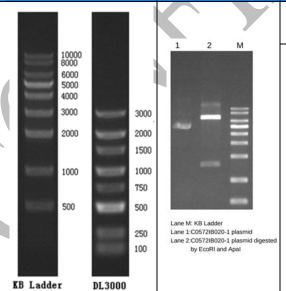


CERTIFICATE OF ANALYSIS					
Vector Name	pCC1FOS	Catalog	V	7008674	
Project/ Lot No.	C0572IB020-1 /PN99239	Size	8,	8,139 bp	
Quantity	10 ug	Resistance	С	Chloramphenicol	
QC Results					
Test Items	Specifications	Specifications			
Insert Sequence	Insert sequence re	Insert sequence results consistent with target			
Vector Sequence	Flanking sequence	Flanking sequence consistent with expected			

ORF Across Junction Correct and consistent with target N/A Expected fragment sizes observed **Restriction Digest** Pass **PCR** Amplification Correct without non - specific bands Pass Actual yield (by A 260) $5\mu g/5\mu g$ Concentration (n/a if lyophilized) N/A Purity (A 260/A280 = 1.8 - 2.0) Pass **DNA Quantity/Quality** # of Tubes 2 Matrix TE (lyophilized) Verified, <0.1 EU/μg (Endo-Free Preps Only) **Endotoxin Test** N/A Clear, no visible particles Pass **Appearance** Label Correct and white Pass **Comments**

Restriction Digestion Map



Lane 1: Plasmid

Lane 2: Plasmid Digested with BamHI

Lane M: DNA Marker

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Certified by: Chou Fang

Date: Mar/14/2023

Valid until: Mar/13/2025



Note

BAC and fosmid clones are highly suitable for modification by recombineering but, because they are present at low (1-2) copies per cell, the DNA is difficult to isolate in high yield and purity. To overcome this limitation vectors, e.g. pCC1BAC/pCC1FOS, have been constructed that contain the additional replication origin, oriV, which permits copy-number to be induced transiently when propagated in a suitable host strain, e.g. EPI300, that supplies the cognate trans-replication protein TrfA.

Protocol for EPI400/EPI300

- a) Add 4 ml LB media into each test tube. Inoculate each tube with bacterial culture with antibiotic at the proper concentration.
- b) Incubate the tubes at 37°C, shaking overnight.
- c) Dilute the starting culture (from step b) 1:10 into antibiotic-supplemented fresh media.
- d) Supplement induction solution with a ratio of 1:1000, and grow the culture at 37°C for 4 h with vigorous shaking (approx. 250 rpm).
- e) Isolate DNA from the induced culture cells as per the protocol provided

Vector Map

