Enzyme-Based Listericidal Nanocomposites

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Cell lytic enzymes represent an alternative to chemical decontamination or use of antibiotics to kill pathogenic bacteria, such as listeria. A number of phage cell lytic enzymes against listeria have been isolated and possess listericidal activity; however, there has been no attempt to incorporate these enzymes onto surfaces. We report three facile routes for the surface incorporation of the listeria bacteriophage endolysin Ply500: covalent attachment onto FDA approved silica nanoparticles (SNPs), incorporation of SNP-Ply500 conjugates into a thin poly(hydroxyethyl methacrylate) film; and affinity binding to edible crosslinked starch nanoparticles via construction of a maltose binding protein fusion. These Ply500 formulations were effective in killing L. innocua (a reduced pathogenic surrogate) at challenges up to 10^5 CFU/ml both in non-growth sustaining PBS as well as under growth conditions on lettuce. This strategy represents a new route toward achieving highly selective and efficient pathogen decontamination and prevention in public infrastructure.

Results

6x-His-Ply500 was produced in E. coli and purified by Ni-NTA chromatography (see details in Supplementary information). The purified protein (33.4 kDa) was stored in Phosphate Buffered Saline (PBS) at 4°C at...
concentrations <1 mg/ml to avoid aggregation. Unlike many phage endolysins\textsuperscript{2}, Ply500 was active and stable as a monomer (90 ± 3% of total purified enzyme; see details in Supplementary Information and Supplementary Fig. S1). The monomer was stable for at least 7 days when stored in solution or as a lyophilized powder at 4 °C, with <5% of monomer converted into dimer. The catalytic activity of Ply500 was initially assessed via breakdown of isolated cell walls of \textit{L. innocua} (Fig. 1a), with optimum assay conditions of pH 8.0 in 250 mM NaCl and 50 mM Tris buffer, translating into a specific activity of 3000 ± 300 U/mg. The rate of turbidity decrease was essentially linear with enzyme concentration in a range of 0.5 to 2.0 µg/ml (Fig. 1a, Inset). Irrespective of the concentration of Ply500 used, the final turbidity obtained after lysis was ~60% of the initial value, indicating that the enzyme is rather specific and cannot cleave the cell wall peptidoglycan into extremely small, and more soluble fragments.

To date, Ply500 has not been employed for actual cell killing assays. Therefore, to obtain accurate cell killing activity, we proceeded to establish a protocol for the antimicrobial/plating assay of Ply500. An appropriate amount of filter-sterilized enzyme (1–20 µg) was incubated with \textit{Listeria} (10⁶ CFU/ml) in 1 ml of sterilized PBS buffer containing 0.1% (v/v) Tween 80 followed by plating on BH1 agar plates (Fig. 1b). Ply500 was highly active against \textit{L. innocua}; for a 10⁶ CFU/ml \textit{L. innocua} cell challenge, a 3-log and close to 5-log reduction was obtained after 24 h using 10 and 20 µg/ml of enzyme, respectively (Fig. 1b). To ascertain whether the residual \textit{L. innocua} cells were naturally resistant to Ply500, we performed six repeated cycles of killing, outgrowth, and killing and observed no resistance against Ply500 (see details in Supplementary Information).

Endolysins require both the N- and C-domains to be in proper orientation to access the substrate (bacterial cell wall) for activity\textsuperscript{10}. Thus, even small limitations in accessibility and enzyme conformational deformation due to immobilization would likely result in very low observed activity. For these reasons, we hypothesized that immobilization onto a surface that is similar in size to the enzyme itself, yet possessing the virtues of an immobilization support, would yield highly active and stable Ply500 formulations. To that end, we chose silica nanoparticles (SNPs) as the conjugation material. SNPs are cost effective, inert and are considered as GRAS by the U.S. Food and Drug Administration\textsuperscript{23}, which enable them to be used in the food industry\textsuperscript{24}. SNPs were functionalized with NCO and methacrylate groups (Fig. 2a), and the methacrylate functionalization was qualitatively confirmed through reaction of the methacrylate group double bond with Br\textsubscript{2}, followed by X-ray photoelectron spectroscopy (XPS) analysis of the Br signature (see details in Supplementary Information; Supplementary Fig. S2). Titration of the functionalized SNPs with benzylamine indicated that functionalization of both NCO and methacrylate groups is 0.1 ± 0.01 mmol/g SNPs (see details in supplementary information). The enzyme loading was 0.14 ± 0.03 mg Ply500/mg SNPs. Dynamic light scattering was used to determine the size of SNPs in PBS buffer (containing 0.1% (v/v) Tween 80) at each step of modification (see details in Supplementary Information; Supplementary Table S1). Modification with NCO and methacrylate SNPs resulted in formation of SNP aggregates of 65 ± 18 nm (Supplementary Table S1). Immobilization of Ply500 led to further crosslinking and increased aggregate size of 81 ± 29 nm (Supplementary Table S1).

We evaluated the effect of covalent attachment on the bactericidal activity of Ply500 (Table 1). Ply500-SNP conjugates showed ~33% retention of activity, as determined by the cell plating assay (on basis of number of \textit{L. innocua} cells killed per mg of Ply500 in the conjugates) (Table 1). While free Ply500 (20 µg/ml) showed complete cell killing against a challenge of 10⁶ CFU/ml in 24 h, a similar bactericidal effectiveness required Ply500-SNP conjugates containing 60 µg/ml of Ply500 (Table 1), indicating some loss of enzyme activity upon immobilization. The SNP-no-enzyme control was ineffective in killing \textit{L. innocua}. The SNP-Ply500 conjugates were substantially more stable than the free enzyme at both 4 and 25 °C, temperatures representative of storage and operational conditions, respectively (see details in supplementary information). At 4 °C the free enzyme lost 50% of its log killing (calculated on the basis of log reduction per mg of Ply500) after 40 days, whereas the SNP-Ply500 conjugates retained >95% of its initial log killing during this time (Fig. 2b). More striking was the stabilization at 25 °C, where the native enzyme was completely deactivated after 15 days, while with the nanoparticle conjugates retaining >95% log killing during this time (Fig. 2c). The increased stability of SNP-Ply500 conjugates obtained was in agreement with previous reports, which demonstrate that protein stability is enhanced by the highly curved surfaces of nanosupports\textsuperscript{19,24–26}. The retention of bactericidal activity and high storage stability were sufficiently encouraging to proceed with incorporating these conjugates into a polymer film.

The Ply500-SNP conjugates containing methacrylate groups were co-polymerized with HEMA and polyethylene glycol dimethacrylate (PEGDMA), as crosslinker, in the presence of ammonium persulphate (APS) and tetramethyl ethylenediamine (TEMED) as initiators (Fig. 2a). The Ply500-SNP conjugates were dispersed throughout the film, as shown via scanning electron microscopy (SEM) (Fig. 2a). The polymer film containing Ply500 conjugates was then tested for bactericidal activity against \textit{L. innocua} by incubating the film in 2 ml of a suspension containing 10³ CFU/ml for 24 h (Table 1). This incubation resulted in a 1.6-log reduction in the number of \textit{L. innocua} cells, whereas the control film with only SNPs (i.e., without Ply500) was devoid of antimicrobial activity. Next, we confirmed that the activity of the film was not due to leaching of Ply500 into the solution. Specifically, the Ply500-SNP containing film was incubated in PBS (with 0.1% (v/v) Tween 80) for 24 h, the film was removed, and the solution was tested for free protein leached using the BCA assay. No free protein was observed in the washes; however, the detection limit of the BCA assay is not sufficiently low to truly assess for leached enzyme. A more sensitive
antimicrobial activity assay, therefore, was performed on the washes, which also yielded no evidence of protein leaching from the film. To ensure that Ply500-SNP conjugates were not simply leaching in an inactive form, we used a highly sensitive surrogate fluorescent protein, green fluorescent protein (GFP), incorporated into pHEMA in a manner identical to that of Ply500 (see details in Supplementary information, Supplementary Fig. S3). Specifically, GFP was immobilized onto the SNPs, with a measured loading of 0.04 mg/mg SNPs (Supplementary Fig. S3(a)). Then the polymer film was cast with the embedded GFP-SNP conjugates. The resulting film containing GFP-SNP conjugates was incubated in PBS (with 0.1% (v/v) Tween 80), respectively, for 24 h and the film was then removed from the buffer. The resulting fluorescence of the buffer solutions was no higher than that for controls (polymer films with SNP without GFP), indicating no significant protein leaching from the film (Supplementary Fig. S3 (b)). Collectively,

Table 1 | Anti-listeria activity of different Ply500 formulations against *L. innocua* assessed after incubation with 1 mL of a suspension containing 10⁵ CFU/ml

| Ply500 Formulation           | Log CFU reduction after 24 h | Retention of native enzyme activity (plating assay) as a percentage relative to native enzyme | Retention of native enzyme activity (cell wall assay) as a percentage relative to native enzyme |
|------------------------------|------------------------------|------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------|
| Ply500 (20 µg/ml)            | 4.7 ± 0.05                   | 100%                                                                                     | 100%                                                                                     |
| Ply500-SNP (60 µg/ml)        | 4.6 ± 0.02                   | 33                                                                                       | n.d.                                                                                     |
| Ply500 film (500 µg/ml)      | 1.6 ± 0.12                   | 3.9 ± 0.05                                                                               | n.d.                                                                                     |
| FP (20 µg/ml)                | 0.3 ± 0.02                   | 51 ± 4                                                                                   | 4.5 ± 0.5                                                                               |
| FP10 (20 µg/ml)              | 2.5 ± 0.30                   | 99 ± 0.5                                                                                 | 32 ± 4.0                                                                                |
| FP10-starch nanoparticle (80 µg/ml) | 2.9 ± 0.18                 | 24 ± 0.02                                                                                | n.d.                                                                                     |

*Listeria cells were grown in BHI media for 7 h. Standard deviations were calculated from triplicate measurements. No significant killing was observed with controls of SNP-NCO, overnight wash of Ply500-SNP, Film (without Ply500), overnight wash of Ply500 film, MBP, starch nanoparticles, and MBP-starch nanoparticles. The concentration (in µg/mL) in column one represents the Ply500 content of the formulation.

*n.d.: not determined.

**Figure 2** | (a) Schematic representation of covalent immobilization of Ply500 on silica nanoparticles followed by co-polymerization with HEMA to form an anti-listeria polymeric film. Storage stability of native Ply500 (20 µg) (●) and Ply500-SNP (30 µg) (□) in sterile PBS at (b) 4°C and (c) 25°C. The amount (in µg) represents the Ply500 content of the formulation. Error bars represent standard deviations from triplicate measurements.
these results indicate that the anti-listeria activity of the films was a result of Ply500 activity within the film.

Ready to eat (RTE) foods, which are stored at 4°C and are not cooked properly prior to consumption, are the main source of listeria outbreaks. Cross-contamination in RTE foods comes from adhered cells or listeria biofilms in cold storage equipment. We therefore examined adherence and growth of L. innocua cells on Ply500-containing polymer films. The cells were allowed to adhere and grow on the pHEMA polymer films for 48 h at the typical food storage and processing temperature of 4°C. No L. innocua cells were found on Ply500-SNP-containing films as compared to 50 ± 10 cells found on the control (no-enzyme) film. Hence, the SNP-Ply500-containing pHEMA films were able to prevent the viability of adhered listeria cells.

An additional challenge in the food industry is the need for safety in packaging and food transport. Indeed, nearly 40% of the food produced in the world is never eaten27,28. Therefore, modification in packaging and food transport. Indeed, nearly 40% of the food cells.

The purified MBP-Ply500 (FP) fusion protein was constructed to contain an N-terminal MBP fusion and a C-terminal 6x-His Tag (Fig. 3). Following optimization of enzyme induction and expression in BL21 cells, the fusion protein was purified using Ni-NTA chromatography (see details in Supplementary Information, Supplementary Fig. S4). The purified MBP-Ply500 (FP) was then evaluated for activity against L. innocua using the viable plate assay, as was used for native Ply500 (Table 1). Free Ply500 (20 μg/ml) challenged with 10^6 CFU/ml resulted in a nearly 5-log reduction after 24 h. The fusion protein, however, was nearly inactive with only a 0.3-log reduction at the same Ply500 concentration and incubation time. We speculated that the decreased activity of the fusion protein could be due to steric hindrance by the larger MBP (44 kDa). To overcome this potential steric limitation, we hypothesized that a short linker between the MBP and Ply500 could relax this steric constraint.

Our approach took advantage of the presence of a 10 amino acid putative linker between the catalytic and binding domains of Ply500 (Pro-Ala-Ala-Thr-Gln-Asn-Thr-Asn-Thr-Asn)36, and we prepared a modified fusion protein containing this 10-mer between the MBP and Ply500 components of the fusion protein (FP10). The insertion of a linker improved the cell lytic activity of the fusion protein (Table 1), with 2.5-log reduction observed after 24 h of incubation with 10^6 CFU/ml L. innocua. We then proceeded with affinity immobilization of the FP10 onto crosslinked starch nanoparticles. The size of starch nanoparticles was 80 ± 34 nm (as determined by SEM (Supplementary Fig. S5)). A loading of 0.11 ± 0.02 mg protein/mg of starch nanoparticles was obtained for FP10. We then evaluated the effect of affinity immobilization on the bactericidal activity of FP10 (Table 1); 24% retention in activity (on the basis of the plating assay, e.g., number of listeria cells killed per mg of Ply500) was observed for FP10-starch conjugates, with 2.9-log reduction observed after 24 h of incubation with 10^6 CFU/ml L. innocua (Table 1). A control with only MBP attached to starch nanoparticles was inactive. A control consisting of free FP10 at 80 μg/ml of Ply500 could not be performed due to rapid formation of visible aggregates in the presence of listeria cells.

To evaluate an actual food material, we tested the efficacy of different Ply500 formulations against exponentially growing L. innocua inoculated onto iceberg lettuce leaves. L. innocua grew on the lettuce; after 24 h at 25°C the cell density was ca. 10^4 CFU. Treatment of the contaminated lettuce with free Ply500 (50 μg) or SNP-Ply500 conjugates (200 μg) resulted in complete killing of the contaminating L. innocua (Table 2). Less striking results were obtained with the MBP-fusion Ply500, both in the free form and adsorbed to starch. Nonetheless, nearly 1-log unit reduction in CFU was achieved with the FP10-starch nanoparticles.

Discussion

In line with the major requirement of surface-active decontamination methodologies for the food industry, the primary goal of our work was to develop stable and active surface incorporated phage

Table 2 | Anti-listeria activity of different Ply500 formulations on lettuce inoculated with L. innocua cells

| Ply500 Formulation | log CFU after 24 h |
|--------------------|-------------------|
| Cell control       | 1.7 ± 0.09        |
| Lette + listeria control | 4.0 ± 0.05 |
| Ply500 (50 μg)     | No cells*         |
| Ply500-SNP (200 μg)| No cells*         |
| FP10 (42 μg)       | 2.1 ± 0.05        |
| FP10-starch nanoparticle (80 μg)| 2.8 ± 0.14 |

*Listeria cells were grown in BHI media for 7 h and lettuce was inoculated with 50 L. innocua cells. Standard deviations were calculated from triplicate measurements. No significant killing was observed with controls of SNP-NCO, MBP, starch nanoparticles, MBP-starch nanoparticles. The amount (μg) represents the Ply500 content of the formulation.

*No viable L. innocua cells were observed in the plating assay.
lytic enzyme formulations. We demonstrated the potency of Ply500 by establishing a cell plating assay for *L. innocua* cell killing. Listeriosis treatment includes antibiotics, such as penicillin, ampicillin, and amoxicillin, but their overuse often leads to antibiotic resistance. To evaluate whether Ply500 use could lead to gained resistance against *L. innocua* against Ply500 lytic activity, we repeatedly (6 times) exposed *L. innocua* to Ply500 with the expectation that residual viable cells would be more resistant to Ply500 than the original starting stock. However, no change in the log-killing was observed after each round of Ply500 treatment, which suggests that we were not selecting Ply500 resistant strains of *L. innocua*. These results are not surprising. Accelerated evolution of *Bacillus cereus* was performed by Fischetti et al. who found no gained resistance of the bacterium upon treatment with a bacillus phage endolysin. While we cannot rule out the ultimate formation of gained resistance of *L. innocua* to Ply500, our treatment indicates that we were not selectively isolating more resistant strains of *L. innocua*.

For surface-based applications of cell-specific endolysins in preventing contact-based bacterial infections, the enzymes must be surface immobilized with long shelf-life and operational reusability. Due to their chimeric structure and requirement of both catalytic and binding domains to be in the proper orientation for lytic activity, the main challenge of this work was to obtain active enzyme after its incorporation onto surfaces. To achieve this goal, we covalently immobilized Ply500 onto FDA approved nanoscale silica particles. Immobilization of Ply500 onto SNPs resulted in enhanced storage stability at 4 °C and operational stability at 25 °C. This stabilization enabled us to test Ply500-SNP conjugates against *L. innocua* cells not only in buffer but also under exponentially growing conditions on iceberg lettuce leaves. Moreover, we developed an effective listericil nanocomposite polymer film based on Ply500-SNP conjugates. These films were capable of complete killing of listeria cells on contact and was able to prevent growth of listeria cells at 4 °C. This result makes SNP-Ply500-containing PHHEMA films potential candidates for applications as anti-listeria coatings of cold storage or food processing equipment, such as a thaw and cold rooms, conveyor belts, grinders, commercial kitchens, and packaging materials. Finally, we exploited affinity immobilization of Ply500 on edible crosslinked starch nanoparticles by constructing a MBP-Ply500 fusion protein. These conjugates also exhibited listericidal activity. These Ply500-starch conjugates may find further applications in antimicrobial packaging systems like spraying/ coating of antimicrobial enzyme-starch conjugates on food before packaging or incorporation into packaging materials. By extended our approach to other cell lytic enzymes, our findings may encourage broader development of highly efficient bactericidal surfaces for applications in the food industry, healthcare, and other common infrastructures.

### Methods

**Listeria, plasmids and culture conditions.** *Listeria innocua* ATCC® 33090™ was used as the cell target in these studies. One shot® Top 10 and One Shot® BL21(DE3) chemically competent *E. coli* cells were obtained from Invitrogen and were used for cloning and overexpression of proteins, respectively. *L. innocua* was grown in Brain Heart Infusion (BHI) medium (BDC, MD, USA) at 37 °C with shaking at 220 rpm. From the grown culture, 1 ml was removed and placed in an Eppendorf tube and centrifuged at 10,000 g for 1 min at room temperature to obtain a pellet. The pellet was then washed twice with sterilized PBS to remove excess medium. After washing, the pellet was again resuspended in 1 ml of sterilized PBS. To obtain an approximate measure of cell density in terms of colony forming units (CFU/ml), the optical density of the microbial suspension was measured at 600 nm (with a conversion factor of 1 absorbance unit corresponding to 10⁶ CFU/ml). The bactericidal efficiency of all the enzyme formulations was determined by using a diluted suspension containing 10⁶ CFU/ml. Unless otherwise indicated, solvents and reagents were obtained from Sigma-Aldrich.

**Cloning procedures.** Standard cloning techniques were used for construction of plasmids encoding Ply500 and Ply500-based fusion proteins. The enzymes and kits used for cloning were used according to the manufacturer’s instructions.

**Construction of MBP-Ply500 (pEP) plasmid.** Ply500 gene was amplified by PCR with *Taq* polymerase (Green Master Mix, Promega), using primers Ply500_BamHI_FP (5’ CGG CGG ATC GAT GGC ATT AAG AGA GGG ATG) and Ply500_SalI_RP (5’ CGG CGC GTG CAT GTG GAT GTT GTT GAT GTT). The amplified gene was then purified by agarose gel extraction using QIAquick Gel extraction kit (Qiagen). The purified Ply500 gene and pMBP3C vector containing MBP gene were double digested with *BamHI* and *SalI* (New England Biolabs) separately for 3 h at 37 °C. After digestion, the Ply500 gene and MBP plasmid were again purified and the concentration of purified digested gene and plasmid was determined spectrophotometrically (Nanodrop ND-1000). The Ply500 gene was ligated into predigested pMBP3C using T4 ligase (New England Biolabs). The ligated plasmids were transformed into Chemically Competent One Shot® Top 10 cells, which were then cultured overnight at 30 °C. The transformed cells were then plated onto ampicillin agar plates and incubated at 37 °C for 16 h. Plasmids were isolated (Miniprep Kit, Sigma-Aldrich) and sequenced for insertion of Ply500. After confirmation by sequencing, the plasmid containing MBP-Ply500 gene was transformed into competent One Shot® BL21(DE3) *E. coli* cells for overexpression of fusion protein.

**Construction of pMBP-linker₆-Ply500 (pFP₆).** Site directed mutagenesis of plasmid containing MBP-Ply500 was performed (using QuickChange® XL Site-directed Mutagenesis Kit, Agilent) to introduce a 10 amino acid linker between MBP and Ply500. The composition of linker chosen was the same as that of a natural linker that exists between the catalytic and binding domains of Ply500 at positions 154–163; Pro- Ala-Ala-Thr-Gln-Aln-Thr-Asn-Aln. The primers used were: FP- 5’ GGCG GGA TCC CCT GCT GCA ACA CAA TAT AAC ATG GCA TTA ACA GAG and RP: 5’ CTT TGT TAA TCA TGT AGT TTT ATG GTT TGC TGG AGG GGA TCG TCC GGG.

**Plating assay of Ply500 based enzyme formulations.** Antimicrobial activity of native and immobilized Ply500 was determined by adding an appropriate amount of enzyme formulation (native Ply500 and MBP fusion proteins (20 µg), SNP-Ply500 (60 µg) and FP₆-starch nanoparticles (80 µg) in 1 ml of PBS (with 0.1% (v/v) Tween 80) containing 10⁹ CFU/ml of *L. innocua*. The presence of surfactant prevents the aggregation of cells and their adhesion to the Eppendorf/petri dish. Following 24 h incubation at 25 °C with shaking at 200 rpm, an aliquot was removed and spread onto BHI agar plates. The plates were then incubated for 18 h at 37 °C after which CFU's were measured. To evaluate bactericidal effectiveness of Ply500-containing films, a 4.2 × 3.0 × 0.05 cm cutout was subjected to a cell challenge of 10⁹ CFU/ml in 2 ml of PBS buffer containing 0.1% (v/v) Tween 80 in a sterilized Petri dish. CFU determination was performed as described above.

**Modification of silica nanoparticles.** Five grams of SNPs (10–20 nm, Sigma-Aldrich) were suspended in 120 ml of 2 M HCl (diluted in 5 l water-ethanol) and heated to 70 °C for 6 h. The acid treated SNPs were centrifuged at 4000 rpm for 20 min and washed with excess cold water to remove acid and ethanol. SNPs were finally washed with ethanol and dried under vacuum. The product (1 g SNPs) was reacted with 1 ml of a 1:1 mixture of 3-(trimethoxysilyl) propyl methacrylate and 1 ml of 3-(trimethoxysilyl) propyl isocyanate in 18 ml of anhydrous cyclohexane under nitrogen. The reaction was carried out overnight at 25 °C. The functionalized SNPs were washed (2×) with ethanol and dried under vacuum. The resulting functionalized SNPs were stored at 4 °C and protected from light by wrapping the vial with aluminium foil. For control experiments, SNPs were also functionalized with 3-(trimethoxysilyl) propyl isocyanate (SNP-NCO), but without enzyme attachment.

**Immobilization of Ply500 onto SNPs.** The Ply500 was covalently immobilized onto SNPs. Briefly, 50 µg of functionalized SNPs (methacrylate-SNP-NCO) were incubated with 5 ml of Ply500 (0.8 mg/ml) in Tris-HCl buffer (pH 8.0, 250 mM NaCl) at 4 °C for 10 h. The free amino groups on the enzyme were allowed to react with NCO groups and double bonds on SNPs. After immobilization, SNPs were washed extensively with cold Tris-HCl buffer (pH 8.0, 250 mM NaCl) and 1% Triton X (10 min) at 4 °C to remove unbound enzyme. Supernatants and washes were collected to quantify enzyme loading using BCA assay (Pierce, Rockford, IL).

**Preparation of biocatalytic polymeric film.** The biocatalytic film was prepared by copolymerizing Ply500-SNP conjugates with 2-hydroxyethyl methacrylate (HEMA) and methyl methacrylate (MMA) with double bonds on SNPs. The immobilized SNPs were washed extensively with cold Tris-HCl buffer (pH 8.0, 250 mM NaCl) and 1% Triton X for 2 h in gel casting mode (Dimensions, Bio-Rad). The polymeric biocatalytic film thus formed was washed with excess of PBS at 4 °C, cut into size of 2.7 × 4.2 × 0.05 cm and further used for anti-listeria activity. The negative control film was also prepared using methacrylate-SNP-NCO.
Anti-listeria activity of Ply500 film against adhered L. innocua cells at 4°C. Polymer film (2 × 2 cm) was mounted on a glass slide and kept on damp sterile kimwipe tissues in a Petri dish. Polymer film was inoculated with approximately 1000 L. innocua cells and incubated at 4°C for 5 h to allow adhesion to occur. Non-adherent bacteria were removed by washing gently with 5 ml of sterile PBS. The film is incubated at 4°C for 24 h after applying 100 μl of BH media to provide nutrients for the adhering bacteria. After one day incubation, the films were rinsed with sterile PBS and fresh media was added followed by further incubation at 4°C for 24 h. After 2 days, the film was again washed with sterile PBS. To count viable adhered cells, the film was suspended in 1 ml of sterile PBS (with 0.1% (v/v) tween 80) and ultrasonicated for 5 min. The film dispersed to form a homogenous suspension and the plating assay was performed on Palcam listeria-selective agar plates (EMD Chemical In., NJ, USA). A control of listeria cells was ultrasonicated for 5 min. No effect of ultrasonication on listeria cells was observed. Also, due to dispersion of film during ultrasonication, listeria cells were exposed to Ply500-SNP conjugates which were initially buried in the film. So, a control of listeria cells along with same amount of Ply500-SNP present in entire film (2 × 2 cm) was suspended in 1 ml of sterile PBS and was ultrasonicated for 5 min. No significant cell killing was observed by SNP-Ply500 conjugates after ultrasonication for 5 min.

Affinity immobilization of Ply500 on crosslinked starch nanoparticles. Crosslinked starch nanoparticles (10 mg) were weighed in an Eppendorf tube and 1 ml of 1 mg/ml protein solution (in PBS buffer) was added. The suspension was incubated overnight at 4°C with shaking at 200 rpm. After affinity immobilization of proteins onto the starch nanoparticles, the suspension was centrifuged at 14,000 g for 10 min and supernatant was collected. The starch nanoparticles were then washed five times with cold PBS (1 ml) buffer for 30 min each. The supernatant and washes were assayed for protein concentration and the FP10-starch nanoparticle conjugates activity was assessed using the cell wall assay.

Anti-listeria activity of Ply500 formulations on L. innocua inoculated lettuce leaves. The iceberg lettuce was purchased from a local supermarket. The lettuce leaves were cut into size of 4.0 × 3.0 cm and dipped in 70% (v/v) ethanol for 10 min. After washing three times with sterile PBS, lettuce leaves were inoculated with 20 ml of PBS buffer containing ~50 listeria cells. 100 μl of enzyme formulation was applied on inoculated lettuce followed by incubation at RT for 24 h. Lettuce was washed with 1 ml of PBS buffer (containing 0.1% Tween 80) and aliquots of 50, 100 and 200 ml were plated on Palcam listeria selective agar plates (EMD Chemical).

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Author contributions
J.A.D., R.S.K. and L.S.S. conceived the idea. K.S., N.G., J.S.D. and R.S.K. designed the experiments, analyzed results and wrote the manuscript. K.S. and N.G. performed the experiments. P.D., L.L. and K.K.M. helped in optimization of protein expression and activity. E.P.E. helped in designing molecular biology experiments. All authors discussed the results and commented on the manuscript.

Additional information
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