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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:	Ph.D.™-12 Phage Display Peptide Library Kit
Catalog Number:	E8110S
Packaging Lot Number:	10098420
Expiration Date:	10/2022
Storage Temperature:	-20°C
Specification Version:	PS-E8110S v2.0

Ph.D.™-12 Phage Display Peptide Library Kit Component List				
NEB Part Number	Component Description	Lot Number	Individual QC Result	
S1259AVIAL	-96 glll Sequencing Primer (20-mer)	10066252	Pass	
S1258AVIAL	-28 glll Sequencing Primer (22-mer)	10066251	Pass	
N7024AVIAL	Biotin	10066253	Pass	
N7023AVIAL	Streptavidin, lyophilized	10081107	Pass	
E8111AVIAL	Ph.D.™-12 Phage Display Peptide Library	10085513	Pass	
E4104SVIAL	E.coli K12 ER2738	10055843	Pass	

Assay Name/Specification	Lot # 10098420
Absolute Phage Titer Infection of a mid-log culture of E. coli ER2738 with Ph.D. [™] -12 Phage Display Peptide Library followed by plating, yields \ge 1 x 101 ³ pfu/ml.	Pass
Sequence Verification (DNA) The Ph.D.™-12 Phage Display Peptide Library was sequenced using 5'-CCCATGTACCGTAACACTGAGTTTC-3' as a primer to confirm the correct form of the cloned insert on the displayed peptide, X12-GGG.	Pass
Functional Testing (Panning) A 100-fold representation of the Ph.D. TM -12 Phage Display Peptide Library containing approximately 1011 pfu is diluted in 200 µl TBS and panned against 300 ng β -endorphin monoclonal antibody. The bound phage is affinity captured using magnetic beads and eluted with 1 ml of 0.2M Glycine-HCl, pH 2.2. After three rounds of selection, ≥75% of sequences contain a motif related to the known epitope for the antibody.	Pass
Phage Contamination (Environmental) A 1:100 dilution of an overnight culture of E. coli ER2738 was made in 20 ml LB, to which 10 ³ pfu of Ph.D. [™] -12 Phage Display Peptide Library was added. The flask was	Pass





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incubated at 37°C on a rotating shaker for 5 hours. A 1 ml volume of culture was removed and centrifuged. A volume of culture supernatant equivalent to the initial PFU input was added to a second, 20 ml culture like the first. The final culture supernatant was plated on three LB/IPTG/Xgal plates and then titered. Fewer than 20% clear or white plaques were observed in a minimum of 100 total plaques counted on each plate.	

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

One or more products referenced in this document may be covered by a 3rd-party trademark. Please visit www.neb.com/trademarks for additional information.

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Alicia Bielik Production Scientist 23 Feb 2021

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Michael Tonello Packaging Quality Control Inspector 23 Feb 2021

