

TransforMax EC100 elektrokompente E. coli

Lucigen



10 x 100 µl

TransforMAX™

Artikel-Nr.: 190055 | Lucigen | Herstellernummer: EC10010

370,80 € *

*zzgl. MwSt. [zzgl. Versandkosten](#)

Beschreibung

Produkttyp: Kompetente Zellen

Verpackung: 10 x 100 µl

Applications

- Routine cloning of DNA up to 200 kb.

The highly versatile TransforMax™ EC100™ *E. coli* competent cells are ideal for most cloning applications. The cells provide very high transformation efficiency when tested against a wide range of supercoiled DNAs as well as DNA directly from a ligation reaction (Table 1).

Benefits

- High transformation efficiency with clones of all sizes, including BAC clones (Table 1).
- *lacZΔM15* for blue/white screening of recombinants.
- Restriction minus [*mcrA*, Δ(*mrr-hsdRMS-mcrBC*)] enables efficient cloning of methylated DNA.
- Endonuclease minus (*endA1*) to ensure high yields of DNA.
- Recombination minus (*recA1*) for greater stability of large cloned inserts.

Genotype

F mcrA Δ(mrr-hsdRMS-mcrBC) Φ80dlacZΔM15 ΔlacX74 recA1 endA1 araD139 Δ(ara, leu)7697 galU galK λ⁻ rpsL (Str^R) nupG

TransforMax EC100 Electrocompetent *E. coli*

- Transformation efficiency of >1 x 10¹⁰ cfu/µg of pUC19.

TransforMax EC100 Chemically Competent *E. coli*

- Transformation efficiency of >5 x 10⁸ cfu/µg of pUC19.

DNA	TransforMax™ EC100™ Chemically Competent <i>E. coli</i>	TransforMax™ EC100™ Electrocompetent <i>E. coli</i>
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pUC19	1.4×10^8	1.4×10^{10}
8.1-kb Clone	1.3×10^7	Not tested
13.1-kb Clone	4.3×10^6	1.3×10^9
23.1-kb Clone	9.2×10^5	3.0×10^8
145-kb BAC Clone	Not tested	7×10^7
13.1-kb clone directly from a ligation reaction	2.2×10^5	2.1×10^7

Table 1. Comparison of the transformation efficiencies of TransformMax™ EC100™ *E. coli* with a variety of DNAs. Transformations were performed using 50 µl of competent cells and either supercoiled DNAs of the indicated sizes or a 1-µl aliquot from a standard 10-µl ligation reaction. Results shown are in cfu/µg of DNA and are the average transformation efficiencies obtained from several trials.