

**Bacteriophage BCP01 and Its Endolysin as Tools for Biocontrol and Rapid
Detection of *Bacillus cereus***

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Supplementary Table and Figures

Table S1. Host range of LysBCP01, LysPBC1, BCP01EAD-PBC1CBD and PBC1EAD-BCP01CBD chimeric proteins^a

Bacteria	LysBCP01	LysPBC1	BCP01EAD-PBC1CBD	PBC1EAD-BCP01CBD
<i>Bacillus cereus</i> ATCC 10876	+++	++	+++	+++
ATCC 13061	-	+++	-	-
ATCC 14579	+++	++	++	+
ATCC 21768	+++	+++	+++	++
ATCC 21772	+++	+++	+++	+++
ATCC 27348	++	+++	+++	+++
<i>B. circulans</i> JCM2504	-	+	-	-
<i>B. licheniformis</i> JCM 2505	++	+	-	++
<i>B. megaterium</i> JCM 2506	+	+	-	-
<i>B. mycoides</i> ATCC 6452	-	++	-	-
<i>B. pumilus</i> JCM 2508	-	-	-	-
<i>B. sphaericus</i> JCM2502	+++	+	++	-
<i>B. subtilis</i> ATCC23857	-	+	-	-
<i>B. thuringiensis</i> ATCC10792	+++	++	+++	+++
<i>Enterococcus faecium</i> ATCC27276	-	-	-	-
<i>Listeria monocytogenes</i> ATCC 15313	-	-	-	-
<i>Staphylococcus aureus</i> ATCC 29213	-	-	-	-
<i>Yersinia enterocolitica</i> ATCC 55075	-	-	-	-

^a, The relative lytic activity of endolysin was obtained by measuring the percent drop in OD₆₀₀ in 8 min. -, no lysis; +, limited lysis; ++, medium lysis; +++, rapid lysis.

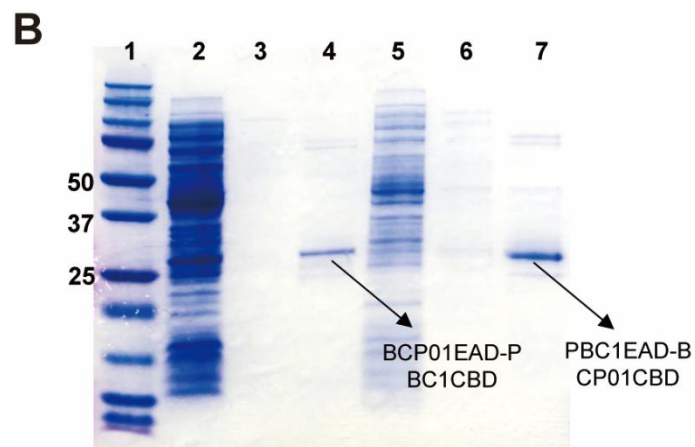
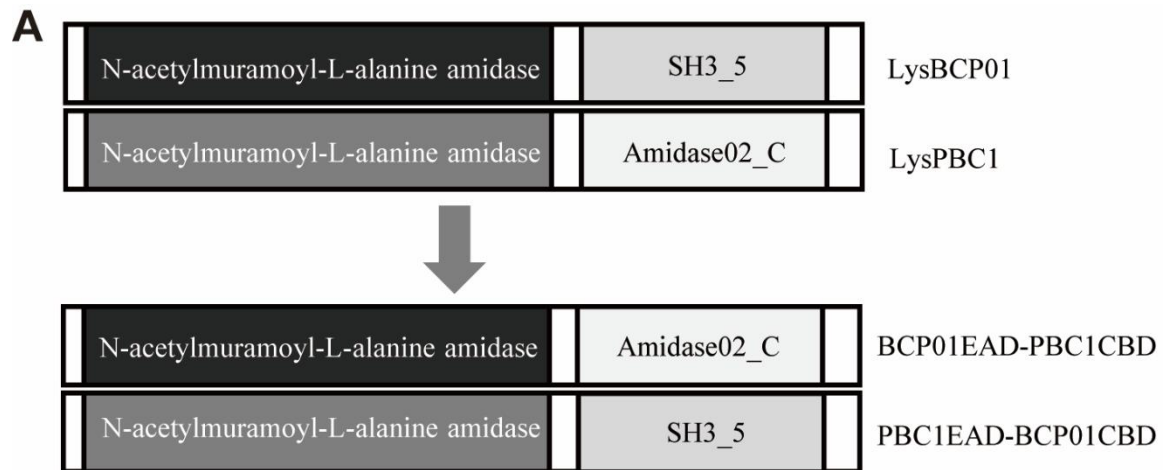


Fig. S1. Construction and purification of chimeric endolysins derived from LysBCP01 and LysPBC1. (A) Schematic diagrams of BCP01EAD-PBC1CBD and PBC1EAD-BCP01CBD. (B) SDS-PAGE analysis of purified chimeric proteins.

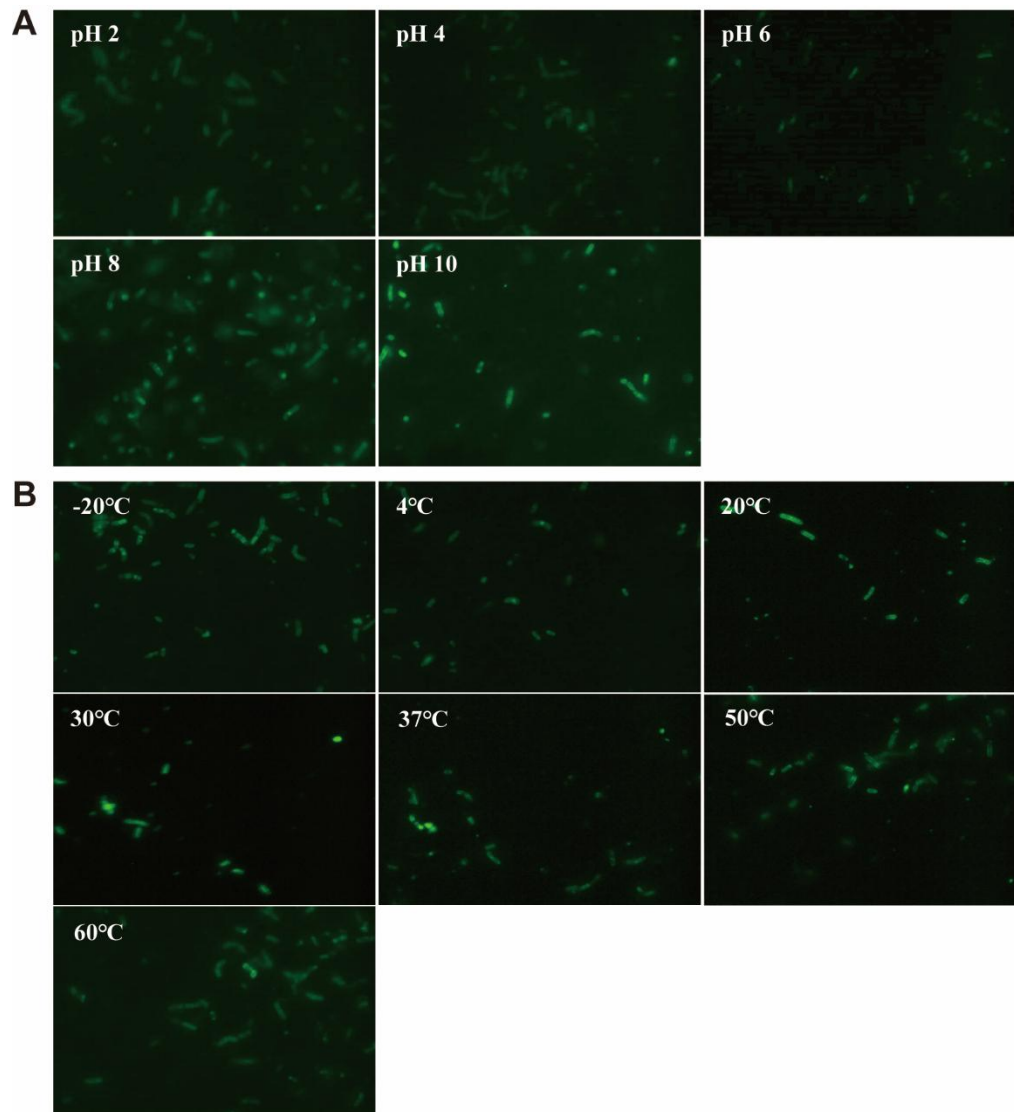


Fig. S2. Stability of EGFP-LysBCP01_CBD binding activity under different pH (A) and temperature (B) conditions against *B. cereus* ATCC 14579.

