



Mix & Go! E. coli Transformation Kit and **Buffer Set**

For generating Mix & Go! chemically competent cells from most E. coli lab strains.

Highlights

- Easy 3 Step Protocol: Produce reliable chemically competent E. coli in less than 45 minutes.
- · Simple 20 sec Transformation: No heat shock! Just add DNA and spread.
- High Transformation Efficiencies: Achieve 108-109 transformants per µg of plasmid DNA.

Catalog Numbers: T3001, T3002



Scan with your smart-phone camera to view the online protocol/video.

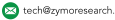






Table of Contents

Product Contents	. 01
Product Description	. 02
Protocol	. 04
Preparation of Mix & Go! Cells	. 04
Fast Transformation of Competent Cells	. 05
Appendix	. 06
Notes for High Efficiency Transformation.	. 06
Media Recipes	. 07
References	. 07
Ordering Information	. 08
Complete Your Cloning Workflow	. 09
Notes	. 10
Guarantee	. 13

Revised on: 8/21/2024

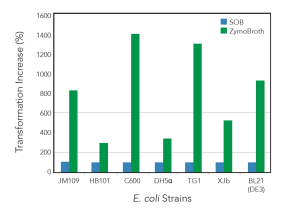
Product Contents

	Mix & Go! E. coli Transformation Kit ¹ T3001 (20 ml)	Mix & Go! E. coli Transformation Buffer Set ² T3002 (60 ml)	Storage
ZymoBroth™	200 ml	(Not Included)	Room Temp.
Wash Buffer (2X)	10 ml	30 ml	0-8°C
Competent Buffer (2X)	10 ml	30 ml	0-8°C
Dilution Buffer	20 ml	60 ml	0-8°C
Instruction Manual	1	1	-

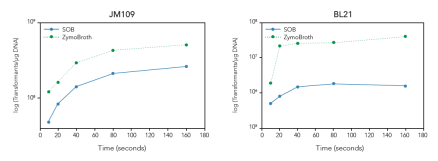
¹ Includes all buffers for making up to 20 ml *Mix & Go! E. coli.* **ZymoBroth™** growth medium is included. ² Includes all buffers for making up to 60 ml *Mix & Go! E. coli.* **ZymoBroth™** growth medium is <u>not</u> included.

Product Description

The *Mix & Go! E. coli* Transformation Kit and Buffer Set are simple methods for generating *Mix & Go!* chemically competent *E. coli* for rapid, reliable, and highly efficient DNA transformation. The methods eliminate the requirement of heat-shocking and related procedures. Instead, transformation can be performed by adding DNA to prepared *Mix & Go!* cells and spreading the mixture directly to a culture plate. Transformation efficiencies typically range from 10⁸–10⁹ transformants/µg of pUC19 DNA but can vary depending on the strain of *E. coli*. Most *E. coli* strains respond well to the *Mix & Go!* preparation method and demonstrate fast transformation kinetics (see figures below).



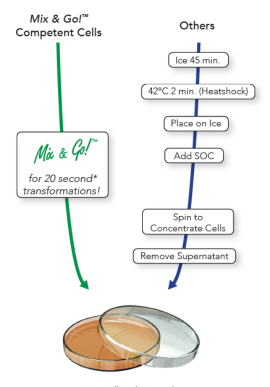
Transformation efficiencies of strains generated with ZymoBroth™ and SOB media. ZymoBroth™ dramatically increases the transformation efficiencies of a broad range of *E. coli* strains. Generally, ZymoBroth™ enhances transformation efficiencies better for difficult-to-transform strains.



Transformation kinetics. *Mix* & *Go!*™ *E. coli* prepared with ZymoBroth™ display fast transformation kinetics and high transformation efficiencies. pUC19 DNA was used for transformation and the data are the averages of three individual experiments.

The procedures are easy. Simply culture the *E. coli* strain of your choice in **ZymoBroth**[™] medium (or SOB), wash and then resuspend the cells in the provided uniquely formulated buffers. The cells are now ready for transformation!

The *Mix & Go! E. coli* Transformation Kit (T3001) includes all buffers for making up to 20 ml *Mix & Go! E. coli* from your favorite lab strains. **ZymoBroth™** growth medium is included. The *Mix & Go! E. coli* Transformation Buffer Set (T3002) includes all buffers for making up to 60 ml *Mix & Go! E. coli* from your favorite lab strains, but **ZymoBroth™** growth medium is <u>not</u> included.



*Ampicillin selection only

Protocol

The following procedure is for a 50 ml *E. coli* culture in **ZymoBroth™** (supplied with T3001 only) or SOB medium (see appendix for recipe); however, the volume can be adjusted according to your specific requirements.

Preparation of Mix & Go! Cells

 Use 0.5 ml of fresh, overnight *E. coli* culture grown in LB to inoculate 50 ml **ZymoBroth™** or SOB medium in a 500 ml culture flask. Shake culture vigorously (150 - 250 rpm) at 18-26°C until the OD_{600nm} is 0.4 - 0.6.

Buffer Preparation Prior to Harvesting the Cells...

- ✓ The Wash and Competent Buffers are provided as 2X stock solutions. They need to be diluted to 1X by adding an equal amount of Dilution Buffer.
- ✓ To prepare 5 ml of 1X Wash Buffer: Add 2.5 ml Dilution Buffer and 2.5 ml of 2X Stock Wash Buffer.
- ✓ To prepare 5 ml of 1X Competent Buffer: Add 2.5 ml Dilution Buffer and 2.5 ml of 2X Stock Competent Buffer.
- ✓ Please keep these freshly prepared 1X Buffers ice cold. These 1X Buffers are good for 2 days at 0-25°C.

Important! Each step of the following procedure should be done on ice or at 0-4°C.

2. Transfer the culture from Step 1 to ice. After 10 minutes, pellet the cells by centrifugation at 3,000 - 3,700 rpm (i.e., 1,600 - 2,500 x g) for 10 minutes at 0-4°C.

- 3. Remove the supernatant and resuspend the cells gently in 5 ml ice-cold 1X **Wash Buffer**. Re-pellet the cells as in Step 2.
- 4. Completely remove the supernatant and gently resuspend the cells in 5 ml ice-cold 1X **Competent Buffer**.
- 5. Aliquot (on ice) 0.1-0.2 ml of the cell suspension into sterile microcentrifuge tubes. Cells are now ready for transformation with DNA or can be stored below -70°C for transformation at a later time.

Note: The prepared competent cells are referred to as "*Mix & Go*" in the procedures that follow.

Fast Transformation of Mix & Go Competent Cells*

- 1. Add 1-5 μ l plasmid DNA to a tube of thawed *Mix & Go* cells on ice, mix gently for a few seconds (try to keep the added volume of DNA less than 5% of the total).
- 2. Spread 50-100 µl of the mixture onto a pre-warmed (37°C) culture plate containing Ampicillin. Incubate the plate at the appropriate temperature (*e.g.*, 37°C) for the colonies to grow.

^{*}For Ampicillin selection only. For selection with other antibiotics, see notes Section 4 on next page.

Appendix

Notes for High Efficiency Transformation

1. E. coli Strains

Different *E. coli* strains vary in their ability to be transformed with DNA. Strains like JM109, C600, TG1, DH5α, XL10 Gold, and BL21 and its derivatives typically yield the best results when prepared with the *Mix* & *Go! E. coli* Transformation Kit.

2. Incubation Time

The *Mix & Go!* procedure can be used for most transformations using Ampicillin selection and not requiring outgrowth (see Section 4 below). The highest transformation efficiencies can be obtained by incubating *Mix & Go* cells with DNA on ice for 2-5 minutes prior to plating.

3. Prewarming Culture Plates

Chilled plates will decrease *Mix & Go* cell transformation efficiency. It is recommended that culture plates be pre-warmed to >20°C (preferably 37°C) prior to plating.

4. Addition of SOC Medium to Transformation Mixtures (Outgrowth) When selecting with Kanamycin, Tetracycline, etc., an outgrowth performed in SOC medium is required for efficient transformation. In most cases, this step can be omitted when selecting with Ampicillin. After the transformation mixture has incubated on ice for 5-10 min, add 4 volumes of SOC (400 µl of SOC to 100 µl of transformation mixture) and incubate for 1 hour at 37°C with gentle shaking at 200-300 rpm. Afterwards, spread the mixture directly onto pre-warmed culture plates. Reducing agents [e.g., DTT (Dithiothreitol) and 2-ME (β-mercaptoethanol)] are not required for this procedure.

5. Culture Conditions

E. coli cells become highly competent when cultured at 20-33°C prior to preparation. Higher temperatures (*i.e.*, 33°C-37°C) can decrease the transformation efficiency 2 to 10-fold. Also, cells can be harvested at lower densities (e.g., OD_{600nm} 0.2-0.4) and resuspended in smaller volumes (e.g., 1-3 ml vs. 5 ml as recommended in the standard procedure). Cells harvested at lower densities (OD_{600nm} 0.2-0.6) are usually "more competent" than those cells harvested at higher densities ($OD_{600nm} > 0.6$).

Media Recipes

Although SOB has traditionally been used for *Mix & Go* cell preparation, **ZymoBroth™** is now the medium of choice for the generation of *Mix & Go E. coli* that exhibit fast and highly efficient transformation kinetics. However, SOB can still be used in both *Mix & Go! E. coli* Transformation **Kit and Buffer Set** procedures.

SOB Recipe: (1 Liter)

Mix the following ingredients:

- √ 20 g Bacto-tryptone
- ✓ 0.58 g NaCl (or 2 ml of 5M NaCl)
 - √ 10 ml 1M MgCl2
- ✓ 5 g Yeast extract
- ✓ 0.19 g KCl (or 0.5 ml 1M KCl)
- √ 10 ml 1M MgSO4

Add ddH2O to a total volume of 1 liter.

Adjust pH to 6.0-7.0 with NaOH.

Autoclave at 10 psi for 15-20 minutes.

SOC Recipe: (100 ml)

Add 1 ml of a 2 M filter-sterilized glucose solution or 2 ml of 20% (w/v) glucose solution to 100 ml of SOB medium.

References

- Sheridan, P. et al. Phylogenetic Analysis of Anaerobic Psychrophilic Enrichment Cultures Obtained from a Greenland Glacier Ice Core, Appl. Envir. Microbiol., Apr 2003; 69: 2153 – 2160.
- 2. Yokobayashi, Y. et al. From the Cover: Directed evolution of a genetic circuit, PNAS, Dec 2002; 99: 16587 16591.
- 3. Trent, J. et al. A Ubiquitously Expressed Human Hexacoordinate Hemoglobin, J. Biol. Chem., May 2002; 277: 19538 19545.
- Mourez, M. et al. Mapping dominant-negative mutations of anthrax protective antigen by scanning mutagenesis, PNAS, Nov 2003; 100: 13803 – 13808.

Ordering Information

Product Description	Catalog No.	Size
Mix & Go! E. coli Transformation Kit Includes all buffers for making up to 20 ml Mix & Go! E. coli from your favorite lab strains. ZymoBroth™ growth medium is included.	T3001	Prepare up to 20 ml Competent Cells
Mix & Go! E. coli Transformation Buffer Set Includes all buffers for making up to 60 ml Mix & Go! E. coli from your favorite lab strains. ZymoBroth™ growth medium is not included.	T3002	Prepare up to 60 ml Competent Cells

Individual Kit Components	Catalog No.	Amount
ZymoBroth™	M3015-100 M3015-500	100 ml 500 ml
Wash Buffer (2X Stock)	T3001-2-10 T3001-2-30	10 ml 30 ml
Competent Buffer (2X Stock)	T3001-3-10 T3001-3-30	10 ml 30 ml
Dilution Buffer	T3001-4-20 T3001-4-60	20 ml 60 ml

Complete Your Cloning Workflow

✓ Transfection-grade plasmid DNA from a miniprep

ZymoPURE™ Plasmid Miniprep	Size	Catalog No.
ZymoPURE™ Plasmid Miniprep Kit	10 Preps. 50 Preps. 100 Preps. 400 Preps. 800 Preps.	D4208T D4309 D4210 D4211 D4212
ZymoPURE™ 96 Plasmid Miniprep Kit	2 x 96 Preps. 4 x 96 Preps.	D4214 D4215

✓ 18 Minute Endotoxin-Free Midi & Maxipreps

ZymoPURE™ II Plasmid Prep Kits	Size	Catalog No.
ZymoPURE™ II Plasmid Midiprep Kit	25 Preps. 50 Preps.	D4200 D4201
ZymoPURE™ II Plasmid Maxiprep Kit	10 Preps. 20 Preps.	D4202 D4203
ZymoPURE™ II Plasmid Gigaprep Kit	5 Preps.	D4204

✓ Simple 20 second High Efficiency Transformations

Mix & Go! Competent Cells	Size	Catalog No.
DH5α	10 x 100 μl aliquots 96 x 50 μl aliquots 96 x 50 μl aliquots PCR Plate	T3007 T3009 T3010
JM109	10 x 100 μl aliquots 96 x 50 μl aliquots	T3003 T3005
Zymo10B	10 x 100 μl aliquots 96 x 50 μl aliquots	T3019 T3020
HB101	10 x 100 μl aliquots 96 x 50 μl aliquots	T3011 T3013
TG1	10 x 100 μl aliquots	T3017

✓ Recover ultra-pure highly concentrated DNA from PCR & other sources

DNA Clean & Concentrator™	Size	Catalog No.
DNA Clean & Concentrator™-5	50 Preps. 200 Preps.	D4003 D4004
ZR-96 DNA Clean-Up Kit™	2 x 96 Preps. 4 x 96 Preps.	D4017 D4018

✓ Rapid extraction of ultra-pure DNA from agarose gels

Zymoclean Gel DNA Recovery [™]	Size	Catalog No.
Zymoclean™ Gel DNA Recovery Kit	50 Preps. 200 Preps.	D4001 D4002

Notes			

Notes			

Notes			



100% satisfaction guarantee on all Zymo Research products, or your money back.

Zymo Research is committed to simplifying your research with quality products and services. If you are dissatisfied with this product for any reason, please call 1(888) 882-9682.

Integrity of kit components is guaranteed for up to one year from date of purchase. Reagents are routinely tested on a lot-to-lot basis to ensure they provide the highest performance and reliability.

This product is for research use only and should only be used by trained professionals. It is not for use in diagnostic procedures. Some reagents included with this kit are irritants. Wear protective gloves and eye protection. Follow the safety guidelines and rules enacted by your research institution or facility.

[™] Trademarks of Zymo Research Corporation DH5α is a trademark of Life Technologies. XL10 Gold is a trademark of Stratagene.



The **BEAUTY** of **SCIENCE** is to Make Things **SIMPLE**®