



# SS320 Phage Display ElectroCompetent Cells

# Manual

Catalog #	1173-12	1173-24	1274-24	1274-48
Package Size	6x50 μl	12x50 μl	6x100 μl	12x100 μl



# Important!

# -80°C Storage Required

- \* Immediately inspect packages
- \* Freeze upon receipt

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# **SSS320** Phage Display ElectroCompetent Cells

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## **Description:**

Intact Genomics (ig®) SS320 (MC1061F\') phage display electrocompetent cells are suitable for protein expression, general cloning, blue/white screening, M13 phage work and phage display protein expression.

## **Product Components:**

- ig® SS320 phage display electrocompetent cells
- pUC19 Control
- · Recovery medium

#### **Storage:**

ig® SS320 phage display electrocompetent cells: -80 ºC

pUC19 control DNA: -20 ºC

Recovery medium: 4 ºC

#### **Genomic Features:**

Intact Genomics SS320 phage display electrocompetent cells have the following features:

- $>5 \times 10^{10}$  cfu/µg efficiency with electroporation.
- Non-amber suppressor strain (sometimes called MC1061F\')

# **Genotype:**

F'[proAB+ lacIq lacZ $\Delta$ M15 Tn10 (tetr)] hsdR mcrA0 araD139  $\Delta$ (araA-leu)7697  $\Delta$ lacX74 spoT1 galK  $\lambda$ e14- galE rpsL thi

## **Quality Control:**

Transformation efficiency is tested by using the pUC19 control DNA supplied with the kit and using the protocol given below. Transformation efficiency should be >5 x  $10^{10}$  CFU/µg pUC19 DNA. Untransformed cells are tested for appropriate antibiotic sensitivity.



#### **General Guidelines:**

Follow these guidelines when using Intact Genomics SS320 phage display electrocompetent cells:

- Handle competent cells gently as they are highly sensitive to changes in temperature or mechanical lysis caused by pipetting.
- Thaw competent cells on ice and transform cells immediately following thawing. After adding DNA, mix by tapping the tube gently. Do not mix cells by pipetting or vortexing.

**Note:** A high-voltage electroporation apparatus such as Bio-Rad Gene Pulser II #165-2105, capable of generating field strengths of 16 kV/cm is required.

#### **Calculation of Transformation Efficiency:**

Transformation Efficiency (TE) is defined as the number of colony forming units (cfu) produced by transforming 1µg of plasmid into a given volume of competent cells.

TE = Colonies/ $\mu$ g/Dilution

Transform 1  $\mu$ l of (10 pg/ $\mu$ l) pUC19 control plasmid into 50  $\mu$ l of cells, add 950  $\mu$ l of Recovery Medium. Dilute 10  $\mu$ l of this in 990  $\mu$ l of Recovery Medium and plate 50  $\mu$ l. Count the colonies on the plate the next day. If you count 100 colonies, the TE is calculated as follows:

Colonies = 100  $\mu$ g of DNA = 0.00001 Dilution = 50/1000 x 10/1000 = 0.0005 TE = 100/.00001/.0005 = 2.0x10<sup>10</sup>



#### **Transformation Protocol:**

Use this procedure to transform ig® SS320 phage display electrocompetent cells. Do not use these cells for chemical transformation.

- 1) Place sterile cuvettes and microcentrifuge tubes on ice.
- 2) Remove competent cells from the -80 °C freezer and thaw completely on wet ice (10-15 minutes).
- 3) Aliquot 1  $\mu$ l (1 pg-10 ng) of DNA to the chilled microcentrifuge tubes on ice.
- 4) When the cells are thawed, add 25  $\mu$ l of cells to each DNA tube on ice and mix gently by tapping 4-5 times. For the pUC19 control, add 1  $\mu$ l of (10 pg/ $\mu$ l) DNA to the 25  $\mu$ l of cells on ice. Mix well by tapping. Do not pipette up and down or vortex to mix, this can harm cells and decrease transformation efficiency.
- 5) Pipette 26 µl of the cell/DNA mixture into a chilled electroporation cuvette without introducing bubbles. Quickly flick the cuvette downward with your wrist to deposit the cells across the bottom of the well and then electroporate.
- 6) Immediately add 974  $\mu$ l of Recovery Medium or any other medium of choice to the cuvette, pipette up and down three times to re-suspend the cells. Transfer the cells and Recovery Medium to a culture tube.
- 7) Incubate tubes at 37 °C for 1 hour at 210 rpm.
- 8) Dilute the cells as appropriate then spread 20-200 μl cells onto a pre-warmed selective plate. For the pUC19 control, plate 50 μl of diluted transformants onto an LB plate containing 100 μg/ml ampicillin. Use sterilized spreader or autoclaved ColiRoller™ plating beads to spread evenly.
- 9) Incubate the plates overnight at 37 °C.



#### **Related Products:**

- TG1 Phage Display ElectroComp. Cells (Cat.# 1264-24)
- ig® 5-Alpha Chemically Comp. Cells (Cat.# 1031-12)
- T4 DNA Ligase (Cat.# 3212)
- i7<sup>®</sup> High Fidelity DNA Polymerase (Cat.# 3254)

# **Ordering Information:**

- Order online within the USA. Place orders on www.intactgenomics.com using our secure Shopping Cart.
- Order by email, phone, or fax.

Email: sales@intactgenomics.com

Phone: (314) 942-3655 | Toll-free: 855-835-7172 | Fax: (314) 942-3656

• Order via our distributors.



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