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240 County Road Ipswich, MA 01938-2723 Tel 978-927-5054 Fax 978-921-1350 www.neb.com info@neb.com

## New England Biolabs Certificate of Analysis

| Product Name:          | Quick Ligation™ Kit  |
|------------------------|--|
| Catalog Number:        | M2200S   |
| Unit Definition:       | N/A  |
| Lot Number:            | 10013771   |
| Expiration Date:       | 06/2020  |
| Storage Temperature:   | -20°C  |
| Storage Conditions:    | 10 mM Tris-HCl , 50 mM KCl , 1 mM DTT , 0.1 mM EDTA , 50 % Glycerol,<br>(pH 7.4 @ 25℃) |
| Specification Version: | PS-M2200S/L v1.0   |

| Quick Ligation™ Kit Component List |                                 |            |                      |  |
|------------------------------------|---------------------------------|------------|----------------------|--|
| NEB Part Number                    | Component Description           | Lot Number | Individual QC Result |  |
| M2200SVIAL                         | Quick Ligation™ Kit             | 10009400   | Pass                 |  |
| B2200SVIAL                         | Quick Ligation™ Reaction Buffer | 0021804    | Pass                 |  |

| Assay Name/Specification   | Lot # 10013771 |
|--|----------------|
| <b>Endonuclease Activity (Nicking)</b><br>A 50 $\mu$ I reaction in NEBuffer 1 containing 1 $\mu$ g of supercoiled PhiX174 DNA and a<br>minimum of 2000 units of Quick Ligase incubated for 4 hours at 37°C results in <10%<br>conversion to the nicked form as determined by agarose gel electrophoresis.  | Pass           |
| <b>Exonuclease Activity (Radioactivity Release)</b><br>A 50 $\mu$ I reaction in NEBuffer 1 containing 1 $\mu$ g of a mixture of single and<br>double-stranded [ <sup>3</sup> H] E. coli DNA and a minimum of 2000 units of Quick Ligase<br>incubated for 4 hours at 37°C releases <0.1% of the total radioactivity.  | Pass           |
| <b>Functional Testing (Ligation and Transformation)</b><br>After a five-minute ligation of linearized, dephosphorylated LITMUS 28 or pUC19<br>(containing either blunt [EcoRV] or cohesive [HindIII] ends) and a mixture of<br>compatible insert fragments, transformation into chemically competent E. coli DH-5<br>alpha cells yields a minimum of 1 x 10e6 recombinant transformants per µg plasmid<br>DNA. | Pass           |
| <b>Ligation and Recutting (Terminal Integrity, Digested DNA)</b><br>A 20 µl reaction in 1X T4 DNA Ligase Reaction Buffer containing 2 µg of Lambda<br>DNA-HindIII Digest and a minimum of 4000 units of Quick Ligase incubated for 16<br>hours at 37°C results in >95% ligation of the DNA fragments as determined by agarose  | Pass           |





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| gel electrophoresis. Of these ligated fragments, >95% can be recut with HindIII.   |                |
| <b>Non-Specific DNase Activity (16 Hour)</b><br>A 50 µl reaction in NEBuffer 1 containing 1 µg of CIP-treated Lambda-HindIII DNA and<br>a minimum of 2000 units of Quick Ligase incubated for 16 hours at 37°C results in a<br>DNA pattern free of detectable nuclease degradation as determined by agarose gel<br>electrophoresis.  | Pass           |
| Protein Purity Assay (SDS-PAGE)<br>Quick Ligase is ≥ 95% pure as determined by SDS-PAGE analysis using Coomassie Blue<br>detection.  | Pass           |
| <b>qPCR DNA Contamination (E. coli Genomic)</b><br>A minimum of 2000 units of Quick Ligase is screened for the presence of E. coli<br>genomic DNA using SYBR® Green qPCR with primers specific for the E. coli 16S rRNA<br>locus. Results are quantified using a standard curve generated from purified E. coli<br>genomic DNA. The measured level of E. coli genomic DNA contamination is $\leq$ 1 E. coli<br>genome. | Pass           |
| <b>RNase Activity (Extended Digestion)</b><br>A 10 µl reaction in NEBuffer 4 containing 40 ng of a 300 base single-stranded RNA<br>and a minimum of 1 µl of Quick Ligase is incubated at 37°C. After incubation for 16<br>hours, >90% of the substrate RNA remains intact as determined by gel electrophoresis<br>using fluorescent detection.   | Pass           |
| <b>Single Stranded DNase Activity (FAM-Labeled Oligo)</b><br>A 50 µl reaction in CutSmart® Buffer containing a 20 nM solution of a fluorescent<br>internal labeled oligonucleotide and a minimum of 10,000 units of Quick Ligase<br>incubated for 16 hours at 37°C yields <5% degradation as determined by capillary<br>electrophoresis.   | Pass           |
| <b>DNase Activity (Labeled Oligo, 3' extension)</b><br>A 50 µl reaction in CutSmart® Buffer containing a 20 nM solution of a fluorescent<br>labeled double-stranded oligonucleotide containing a 3' extension and a minimum of<br>10,000 units of Quick Ligase incubated for 16 hours at 37°C yields <5% degradation<br>as determined by capillary electrophoresis.  | Pass           |
| DNase Activity (Labeled Oligo, 5' extension)<br>A 50 µl reaction in CutSmart® Buffer containing a 20 nM solution of a fluorescent<br>labeled double-stranded oligonucleotide containing a 5' extension and a minimum of<br>10,000 units of Quick Ligase incubated for 16 hours at 37°C yields <5% degradation<br>as determined by capillary electrophoresis.   | Pass           |





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| Double Stranded DNase Activity (Labeled Oligo)                                     | Pass           |
| A 50 µl reaction in CutSmart® Buffer containing a 20 nM solution of a fluorescent  |                |
| labeled double-stranded oligonucleotide containing a blunt end and a minimum of    |                |
| 10,000 units of Quick Ligase incubated for 16 hours at 37°C yields <5% degradation |                |
| as determined by capillary electrophoresis.  |                |

This product has been tested and shown to be in compliance with all specifications.

Klorengen

Mary Lorenzen Production Scientist 29 Jun 2018

Michae 11.

Michael Tonello Packaging Quality Control Inspector 12 Jul 2018

