Using Positively Charged Magnetic Nanoparticles to Capture Bacteria at Ultralow Concentration

Zhiming Li 1,2*, Jinyuan Ma 3, Jun Ruan 4 and Xuan Zhuang 2,5*

Abstract

Detecting bacteria at low concentrations without time-consuming culture processes would allow rapid diagnoses. Since electrostatic attraction exists between negatively charged bacterial cells and positively charged magnetic nanoparticles (NP+), capture of bacteria holds great promise towards achieving this goal. Here, we present a rapid and highly efficient approach to capture Escherichia coli, which was used as a model for gram-negative bacteria. Capture of E. coli at very low concentrations of 10 and 100 CFU/mL using NP+ is rapidly and efficiently achieved within 1 h. Moreover, the capture efficiency of NP+ was over 90% by analyzing the number of bacterial colonies on the plate. Optical and transmission electron microscopy confirmed the bacterial capture abilities of electrically charged nanoparticles (NPs). In contrast, negatively charged magnetic nanoparticles (NP−) did not show affinities towards E. coli. These results showed that bacterial cells, such as E. coli, carry a negative charge. Unlike a ligand-dependent capture system, our designed NP+ has potentials to capture a broad range of bacteria via electrostatic attractions.

Keywords: Bacterial charge, Electrostatic attraction, Magnetic nanoparticles, E. coli

Background

Infectious diseases are among the world’s most pressing health challenges. Microbial contamination of water resources is a major threat to public health. Escherichia coli (E. coli), a gram-negative bacterium, is very common in contaminated water and food. Some strains of E. coli can even cause serious bacterial infections. Bacteria at low concentrations are difficult to detect and usually require a pre-enriching process before further analysis. Culture-based microbiological methods are laborious and may take several days. Additionally, some bacterial strains may enter a viable but non-culturable state where they are viable but not culturable on routine agar, which impedes their detection by culture-based methods [1]. Inversely, rapid capture and decontamination of bacterial pathogens could provide real-time results to mitigate infectious disease outbreaks.

A variety of materials are developed for rapid capture and removal of bacteria from the contaminated source. Carbon nanotubes and resin-linked oligoacyllysine bead have been used to remove the bacteria from water [2, 3]. Magnetic nanoparticles, which can be conveniently separated from various resources by the employment of magnetic process, were widely used for bacteria detection and decontamination after functionalized with organic molecules [4–6]. The magnetic-based techniques have the advantages of target capture by time-saving (common separation time within 1 h), high recovery, possible automation, and scale-up separation [7]. The efficiency and selectivity of magnetic separation largely depends on the ligands, but sometimes it is hard to obtain a ligand with high affinity and specificity to the target. Therefore, it is necessary to develop a bacterial capture system with ligand-independent magnetic nanoparticles to capture the bacteria, especially under low concentrations.

Many scientists have investigated the nature of the electric charge of bacteria. Bechhold (1904) was the first to find the fact that bacterial cells carry a negative charge [8]. While it was already known that the large...
populations of bacterial cells tended to maintain a negative charge, little is known about the electrophysiology of bacteria at the level of single cells. In 2011, Cohen et al. revealed electrical spiking in *E. coli* at up to 1 Hz using a fluorescent voltage-indicating protein [9]. Since many kinds of bacterial cell walls are negatively charged, positive charged nanoparticles can strongly interact with a broad spectrum of bacteria via electrostatic interactions.

To take advantage of magnetic nanoparticles and negative charge of individual bacteria for fast pathogen detection, we designed a system to capture bacteria under low concentrations. Positively charged magnetic nanoparticles were fabricated by polyethylenimine (PEI), which is composed of abundant amine groups. Then we investigated the affinity of PEI functionalized nanoparticles against *E. coli*. This innovative method provides efficient binding to the bacteria by electrostatic interactions.

**Materials and Methods**

**Nanomaterials**

Iron (III) chloride hydrate (FeCl₃·6H₂O), ammonium hydroxide (NH₄OH, 28 wt%), hydrochloric acid (37 wt% aqueous solution), ethylene glycol, and sodium acetate were purchased from Shanghai (China) Reagent Company. Tetraethyl orthosilicate (TEOS), (3-aminopropyl)triethoxysilane (APTES), and fluorescein tetramethylrhodamine (TRITC) were purchased from Sigma-Aldrich (USA). Branched poly(ethylene imine) (PEI, 99%, *M*ₙ = 10,000) was purchased from Alfa Aesar. All the solutions were prepared using Milli-Q deionized water (18.2 MΩ cm at 25 °C resistivity).

**NP Syntheses**

Fe₃O₄ nanoparticles were prepared by a solvothermal reaction [10]. Briefly, 0.081 g of FeCl₃·6H₂O was dissolved in 30 mL of ethylene glycol under magnetic stirring. Then, 0.3 g of polyacrylic acid (PAA) and 1.8 g urea were added to this solution. After being stirred for 30 min, the solution was heated at 200 °C for 12 h by using a Teflon-lined stainless-steel autoclave. When cooled to room temperature, a black product, namely magnetic nanoparticle cores, was collected by a magnet. Followed by washing with ethanol and deionized water each three times, the Fe₃O₄ nanoparticles were treated with 0.15 M HCl under sonication for 15 min and then were coated with silica via hydrolysis and TEOS.

To prepare the negatively charged fluorescent magnetic nanoparticles (NP⁻), APTES-TRITC (C₃₃H₄₄N₃O₆Si) complex was first reacted under dark conditions overnight in ethanol. The complex was then grafted to the Fe₃O₄ nanoparticles through reaction between APTES and hydroxyl groups on the Fe₃O₄@SiO₂ nanoparticle. Subsequently, 30 μL of TEOS was added and reacted for 24 h in the dark. Followed by washing with ethanol and deionized water each three times, fluorescent NP⁻ were produced. Through the modification of NP⁻ with the polycation polymer PEI, the positively charged magnetic nanoparticles (NP⁺) were finished.

**NP Characterization**

Transmission electron microscopy (TEM) studies were performed by a TECNAI F−30 high-resolution transmission electron microscope operating at 300 kV. The particle size and zeta potential of NPs were determined by Malvern Zeta Nano series (Westborough, MA). Fluorescence was examined with a Carl Zeiss LSM5 EXITER laser scanning confocal microscope (Zeiss, Jena, Germany).

**Bacteria Preparation**

The gram-negative strain *E. coli* BL21 were used as the model bacteria. *E. coli* was cultivated in 100 mL of Luria Broth growth medium [11]. The bacteria were cultured in a thermostatic incubator at 200 rpm, 37 °C for 16 h. Subsequently, 100 μL of the bacterial culture was removed, diluted with the medium 1 × 10⁵ times, coated on the agar, and cultured at 37 °C for 24 h. The number of colony-forming units was counted. The remaining bacterial cells were harvested by centrifugation at 5000 rpm for 5 min, washed thrice with 1× PBS (10 mM, pH 7.4), and diluted to concentrations of approximately 1 × 10³ colony-forming unit (CFU)/mL. For safety considerations, all of the bacterial samples were placed in an autoclave at 121 °C for 20 min to kill bacteria before disposal and all glassware in contact with the bacteria was sterilized before and after use.

**Bacteria Capture Experiment**

All the batch capture studies were conducted in sterilized 1× PBS buffer. Forty microliters of NPs (1 μg/μL)
were dispersed in the sterilized saline under ultrasonication for 10 min, and then 1 mL of the bacterial solution (approximately $10^3$ CFU/mL) was added into the suspension. After incubation of 10 min, the NP-bounded bacteria were captured via a permanent magnet onto the wall of the vial, and free bacteria were removed with the wash solution. The captured bacteria were released by removing the magnet and resuspended in PBS. For microscopic analysis, an aliquot of bacteria was spread onto slides and stained with Hema-3 (Fisher Diagnostics). For immunofluorescence analysis, an aliquot of bacteria was spread onto slides and stained with 4′,6-diamidino-2-phenylindole (DAPI).

**Bacterial Capture Efficiency of NPs at Different Concentrations**

One milliliter of bacterial suspension (approximately $2 \times 10^2$ CFU/mL) was incubated with different amounts (5, 10, 20, 30, 40, 50, 75, and 100 μg/mL) of NP+ or NP− for 10 min. After magnetic separation by the nanoparticles, total solution was then sampled and analyzed for bacterial concentration via a plate counting method. The bacterial-capture efficiency of

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**Fig. 2** Characterization of the nanoparticles. **a** Transmission electronic microscopy (TEM) image of positively charged nanoparticles (NP+) and negatively charged nanoparticles (NP−). **b** Dynamic light scattering size and distribution of NPs. **c** Zeta potential distributions of NPs.
the NPs was tested by counting the number of CFU on the LB-agar plates.

**Capability of NPs to Capture Bacteria at Low Concentrations**

Forty micrograms of NP+ or NP− was incubated with 1 mL of bacterial suspension at very low concentrations (10 and 10^2 CFU/mL). After magnetic separation by the nanoparticles, total solution was then sampled and analyzed for bacterial concentration via a plate counting method. The bacterial-capture efficiency of the NPs was tested by counting the number of CFU on the LB-agar plates.

**Statistical Analysis**

Results were expressed as mean ± standard deviation (SD) as indicated in the figure legends. A two-way analysis of variance (ANOVA) with proper hoc analysis was calculated using GraphPad Prism software with P values < 0.05 considered statistically significant.

**Results**

**Characterization of Magnetic NPs**

The schematic diagram for preparation of the surface-charged magnetic composite nanoparticles is displayed in Fig. 1. The Fe₃O₄ nanoparticles are conjugated with APTES to form a thin layer of SiO₂ shell on the surface of nanoparticles upon reaction with TEOS and NH₄OH. To visualize and quantify captured cells directly, the APTES-TRITC complex is initially reacted, followed by grafting onto the surface of the Fe₃O₄@silica composites through a classical sol-gel reaction. Abundant SiOH groups govern the overall surface of this product, exhibiting a strong negative surface charge, namely negatively charged magnetic nanoparticles (NP−). For positively charged magnetic nanoparticles (NP+), PEI molecules are used to cover and modify the surface of NP−. The modified product shows a strong positive surface charge due to the abundant presence of amine groups.

As shown in Fig. 2a, TEM demonstrated that magnetic composite nanoparticles had a diameter of 450 nm, which were composed of uniformed SiO₂ coating (size of 60 nm). Dynamic light scattering (DLS) of the particles in Fig. 2b displayed a narrow size distribution with an increased average diameter after surface functionalization. The maximum sizes of the composite nanoparticles with the positive and negative charges are 620 and 700 nm, respectively. Nanoparticles measured by DLS are usually larger than those measured by TEM. This is because the DLS-assessed size is influenced by Brownian motion and depends on the ambient temperature, the dynamic radius of the nanoparticle, and the extent of nanoparticle agglomeration triggered by a static environment via the occurrence of confliction [12]. Figure 2c shows the zeta potential distributions of the negative and positive nanoparticles. In deionized water (pH 7.0), the zeta potentials of the NP− and NP+ are −26.6 mV and +28.1 mV, respectively. The pH dependences of zeta potentials for NP+ are depicted in Additional file 1: Figure S1. These results indicated that the surface-charged nanoparticles are well dispersed in aqueous solution under neutral conditions, which could be applied for cell capture.

**Ability of Magnetic NPs to Capture E. coli**

Figure 3 shows the general experimental procedure of bacteria capture. NPs were mixed with a solution of bacteria and incubated at room temperature for 10 min. Subsequently, we used a permanent magnet to capture the “magnetized” bacteria (magnetic nanoparticles bounded to the cell surface) onto the wall of the tube. After the removal of the remaining solution and washing the aggregates by PBS (with a magnet outside), we transferred the aggregates to a slide for microscopic analysis. As shown in the scheme, NP+ provides sufficient electrostatic responsiveness to quickly enrich E. coli. On the contrary, the bacteria are removed with rinsing PBS in the NP− experiment.

Optical images for the nanoparticles are shown in Fig. 4. Optical image of NP− showed the monodisperse and uniformly distributed particles. However, NP+ tended to agglomerate. The size distribution of NP+ is obviously wider than that of NP−. These results demonstrated that NP+ and NP− had a completely different pattern of interaction with E. coli. To further investigate the bacterial affinity of NP+, the localization NP+ in E. coli was examined using fluorescence analysis.
shows that captured E. coli are positive for both DAPI (blue color) and TRITC (red color). In order to clearly confirm the affinity of NP+ for bacteria, TEM technology was used. In Fig. 5b, a number of NP+ were observed to aggregate on the bacterial cell wall. These results suggested that NP+ have a strong affinity for bacteria.

Detection of Low Concentrations of E. coli
To further characterize the dynamics of the bacteria and NP+ interactions, we incubated E. coli at a constant number (2 × 10^2 CFU/mL) with various concentrations of NPs ranging from 5 to 100 μg/mL. The NP-bound bacteria were then magnetically captured and separated. The magnetic capture efficiencies of bacteria by NP+ and NP− are plotted as shown in Fig. 6. The number of E. coli captured by NP− is only 12% even at a high concentration of 100 μg/mL. In contrast with NP−, NP+ showed significant bacterial capture capacities and achieved 81% of capture efficiency with 40 μg/mL (P < 0.001). As can be seen from the LB-agar plates, a dose-dependent increase of bacterial colonies with NP+ was demonstrated.

In order to confirm NP+ affinity for E. coli at ultralow concentration, we mixed 40 μg NP with 1 ml PBS solution containing only 10 and 100 CFU of E. coli. Figure 7 shows photographs of the resulting colonies in agar plates for all samples. As expected, the NP+ indeed captured E. coli at an ultralow concentration, while the bacterial colonies were not obvious in plates using NP−.
The few colonies in the plate might be attributed to the non-specific affinity of nanoparticles. Further analysis indicated that over 90% bacterial capture efficiencies were obtained at an ultralow concentration (10 and 100 CFU/mL) using NP+. By contrast, the capture efficiency is less than 4% with NP− at the same conditions and has a significant difference (P < 0.001). These results suggest NP+ have a strong affinity for bacteria, which could be explained by the electrostatic attractions. To investigate the broad-spectrum bacterial capture properties, we employed three gram-positive bacteria (Bacillus subtilis, Staphylococcus aureus, and Lactococcus lactis) as models. As illustrated in Additional file 1: Figure S2, NP+ have a higher adsorption capacity for bacilli (E. coli and B. subtilis) than staphylococci (S. aureus) and streptococci (L. lactis). In addition, we also found that negatively charged molecules, such as 3-bromopyruvate (3-BP) and DNA, could interfere with the bacterial capture effect. The dead E.coli is invalid for such system (Additional file 1: Figure S3).

Discussion
It is known that both gram-negative bacteria (such as, E.coli) and gram-positive bacteria (such as B. subtilis) much more easily interact with positively charged particles than negatively charged particles via electrostatic attractions [13, 14]. This was also found in our study. One advantage in our capture system is that the PEI-functionalized nanoparticles have more amine groups, which were able to capture bacteria at ultralow concentrations. To date, there is a few general and satisfactory assays that could detect bacteria at concentrations of less than 10^2 CFU/mL without pre-enriching bacteria via a culture process. This study displayed a simple assay that uses electrically magnetic nanoparticles to capture and detect gram-negative bacteria (the organisms have a cytoplasmic membrane, a cell wall, and an intact outer membrane) within 1 h at a concentration of 10 CFU/mL.

In our experiment, we found that the surface charge of NPs can influence bacterial capture efficiencies. Here, the effects of charge at the surface of NPs on the bacterial capture efficiencies were studied using E.coli as a model bacteria. When NP+ were used in the capture assay, they exhibited efficient adsorptive ability of the bacteria. The bacterial capture efficiencies increased with the dosage of NP+. TEM microscopy shows macroscopic aggregates composed of nanoparticles and bacterial cells. In contrast, NP−, even at high concentrations, displayed low bacterial capture abilities. Overall, these observations

Fig. 6 Capture efficiencies of E. coli by NP+ or NP− at various concentrations indicated. The left image is the photograph of LB-agar plates coated with E. coli captured by NP+ and NP−. The right image shows the bacterial capture efficiency of NP at different concentrations indicated. E. coli (2 × 10^2 CFU in 1 mL PBS solution) without NP incubation was counted as 100% and served as control. *P < 0.05, **P < 0.01, ***P < 0.001.
demonstrated that the NP+ have a significantly higher capture ability than NP−.

Although gram-positive and gram-negative bacteria have differences in their membrane structure, most of them have a negative charge when cultivated at physiological pH values [15, 16]. Cell surface charge of bacterial cells has been characterized by electrostatic interaction chromatography (ESIC) [17]. The cell wall in gram-positive bacteria is mainly composed of a thick layer of peptidoglycan, which is embedded teichoic acid. On the other hand, the gram-negative bacteria have a layer of lipopolysaccharide at the external surface followed by a thin layer of peptidoglycan. The teichoic acid and lipopolysaccharides impart a negative charge to the surface of bacterial cells [18]. Previously, the positively charged silver nanoparticles (AgNPs) displayed the remarkable effectiveness against the microorganisms, including E. coli [19]. They found that the smaller particles are found to have greater antibacterial activity. We considered that larger particles may have a different benefit of bacterial absorbability. Firstly, larger particles difficultly reach the nuclear content of cells to cause the toxicity to the bacteria. Secondly, they can provide a greater surface area and therefore stronger bacterial interaction. Therefore, we designed and applied larger positively charged nanoparticles as a “sponge” agent to capture bacteria.

Conclusions
In conclusion, by PEI-magnetic nanoparticles, we have demonstrated a simple and fast assay to allow E. coli to be captured and analyzed. The existing archives of optical and TEM profiles of bacteria allow easy identification of captured bacteria. The high recovery provided by positively charged magnetic nanoparticles will allow detection of other bacteria strains at ultra-low concentrations.

Additional file

Additional file 1: Figure S1. The pH-dependent zeta potential and capture efficiency of the positive NPs. Figure S2. Effects of NP concentration on the capture efficiencies of four types of bacteria in PBS. Figure S3. Capture efficiency of the positive NPs at the different concentrations of 3-bromopyruvate (3-BP) (A), DNA (B), and the dead bacteria (C). (DOCX 424 kb)

Abbreviations
APTES: 3-Aminopropyl triethoxysilane; CFU: Colony-forming unit; DLS: Dynamic light scattering; E. coli: Escherichia coli; NP: Negatively charged magnetic nanoparticles; NP+: Positively charged magnetic nanoparticles; NPs: Nanoparticles; PAA: Polyacrylic acid; PEI: Polyethylenimine; SD: Standard deviation; TEM: Transmission electron microscopy; TEOS: Tetraethyl orthosilicate; TRITC: Tetramethylrhodamine

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Availability of Data and Materials
All data generated or analyzed during this study are included in this published article.

Authors’ Contributions
ZL is responsible for the conception and design and contribution of reagents and wrote the paper. ZL and JR conducted the experiments. JM
and XZ designed the figures. All authors read and approved the final manuscript.

Competing Interests
The authors declare that they have no competing interests.

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Author details
1Institute of Reproductive Health, Tongji Medical College, Huazhong University of Science and Technology, Wuhan 430030, Hubei, China. 2Department of Urology, the First Affiliated Hospital of Xiamen University, Xiamen 361003, Fujian, China. 3Department of Pharmacy, Shanghai Dermatology Hospital, Shanghai 200443, China. 4College of Life Sciences, Central China Normal University, Wuhan 430079, China. 5Department of Clinical Medicine, Fujian Medical University, Fuzhou 350005, Fujian Province, China.

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