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## Article

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# **Aptamer-assisted tumor localization of bacteria for enhanced biotherapy**

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## **ABSTRACT**

Despite bacterial-mediated biotherapies have been widely explored for treating different types of cancer, their implementation has been restricted by severe side effects and low treatment efficacies, due largely to the absence of tumor-specific accumulation following administration. Here, the conjugation of aptamers to bacterial surface is described by a simple and cytocompatible amidation procedure, which can significantly promote the localization of bacteria in tumor site after systemic administration. The surface density of aptamers can be easily adjusted by varying feed ratio and the conjugation is able to increase the stability of anchored aptamers. Optimal bacteria conjugated with an average of  $2.8 \times 10^5$  aptamers per cell present the highest specificity to tumor cells in vitro, separately generating near 2- and 5-times higher accumulation in tumor tissue at 12 and 60 hours compared to unmodified bacteria. In both 4T1 and H22 tumor-bearing mouse models, aptamer-conjugated attenuated Salmonella show strikingly enhanced antitumor efficacy, along with highly activated immune responses inside the tumor. This work demonstrates how bacterial behaviors can be tuned by surface conjugation and supports the potential of aptamer-conjugated bacteria for both targeted intratumoral localization and enhanced tumor biotherapy.

## **KEYWORDS**

Bacteria; Aptamer; Conjugate; Biotherapy; Tumor

## INTRODUCTION

In the late 19th century, W. Busch and W. Coley have independently utilized *Streptococcus pyogenes* and *Serratia marcescens* to treat cancer, which appears distinct tumor suppression effects in some sarcoma patients.[1, 2] Since then, the exploration of bacterial-based tumor therapies never stops, although the therapeutic outcomes of these treatments remain unstable.[3] Particularly, with recent progress in the fields of immunology and biotechnology, bacteria have attracted increasing attention in tumor biotherapy, both as immune modulators and drug delivery vehicles.[4, 5] The infection by bacteria is able to promote the immunogenicity of tumors, which further induces effective antitumor immune responses.[6] As reported, different immune cell subtypes, such as CD4<sup>+</sup> and CD8<sup>+</sup> T cells, regulatory T cells, and tumor-associated macrophages, can be modulated to generate antitumor immunity by attenuated bacteria.[7] Furthermore, the colonization of bacteria in tumor tissue is capable to trigger the activation of innate immunocytes and the release of proinflammatory factors, which cause the disruption of tumor vasculature as well as the associated thrombosis.[8] Regarding the application of bacteria as drug carriers, the greatest strength is their favorable colonization in tumor site, which is attributed to the intratumor anaerobic condition, eutrophication, and the immunosuppressive microenvironment.[9-11] Namely, the use of bacteria as delivery vehicles can improve the accumulation and penetration of drugs in tumor tissue and hence maximize their therapeutic efficacies. Unfortunately, bacterial-mediated tumor therapy always suffers from low inhibition efficiency of tumor growth and undesirable dose-dependent side

effects, as confirmed in the failure of phase 1 clinical trial of attenuated *Salmonella typhimurium*. [12]

Substantial efforts have been made to tackle these challenges by modifying bacteria either chemically or genetically. [13, 14] To reduce systemic toxicity, several bacterial species including *Listeria*, *Escherichia coli*, and *Clostridium* have been attenuated by removing key virulence factor genes. [15] Alternatively, decoration with various types of substrates on bacterial surface, such as synthetic nanoparticles and therapeutic drugs, has been exploited to enhance antitumor activity. [16-19] We have recently wrapped with extra coatings to retain bacterial viability under unfriendly external conditions. [20-24] For instance, bacteria decorated with a stealth coating can decrease their elimination by macrophages, which subsequently prolongs blood circulation following systemic injection. [25] However, previous modifications by both chemical and genetical approaches lack the ability to endow bacteria with increment in tumor-specific localization after administration. Therefore, new strategies are highly desirable for increasing the tumor-targeting capacity of bacteria, which can in turn increase the safety and antitumor efficacy of bacterial-based therapy.

Aptamers, which are considered as “chemical antibodies”, have been broadly applied as targeting ligands due to their ability to specifically recognize cell-surface targets (such as nucleic acids and proteins), whole cells, and even tissues. [26-28] Similar to antibodies, aptamers exhibit many favorable features including high affinity, excellent specificity, and low immunogenicity. [29] In addition, aptamers can be easily

synthesized in vitro through mature solid state synthetic technology and be chemically modified with various functional groups.[30] By virtue of these unique characteristics, several therapeutic aptamers have been evaluated in clinical trials for their potential in cancer treatment.[31] However, aptamer-based drug delivery has been mainly focused on developing aptamer-drug conjugates and aptamer-functionalized nanoparticles, with aims to enhance antitumor efficacies by increasing the accumulation of drugs in tumors.[32-34] To the best of our knowledge, the use of aptamers to actively deliver living cells to tumor site by a specific ligand-receptor interaction has not been reported yet.

In this work, we describe aptamer-assisted tumor localization of bacteria for enhanced biotherapy (**Figure 1**). Tumor-specific aptamers have been conjugated to the surface of bacteria by a single-step amidation procedure, which produces aptamer-conjugated bacteria (ApCB). The preparation shows negligible influence on bacterial viability, as the obtained ApCB grow similarly to unmodified bacteria. The grafting density of aptamers on bacteria surface can be readily tuned from  $0.7 \times 10^5$  to  $5.7 \times 10^5$  aptamers per cell by controlling feed ratio. More importantly, the attachment is capable of decreasing the enzymatic hydrolysis of aptamers anchored on the surface, as up to 80% of which remains intact after 24 hours exposure to serum. Among a set of ApCB with different grafting densities, bacteria decorated with an average of  $2.8 \times 10^5$  aptamers per cell exhibit the highest specificity to tumor cells in vitro, which separately achieves 2- and 5-fold higher colonization in tumor tissue at 12 and 60 hours post tail vein injection in mice. As a proof-of-concept study,

aptamer-conjugated attenuated *Salmonella Typhimurium VNP20009* (VNP) display significantly enhanced antitumor efficacy in both 4T1 and H22 tumor-bearing mouse models, accompanied with remarkably elicited immune responses within the tumor. Therefore, our work illustrates a unique strategy to manipulate bacterial behaviors by simple surface modification and demonstrates the promising of ApCB for enhanced bacterial-mediated cancer therapy.

## RESULTS AND DISCUSSION

### Preparation and characterization of ApCB

In view of the specificity to bind with nucleolin overexpressing on the membrane of various cancer cells,[35] we selected aptamer AS1411 to prepare ApCB. On the other hand, a well-known probiotic bacterium, *Escherichia coli Nissle 1917* (EcN), was chosen as a model strain due to its ability to colonize in different tumors.[36] By amide condensation, aminated AS1411 was linked to the carboxyl group of *N*-acetylmuramic acid (**Figure 1a**), which extensively exists in the cell wall of Gram-negative bacteria.[37] With the help of 1-ethyl-3-(3-dimethylaminopropyl)carbodiimide (EDC) and *N*-hydroxysuccinimide (NHS), ApCB were prepared by co-incubating EcN with AS1411 in phosphate buffered saline (PBS) at room temperature for 3 hours and the resulting bacteria were purified simply by centrifugation. To enable the measurement by laser scanning confocal microscopy (LSCM) and flow cytometry, AS1411 labelled with fluorochrome Cyanine5 (Cy5) at 5' end of the sequence was used to prepare ApCB.



As visualized in **Figure 1c**, LSCM images showed that EcN expressing green fluorescent protein (GFP) were efficiently decorated with AS1411 (red) under the experimental condition. Successful decoration with AS1411 was further confirmed by flow cytometric analysis. The mean fluorescence intensity of ApCB was around 10-times higher than that of EcN (Supplementary **Figure S1**), indicating that bacteria were conjugated with Cy5-labelled AS1411.

Given that the surface density of aptamers could influence the binding ability of the modified bacteria,[38] we then investigated whether the number of grafted aptamers could be tuned by varying feed ratio. A given amount of  $1 \times 10^8$  colony forming units (CFU) of EcN were incubated with 0, 2, 5, and 10 nmol of AS1411, which were termed as EcN, 2ApCB, 5ApCB, and 10ApCB, respectively. Then, the resulting bacteria were analyzed by flow cytometry. As expected, the fluorescence signals increased with the concentration of AS1411, as revealed clearly by a continuous shift of the curve to a higher intensity (**Figure 1d and e**). These results suggested a concentration-dependent conjugation, which meant that the grafting density of AS1411 could be controlled through changing the feed ratio. To calculate the number of grafted aptamers per cell, EcN conjugated with different concentrations of Cy5-labelled AS1411 were detected by fluorescence spectrophotometer. The number of aptamers decorated on bacteria was quantified by calculating the difference of fluorescent intensity of the aptamer solution after reaction. In good agreement with the result of flow cytometric analysis, the number of attached aptamers increased with the incubation concentration of AS1411, as illustrated in **Figure 1f**. The quantities of

AS1411 conjugated on the surface of 2ApCB, 5ApCB, and 10ApCB were calculated to be  $0.7 \times 10^5$ ,  $2.8 \times 10^5$ , and  $5.7 \times 10^5$  aptamers per cell, respectively.

To investigate whether the preparation procedure and the conjugated aptamers had toxic side effects on bacterial viability, the growth of ApCB was assessed using plate counting. Serial dilutions of EcN, 2ApCB, 5ApCB, and 10ApCB were prepared and then spread onto Luria Bertani (LB) agar plates. After incubation at 37 °C for 24 hours, the viability of ApCB was determined by recording the number of bacterial colonies. As shown in **Figure 1g**, in contrast to EcN, 2ApCB, 5ApCB, and 10ApCB displayed no notable difference in colony number, verifying the uninfluenced bacterial viability after conjugation. In addition, the serum tolerance of an aptamer-based therapeutic agent is critical for in vivo applications.[39] We speculated that the conjugation of aptamers on the surface of bacteria could largely increase the steric hindrance towards the enzymatic hydrolysis of AS1411 by nucleases.[40] To demonstrate this hypothesis, the stability of aptamers decorated on EcN was monitored by tracking the fluorescence intensity of Cy5-labelled AS1411. ApCB were analyzed by flow cytometry after incubation with 90% phosphate-buffered serum at 37 °C for the predetermined time points. On the basis of our previous report, more than 60% of free AS1411 was found to be rapidly degraded after 48 hours incubation.[33] Extraordinarily, the stability of AS1411 grafted to the surface of EcN was greatly improved under the same experimental condition, with more than 70% of the conjugated aptamers remaining intact (**Figure 1h**). These data indicated that the

conjugation could strengthen the resistance of aptamers against serum-mediated degradation, suggesting the potential of ApCB to be utilized in vivo.

### **Increased binding of ApCB with cancer cells**

With the guidance of surface-anchored aptamers, we expected that ApCB could present specific affinity to cancer cells overexpressing the corresponding receptors. To examine the role of attached AS1411 on the invasion of EcN to cancer cells, cellular binding efficiencies of EcN, 2ApCB, 5ApCB, and 10ApCB were assessed by using 4T1 cells, a cell line that overexpresses nucleolin. LSCM imaging showed that the attachment of AS1411 to the cell surface could strengthen the binding of EcN with 4T1 cells (**Figure 2a**). Interestingly, incubation with 5ApCB visualized the highest number of bound EcN on the cell surface. Although there was much more grafted AS1411 on EcN surface, 10ApCB bound less efficiently than 5ApCB. This might be ascribed to the influence of grafting density, such as the accessibility of adjacent aptamers as well as the steric effects of neighboring aptamers on the surface.[41] In accordance with the result of confocal imaging, flow cytometric analysis indicated that the binding efficiency increased with the grafting density of AS1411 increasing from  $0.5 \times 10^5$  to  $2.8 \times 10^5$  aptamers per cell, while decreased with further increasing the number to  $5.5 \times 10^5$  (**Figure 2b**). Under the same culture condition, the percentages of 4T1 cells bound with EcN after incubation with EcN, 2ApCB, 5ApCB, and 10ApCB for 2 hours were about 5%, 15%, 50%, and 30%, respectively (**Figure 2c**). These results demonstrated that decoration with aptamers could promote the

interaction of bacteria with targeting cancer cells via specific ligand-receptor recognition, which depended on the grafting density.

To directly visualize the binding of ApCB with cancer cells, 4T1 cells after co-incubation with EcN and 5ApCB were observed by scanning electron microscopy (SEM), respectively (**Figure 3a**). Representative SEM images showed that more bacteria were attached onto the surface of 4T1 cells after culture with 5ApCB instead of EcN, which was consistent with the results of flow cytometric analysis. To verify the specific binding with cancer cells, EcN and 5ApCB were separately added to the culture medium of 293T cells, a cell line that presents low nucleolin expression.[42, 43] Flow cytometric analysis suggested that there was no significant difference in fluorescence intensity between the binding of 293T cells with EcN and 5ApCB (**Figure 3b** and **c**). Meanwhile, LSCM images displayed that the binding efficiency of 5ApCB with 293T cells was similar to that of EcN, which exhibited limited bacteria attaching onto the surface of 293T cells (**Figure 3d**). These results indicated that the attachment of AS1411 onto bacteria played a critical role for ApCB to specifically bind with cancer cells. Moreover, to further assess the binding efficiency of ApCB with cancer cells in vivo, a 4T1 tumor model was developed and the obtained tumor-bearing mice were intravenously dosed with PBS and  $1 \times 10^7$  CFU of VNP or 5ApCB, respectively. Tumor tissues were sampled and stained with both FITC-labelled anti-*Escherichia coli* and 4',6-diamidino-2-phenylindole (DAPI) at 12 hours post-injection (**Figure 3e**). Clearly, mice injected with 5ApCB presented a largely increased number of bacteria binding with cancer cells compared to EcN,

further demonstrating the specific and enhanced binding of bacteria by conjugation with aptamers on the surface.

Having confirmed the presence of exogenous function that could specifically bind with tumor cells after conjugation, we next turned our attention to evaluate the ability of ApCB to target tumor site. The tumor targeted colonization of ApCB was determined by tumor imaging using a 4T1 tumor-bearing mouse model. Based on the result of in vitro assay, 5ApCB with the highest binding efficiency were used to evaluate the tumor accumulation. To enable visual tracking, luxCDABE engineered 5ApCB were intravenously injected into the tumor-bearing mice through tail vein, while engineered EcN without surface aptamers were administered as a control. At the indicated time points post-injection, the dosed mice were observed by in vivo imaging system (IVIS). As shown in **Figure 4a** and **b**, the average luminescence signals in the tumors obtained by IVIS increased continuously with the time interval increasing from 12 to 60 hours post-administration. Importantly, at each time point, the average intensity of luminescence signals from the tumor site in mice injected with 5ApCB was significantly higher than that of mice treated with unconjugated bacteria. Particularly, at 60 hours after injection, the relative signal intensity was 5-fold higher than that of the control group. To quantify the tumor colonization, the biodistribution of the bacteria was examined at the end point of the animal study. After euthanasia, the heart, kidney, liver, lung, spleen, and tumor tissues were collected and then homogenized before spreading onto LB agar plates for bacterial counting. Accordantly, the bacterial number in the tumor tissue sectioned from mice injected

with 5ApCB reached to 5-times higher than that of the control mice (**Figure 4c**). In contrast to the major organs, the distribution of bacteria in the tumor tissue was extraordinary, with around 99.9% of total bacteria colonizing inside the tumor. The overwhelming aggregation of EcN within the tumor tissue was ascribed to their preference for hypoxic, eutrophication, and the immunosuppressive microenvironment of tumors.[44] Speculatively, in comparison to undecorated EcN, the notably enhanced accumulation within the tumor tissue after injection with 5ApCB was attributed to the improved localization of bacteria following administration, which was resulted from the specific binding with tumor cells. To prove this assumption, the localization of EcN within the 4T1 tumor at a relative short period of time after injection was quantified by plate counting. As confirmed in **Figure 4d** and Supplementary **Figure S2**, the number of bacteria localized into the tumor at 12 hours after dosing with 5ApCB achieved  $1.75 \times 10^7$  CFU per gram tissue, which was 2-fold higher than that of unconjugated EcN. Namely, surface conjugation indeed offered a greater opportunity to localize tumor site through a ligand-receptor interaction between aptamer-decorated bacteria and the tumor cells.

### **Enhanced anticancer efficacy of ApCB**

To validate the versatility of our approach, we translated the surface conjugation of aptamers to VNP, a well-known strain that has been widely used for tumor biotherapy due to its high immunogenicity.[45] VNP were similarly conjugated with AS1411 (Supplementary **Figure S3**) and the antitumor efficacy of the resulting 5ApCB was

assessed in vivo. Mouse subcutaneous tumor model of 4T1 was employed and the obtained tumor-bearing mice were randomly divided into three groups ( $n = 5$ ) that were received with different treatments including PBS, VNP and 5ApCB. Each mouse in VNP and 5ApCB groups was intravenously injected with  $5 \times 10^5$  CFU of bacteria at day 0. A tumor inhibition study by concisely assessing tumor volume demonstrated that both VNP and 5ApCB could inhibit tumor growth compared to PBS group (**Figure 5a** and **b**). Expectedly, 5ApCB group showed significantly enhanced antitumor efficacy in comparison with VNP group, suggesting that the increased tumor localization enabled by surface conjugation of aptamers played an important role in tumor biotherapy. Furthermore, the variation of bodyweight recorded every other day after treatment revealed the limited side effects of the bacteria under the experimental dosage. As displayed in **Figure 5c**, although the bodyweight of mice dropped slightly after injection with the bacteria, the mice in both VNP and 5ApCB groups recovered rapidly as reflected by the return of the bodyweight to the same level of PBS group.

It has been reported that the colonization of *Salmonella* can provoke the immunosuppressive microenvironment of tumors and elicit antitumor immunity.[46] To disclose the bacterial-mediated immune responses, the activation of T cells in the tumor tissue was analyzed at the end of the treatment. Nuclear nonhistone protein Ki67 was used as a maker to detect the infiltration of CD3<sup>+</sup> T cells in the tumor.[47] As shown in **Figure 5d** and Supplementary **Figure S4a**, the proliferation of CD3<sup>+</sup> T cells in mice treated with 5ApCB presented an apparent increase in contrast to both

PBS and VNP groups. Moreover, treatment with 5ApCB significantly elevated the percentage of CD3<sup>+</sup>CD4<sup>+</sup> T cells (**Figure 5e** and Supplementary **Figure S4b**), which suggested that the aptamer-decorated bacteria could boost more efficient tumor infiltration of immunologic effector cells. The percentage of tumor cytokine interferon- $\gamma$  (IFN- $\gamma$ ) and tumor necrosis factor- $\alpha$  (TNF- $\alpha$ ), which predominantly regulates the cell-mediated immune responses,[48] was further examined and the results showed a striking increase in mice treated with 5ApCB (**Figure 5f** and **g**, Supplementary **Figure S4c** and **S5a**). Terminal deoxynucleotidyl transferase dUTP nick end labeling (TUNEL) staining depicted the highest level of tumor cell apoptosis induced by 5ApCB (**Figure 5h** and Supplementary **Figure S5b**). In addition, hematoxylin and eosin (H&E) staining exhibited prominent necrosis of the tumor tissues after treatment with the conjugated bacteria (**Figure 5i**). Taken together, the inhibition of tumor growth could be attributed to the aptamer-mediated enhanced localization in tumor site and the bacterial associated activation of antitumor immune responses.

### **Versatility of ApCB**

To explore the generality of this strategy as a versatile platform for different aptamers, TLS11a was selected and the corresponding tumor model of H22 hepatocellular carcinoma was used to evaluate the effectiveness of conjugating aptamers to bacterial surface.[49, 50] TLS11a based ApCB (T-ApCB) was synthesized similarly by co-incubating amino-functionalized TLS11a with VNP (**Figure 6a**). Details for the



preparation of T-ApCB were described in the Supplementary Information. Correspondingly, T-5ApCB were fabricated by incubation of  $1 \times 10^8$  CFU of VNP with 5 nmol of TLS11a. Flow cytometric analysis showed that H22 cells co-incubated with T-5ApCB presented stronger fluorescence intensity than those cultivated with VNP, demonstrating strengthened affinity of conjugated bacteria to bind with cancer cells (**Figure 6b**). We next evaluated the antitumor efficacy of the resulting T-5ApCB in H22 tumor-bearing mice. After a single intravenous injection of PBS,  $5 \times 10^5$  CFU of VNP or T-5ApCB at day 0 (tumor size  $\sim 100 \text{ mm}^3$ ), both the tumor size and body weight of the treated mice were recorded. As plotted in **Figure 6c** and **d**, T-5ApCB group appeared a significant inhibition of tumor growth in comparison with PBS and VNP groups, which could be explained by that surface conjugation of aptamers increased the intratumoral localization of bacteria. Survival rate of different groups further validated that treatment with T-5ApCB endowed H22 tumor-bearing mice with a significantly extended survival period, as reflected by that near 70% of the treated mice survived successfully in 36 days (**Figure 6e**). While, none of mice dosed with PBS or VNP could survive under the same conditions. In addition, the variation of body weight during treatment revealed that administration with bacteria slightly dropped mouse body weight, which recovered rapidly to the same level of PBS group (**Figure 6f**). The slight fluctuation in body weight implied the limited side effects of 5ApCB at the experimental dosage. Collectively, these results proposed the universality of this approach to prepare diverse ApCB and verified the ability of

ApCB to increase the localization of conjugated bacteria in solid tumors for enhanced biotherapy.

## CONCLUSION

In summary, we have reported a strategy of conjugating aptamers to bacterial surface for endowing bacteria with tumor-specific accumulation following administration. The conjugation is based on a single-step and cytocompatible amidation procedure, which is implementable to diverse Gram-negative bacteria. The grafting density of aptamers on bacterial surface can be simply controlled by tuning feed ratio and the attachment is capable to increase the resistance of anchored aptamers against degradation due to the increase of steric hindrance towards the enzymatic hydrolysis by nucleases. With an optimized number of  $2.8 \times 10^5$  aptamers per cell, the conjugated bacteria exhibit the highest specific affinity with tumor cells in vitro. Strikingly, the optimal bacteria reached near 2- and 5-fold higher colonization in tumor tissue at 12 and 60 hours after systemic administration in vivo. The therapeutic value in tumor biotherapy has been validated by injection with aptamer-conjugated attenuated *Salmonella* in both 4T1 and H22 tumor-bearing mice. As expected, the decorated bacteria display significantly enhanced antitumor efficacy, with effective infiltration of immunologic effector cells and expression of tumor cytokines inside the tumor. This work discloses how bacterial bioactivities can be manipulated by surface conjugation with aptamers. With potential to address the severe side effects and low efficacy associated with bacterial-based therapy, we anticipate the application of this

strategy as a versatile platform to develop various bacterial therapeutics for enhanced tumor treatment.

### **SUPPLEMENTARY DATA**

Supplementary data are available.

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### **AUTHOR CONTRIBUTIONS**

J.L. and W.T. supervised the project. J.L. conceived and designed the experiments with Z.G. Z.G., Z.C., R.L. and K.L. performed all experiments. All authors analysed and discussed the data. Z.G. and J.L. wrote the paper.

### **Conflict of interest**

The authors declare no conflict of interest.

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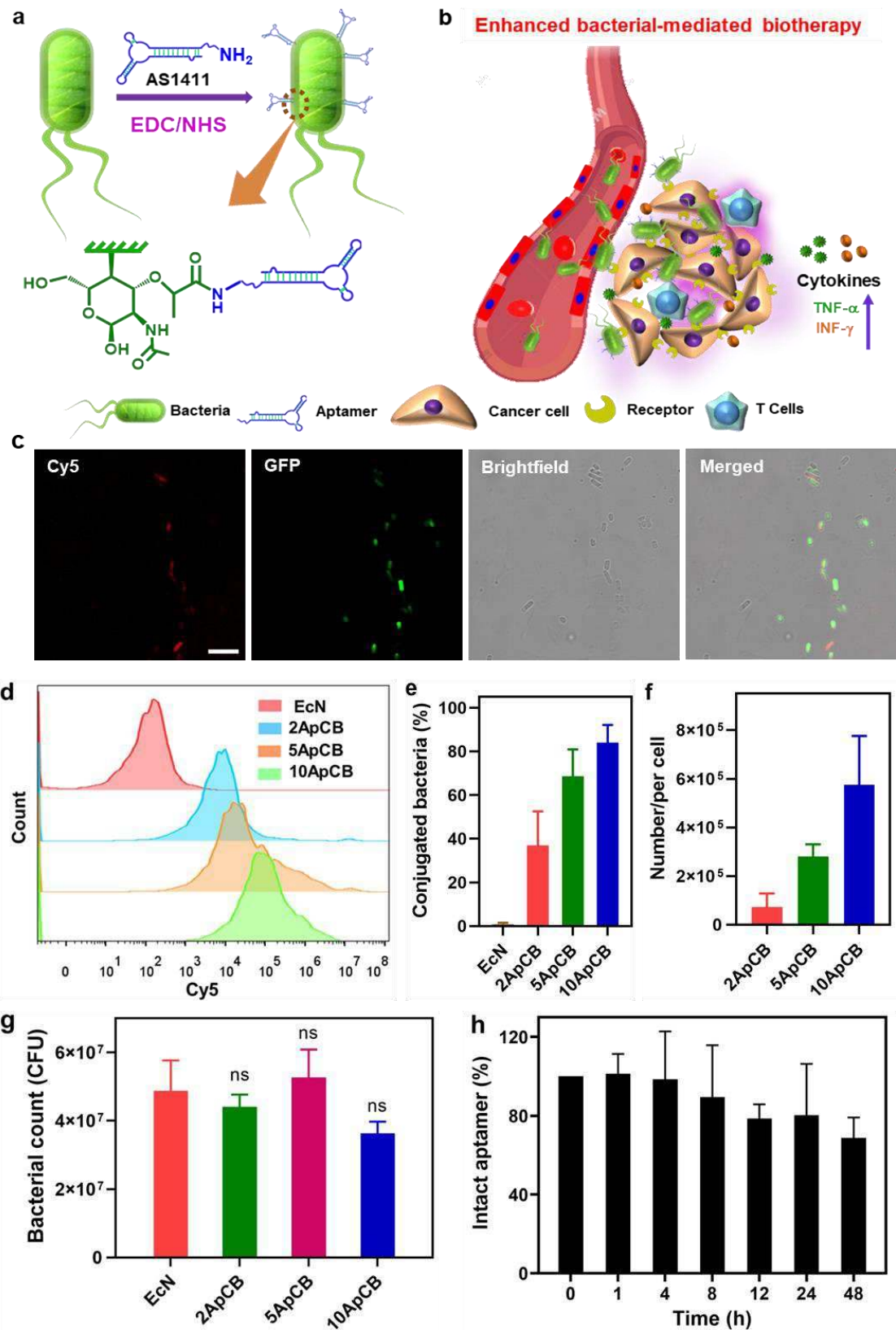
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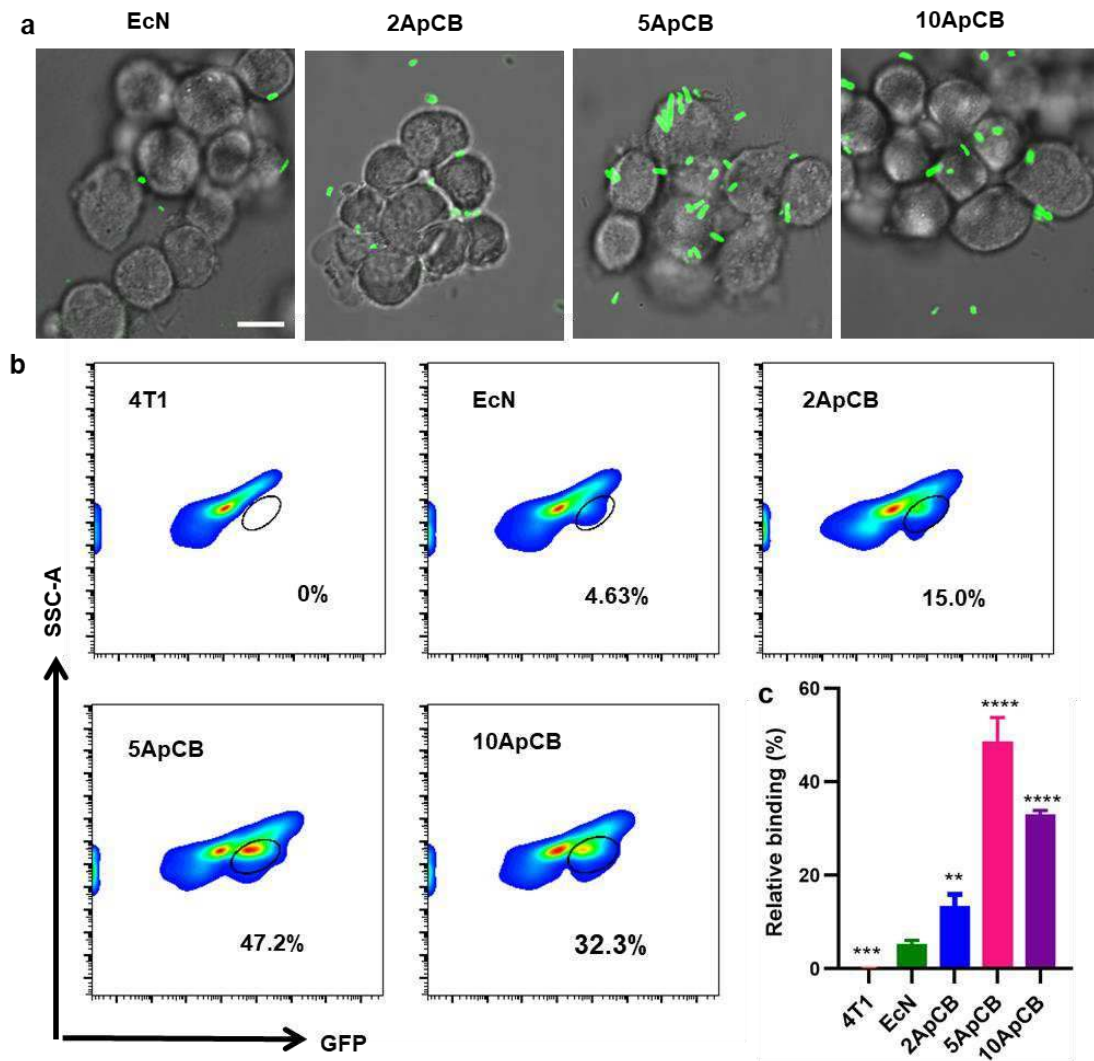
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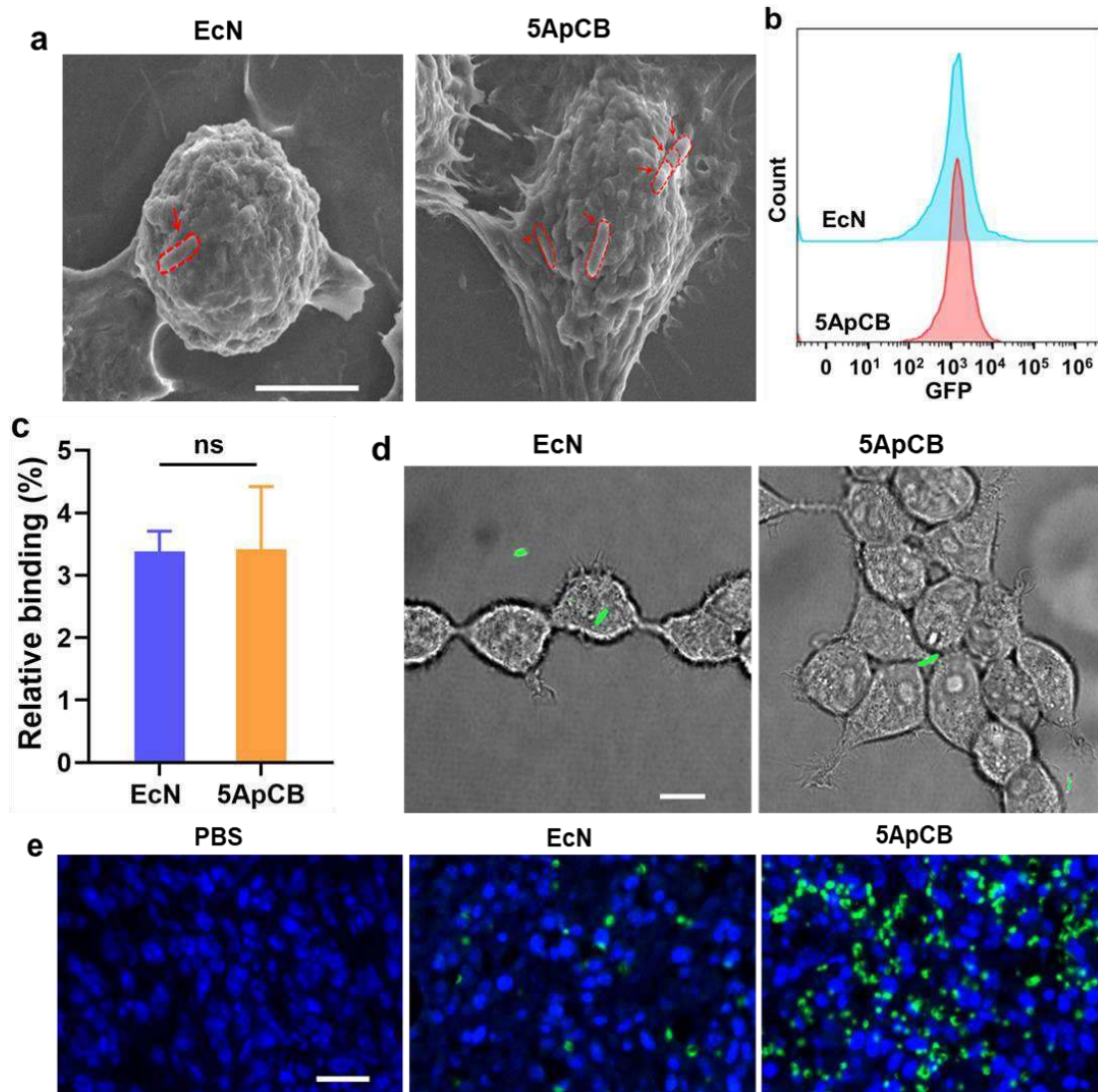


**Figure 1.** Design, preparation and characterization of ApCB. (a) Preparation of ApCB through amide condensation. (b) Aptamer-assisted tumor localization of bacteria for enhanced biotherapy. (c) Typical LSCM images of aptamer-conjugated bacteria. The red and green channels indicate aptamer conjugated with Cy5 and EcN producing

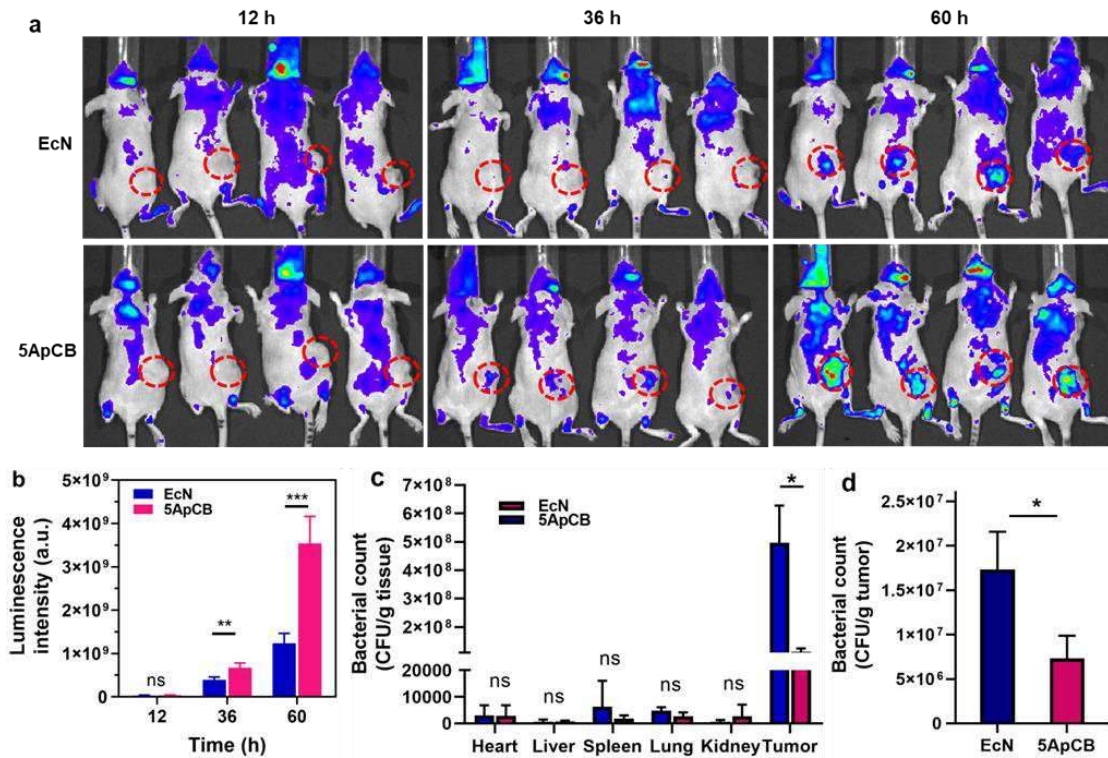
GFP, respectively. Scale bar: 10  $\mu\text{m}$ . (d) Flow cytometric analysis of EcN and EcN conjugated with Cy5-labelled AS1411. (e) Percentages of conjugated EcN under different feed ratios. (f) Average binding number of aptamers on each bacterial quantified by calculating the difference of fluorescent intensity of the aptamer solution after reaction. (g) Bacterial viabilities of EcN, 2ApCB, 5ApCB, and 10ApCB by LB agar plate counting. Plates were incubated at 37 °C for 24 hours prior to enumeration ( $n = 3$ ). ns: no significance. (h) Degradation kinetics of the conjugated AS1411 in 90% phosphate-buffered serum solution at 37 °C. ns: no significance.



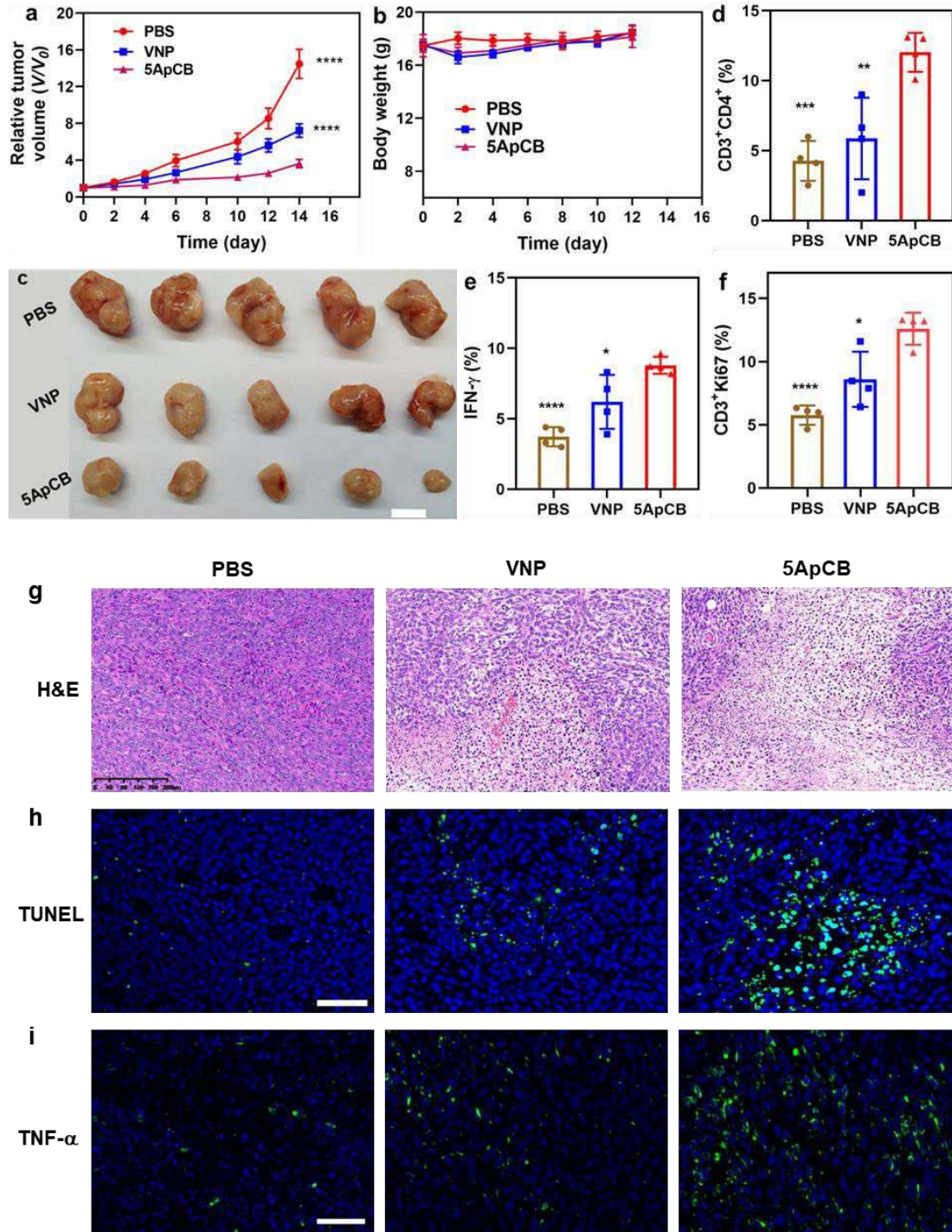
**Figure 2.** Binding of ApCB with 4T1 cells. (a) Representative LSCM images of 4T1 cells after incubation with EcN, 2ApCB, 5ApCB, and 10ApCB at 37 °C for 2 hours, respectively. Cells were rinsed with PBS before observation. The green channel means EcN producing GFP. Scale bar: 10  $\mu$ m. (b) Flow cytometric analysis of 4T1 cells after co-incubation with different conjugated EcN at 37 °C for 2 hours. (c) Percentages of 4T1 cells binding with ApCB after co-incubation at 37 °C for 2 hours. Error bars represent the standard deviation ( $n = 3$ ). Significance was assessed using Student's *t*-test, giving *p*-values, \*\* $p < 0.01$ , \*\*\* $p < 0.001$ , \*\*\*\* $p < 0.0001$ .



**Figure 3.** Specific binding of ApCB with cancer cells. (a) Representative SEM images of 4T1 cells after incubation with EcN and 5ApCB at 37 °C for 2 hours, respectively. Bacteria were circled by red dotted line. Scale bar: 5  $\mu$ m. (b) Flow cytometric analysis of 293T cells after co-incubation with EcN and 5ApCB at 37 °C for 2 hours, respectively. (c) Percentages of 293T cells binding with EcN and 5ApCB after co-incubation at 37 °C for 2 hours, respectively. Error bars represent the standard deviation ( $n = 3$ ). Significance was assessed using Student's  $t$ -test. ns: no significance. (d) Typical LSCM images of 293T cells after incubation with EcN and 5ApCB at 37 °C for 2 hours, respectively. Cells were rinsed with PBS before observation. Green channel means EcN producing GFP. Scale bar: 10  $\mu$ m. (e) Confocal images of tumor tissues sectioned at 12 hours after intravenous injection of  $1 \times 10^7$  CFU bacteria. Green and blue indicate EcN stained with Cyanine3-labelled anti-*Escherichia coli* and nuclei stained with DAPI. Scale bar: 20  $\mu$ m.



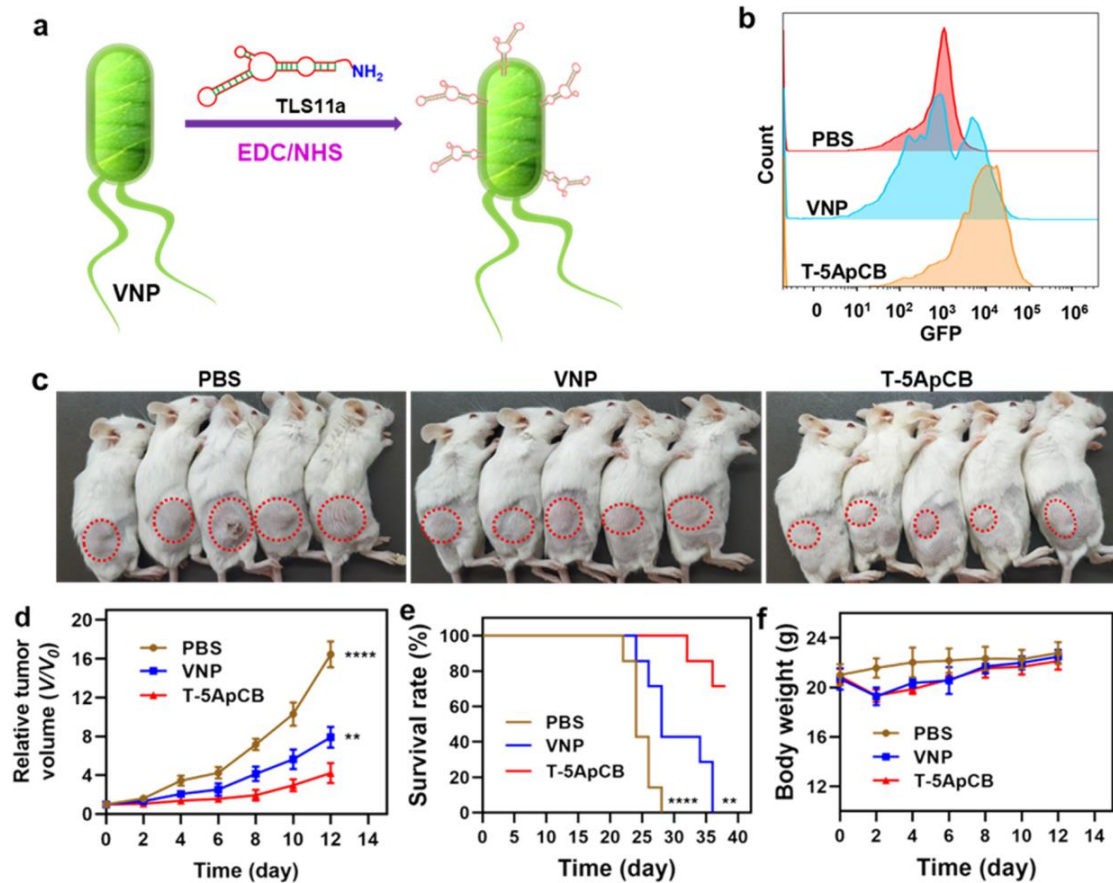
**Figure 4.** Biodistribution of ApCB in tumor-bearing mice. (a) IVIS imaging of 4T1 tumor-bearing mice at 12, 36, and 60 hours after injection with EcN and 5ApCB ( $1 \times 10^7$  CFU, expressing LuxCDABE) through the tail vein. (b) Average intensity of luminescence signals from the tumor site at 12, 36, and 60 hours after injection. (c) Biodistribution of bacteria in the major organs and tumor tissue of mice at 60 hours after injection with  $1 \times 10^7$  CFU of EcN or 5ApCB through the tail vein. (d) Numbers of EcN colonized within the tumor at 12 hours after tail vein injection of EcN or 5ApCB. Error bars represent the standard deviation ( $n = 4$ ). Significance was assessed using Student's *t*-test, giving *p*-values, \* $p < 0.05$ , \*\* $p < 0.01$ , \*\*\* $p < 0.001$ . ns: no significance.



**Figure 5.** In vivo therapeutic efficacy of ApCB. (a) Relative tumor growth after different treatments ( $n = 5$ ). (b) Representative photographs of the tumor tissues harvested from mice at the end of the treatment. Scale bar: 1 cm. (c) Variation of bodyweight during each treatment ( $n = 5$ ). Percentage of (d) Ki67 (gated on CD3<sup>+</sup> cells) inside the tumors after treatment ( $n = 4$ ). (e) Flow cytometric analysis of the population of CD4<sup>+</sup> T cells (gated on CD3<sup>+</sup> cells) within the tumors after treatment ( $n = 4$ ). Percentage of (f) IFN- $\gamma$  (gated on CD3<sup>+</sup> cells) inside the tumors after treatment ( $n = 4$ ). Images of tumor tissues stained with (g) TNF- $\alpha$  and (h) TUNEL. Scale bar: 50  $\mu$ m. (i) Typical H&E staining images of the sectioned tumors after different



treatments. Scale bar: 200  $\mu\text{m}$ . Significance was assessed using Student's *t*-test, giving *p*-values, \**p* < 0.05, \*\**p* < 0.01, \*\*\**p* < 0.001, \*\*\*\**p* < 0.0001.



**Figure 6.** Implementation of ApCB with aptamer TLS11a in a H22 tumor-bearing model. (a) Preparation of ApCB by conjugating VNP with TLS11a via amide condensation. (b) Flow cytometric analysis of H22 cells after co-incubation with PBS, VNP, and T-5ApCB at 37 °C for 1 hour. (c) Digital photos of mice at day 12 after treatment. Tumors were marked in red circles. H22 tumor-bearing mice (inoculated with  $1 \times 10^6$  carcinoma cells) were randomly divided into three groups and intravenously injected with PBS and  $5 \times 10^5$  CFU of VNP or T-5ApCB upon tumor size reaching at  $\sim 100$  mm<sup>3</sup> (defined as day 0). (d) Relative tumor growth after different treatments. (e) Survival curves of H22 tumor-bearing mice after different treatments. (f) Variation of body weight during treatment. Error bars represent the standard deviation ( $n = 7$ ). Significance was assessed using one-way ANOVA with Tukey post hoc test, giving  $p$  values, \*\*  $p < 0.01$ , \*\*\*\*  $p < 0.0001$ .

## Supplementary Files

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