**Description:** ElectroLigase combines T4 DNA ligase and an optimized, ready-to-use 2X reaction buffer containing a proprietary ligation enhancer and no PEG. This combination is specifically formulated to promote robust ligation of all types of DNA ends (blunt, sticky, TA). It is directly compatible, without desalting or purification, with electrocompetent cells used for transformation by electroporation. No thawing of the buffer is required as it maintains a liquid state during storage at −20°C, thereby simplifying reaction set-up. By providing an optimized ratio of enzyme and buffer components, users are able to rapidly ligate all types of DNA ends using a short incubation time at room temperature. Ligations for subcloning can be carried out in small volumes with low concentrations, allowing users to conserve precious DNA samples. These reactions can be used directly, without purification or dilution, to transform many strains of electrocompetent *E. coli*.

* Freezers vary in their actual internal temperatures. Our testing demonstrates that the master mix is liquid at −20°C.

* ElectroLigase is also compatible with chemically competent strains of *E. coli*. Performance is generally around 50% efficiency, when compared to the Blunt/TA Ligase Master Mix (NEB #M0367).

**Source:** Purified from an *E. coli* strain containing a recombinant gene encoding T4 DNA Ligase.

**Applications:**
- Vector construction
- Linker ligation
- Fragment assembly
- Library construction
- TA cloning

**Usage Notes:**
- Cells: Competent cells can vary by several logs in their competence. Perceived ligation efficiency directly correlates with the competence of the cells used for transformation. Always transform uncut vector as a control for comparison purposes.
- DNA: Purified DNA for ligations can be dissolved in dH₂O (Milli-Q® water or equivalent is preferable); TE or other dilute buffers also work well. For optimum ligation, the amount of vector DNA should be 20–100 ng and the insert should be added at a 3-fold molar excess. For ligation volumes greater than 10 µl, increase the volume of *ElectroLigase* Reaction Buffer such that it remains 50% of the reaction. Insert:vector ratios between 2 and 6 are optimal for single insertions. Ratios below 2.1 result in lower ligation efficiency. Ratios above 6:1 promote multiple inserts. If you are unsure of your DNA concentrations, perform multiple ligations with varying ratios.

**Time and Temperature:** Most ligations performed using *ElectroLigase* reach an end point at 60 minutes or less when performed between 4–37°C. Incubation beyond this time provides no additional benefit. Our recommendation for a 25°C (room temperature) incubation was chosen after evaluation of performance at 4°C, 16°C, 25°C, and 37°C. Most conditions reached at least 50% performance within 30 minutes.

**Biology:** Some DNA sequences are not easy to clone. Sequences that form structures, including intact ORF in high-copy vector, toxic genes, etc. have biological effects (intact ORF in high-copy vector, toxic genes, etc.).

**Protocols**

**Ligation Protocol for Subcloning:**

1. Transfer *ElectroLigase* and *ElectroLigase* Reaction Buffer to ice prior to reaction set up. Mix tubes by finger flicking before use.
2. Combine 20–100 ng of vector* with a 3-fold molar excess of insert and adjust volume to 5 µl with dH₂O.
3. Add 5 µl of *ElectroLigase* Reaction Buffer and 1 µl of *ElectroLigase* and pipet up and down 7–10 times to mix.
4. Incubate ligation reaction at room temperature (25°C) for 30 minutes.
5. Inactivate the ligase by incubating the reaction at 65°C for 15 minutes.
6. Chill sample on ice (if to be used within a few hours) or store at −20°C.

* In-house testing has demonstrated that maximal transformation efficiency is achieved using between 20–100 ng of vector (blunt or sticky, including T-vectors) and a corresponding 3-fold molar excess of the insert to be ligated into the vector.

**Transformation Protocol:**

Electrocompetent strains of *E. coli* (commercially available or prepared by user) can be transformed by ligation products prepared using *ElectroLigase*. Chemically competent cells are also compatible, but for maximum performance with chemically competent cells, please consider using the Blunt/TA Ligase Master Mix (NEB #M0367). The following protocol is recommended by NEB. Other protocols can be used but the volume of ligation reaction used should not exceed 5 µl reaction per 50 µl cells.

1. Thaw competent cells on ice.
2. Aliquot 40 µl of cells into a 1.5 ml microcentrifuge tube on ice.
3. Add 2 µl of the ligation reaction to the cells and mix with finger-flicking. Do not vortex the tube.
4. Transfer DNA/competent cell mixture to a precooled electroporation cuvette and follow the manufacturers recommendations for electroporation (e.g. 2500 V, 200 Ω, 25 µF, 2 mm gap cuvette).
5. Add 760 µl recovery media (e.g. SOC) to the cuvette, mix, transfer the transformed cells to a culture tube and incubate for one hour at 37°C with shaking (200–250 rpm).
6. Spread 50 µl of the outgrowth (undiluted or diluted 1:5 with recovery media) onto appropriate antibiotic selection plates and incubate overnight at 37°C.

**Typical Results:**

Transformation efficiencies around 3 x 10⁶ cfu/µg are typically achieved for recombinant blunt-end vectors (vector + insert), using cells with a 5 x 10⁴ calculated efficiency with uncut DNA. Results for TA cloning and standard cohesive end (4 bp overhang) cloning produce even higher numbers, often over 10⁷ cfu/µg. This corresponds to several hundred colonies on a plate when 50 µl of a 1 ml outgrowth is plated at a 1:5 dilution. As all ligation and transformation protocols, many factors affect the calculated transformation efficiency, including purity and integrity of DNA ends, competence of the cells being transformed, media choices, incubation temperatures and times and biological effects (intact ORF in high-copy vector, toxic genes, etc.).
Companion Products Sold Separately:
Blunt/TA Ligase Master Mix
#M0367S  50 rxns
#M0367L  250 rxns

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