1 Phage Resistant (*fhuA2*): T1, an extremely virulent phage requires the *E. coli* ferric hydroxamate uptake receptor for infectivity. Deletion of this gene confers resistance to this type of phage, but does not significantly affect the transformation or growth characteristics of the cell.

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