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Macrophage-red blood cell hybrid membrane-coated ultrasound-responsive microbowls to eliminate pathogens, endotoxins, and heavy metal ions from blood†

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Sepsis is a potentially life-threatening condition triggered by pathogens such as bacteria and toxins. In clinical settings, hemoperfusion has been used as a therapeutic technique for the treatment of sepsis to remove pathogens, toxins and other inflammatory mediators from the bloodstream. However, the existing adsorption process of hemoperfusion is not specific to sepsis-related pathogens or toxins, resulting in unsatisfactory removal effectiveness and the lack of capture selectivity. Herein, we developed an ultrasound-responsive asymmetric polymeric microbowl coated with macrophage and red blood cell (RBC) membranes to selectively eliminate pathogens, toxins, and heavy metal ions from blood. Poly(glycidyl methacrylate) spherical microparticles were first synthesized and then eroded using dibutyl phthalate and styrene to form a microbowl structure. This asymmetric structure could provide acoustic radiation pressure gradients to realize orientation motion under non-invasive ultrasound fields with a velocity of 37.14 $\mu m s^{-1}$ at 20 V_{p-p} , which could increase adsorption efficiency in physiological environments by fuel-free programable movements. In addition, the macrophage and RBC membrane coating endowed the microbowl with desired biocompatibility, enhanced blood circulation and the ability to selectively eliminate endotoxins, bacteria and heavy metal ions. Endotoxin removal assay showed that 92.80% of endotoxin was eliminated within 15 minutes. Owing to the affinity between the red blood cell membrane and heavy metal ions, the removal efficiency reached 92.75% and 93.91% for Pb²⁺ and Hg²⁺, respectively. Meanwhile, over 90% of S. aureus in the blood was eliminated owing to the selective recognition and adhesion of bacteria by macrophage membrane proteins. Overall, this work on ultrasound-responsive asymmetric polymeric microbowls provides a new insight to eliminate pathogens, toxins and heavy metal ions from the bloodstream for sepsis treatment.

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1. Introduction

Sepsis, also referred to as blood poisoning, is a classic disease that is characterised by the dysregulation of the body's response to infection and leads to organ dysfunction, thereby threatening the lives of approximately 50 million people each year. 1,2 Patients with sepsis usually require blood purification

such as hemoperfusion to remove pathogens and related toxins.3-5 Localized infection and pathogen entry into the bloodstream would activate certain pattern recognition receptors on the surface of immune cells and lead to an overwhelming and dysregulated inflammatory reaction throughout the body.6 Additionally, heavy metal ions, such as lead and mercury, can augment the expression of pro-inflammatory cytokines through various mechanisms, alongside the generation of reactive oxygen species, thereby exacerbating the inflammatory response.⁷⁻⁹ As a commonly applied approach for sepsis treatment, hemoperfusion is a technique for blood purification that restores physiological functions by adsorbing harmful components in the blood through hemoperfusion adsorbents in perfusion columns.10 Current hemoperfusion adsorbents are typically composed of activated carbon or resin, both of which offer a large surface area for the physical adsorption of toxins. 11 Recently, protein materials have paved a novel path in the

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design of novel hemoperfusion adsorbents. 12,13 Yang et al. developed a biocompatible protein-polysaccharide complex that exhibited multiple-removal ability for liver and kidney metabolic wastes, toxic metal ions, and antibiotics, which has significantly promoted the development and applications of protein materials in hemoperfusion adsorbents.¹⁴ Nowadays, the development of hemoperfusion adsorbents with high removal efficiency and excellent selective adsorption is particularly crucial. 15 Therefore, it would be of great value to develop a new approach of blood purification to specifically and efficiently eliminate bacteria, endotoxins, and heavy metal ions for enhanced therapeutic outcomes.16

To increase the adsorption efficiency, self-propelled nano/ microparticles guided by external fields have been applied to remove contaminants such as micro/nanoplastics, organic molecules, and oil spills.¹⁷ For example, Katherine et al. developed BiVO₄ microparticles to achieve individual and collective swimming under visible light irradiation, which could selectively adhere with yeast cell walls for fungicidal activity. 18 In addition, Qiu et al. synthesized asymmetric magnetic mesoporous silica nanobottles loaded with Fe₃O₄ nanoparticles to effectively adsorb and remove heavy metal ions with precise directional control and recyclability under external magnetic fields. 19 Compared to the previously reported light and magnetic field as driving forces, ultrasound (US) has been widely applied clinically in a non-invasive manner, which could ensure cellular and tissue integrity with enhanced penetration into deeper tissues and navigation under intricate physiological environments.²⁰ In addition, compared to isotropic symmetric micro/nanoparticles, micro/nanoparticles with asymmetric structures exhibit different properties in a single particle, such as polar and non-polar, hydrophilic and hydrophobic, and anionic and cationic owing to their non-uniform shapes and surface properties, 21,22 which makes it possible to precisely control and direct movements under external fields in clinical settings. Currently, multiple approaches have been developed to fabricate asymmetric microstructures. Most of them were fabricated by template-assisted electrodeposition. For example, Ahmed et al. reported asymmetric metallic and bimetallic nanorods fabricated by electrochemical deposition within the pores of anodic alumina membranes.²³ He et al. reported asymmetric ultrasound-driven nanoswimmers with goldnanoshell-functionalized multilayer tubular structures using a template-assisted electrodeposition protocol.²⁴ For polymerbased asymmetric nanoparticles, our groups have previously synthesized and manipulated the opening concaves of nanobowls via optimizing hydrogen bonding and π - π interactions during the homopolymer self-assembly ²⁵ and co-assembly of homopolymers and peptides.26 However, the existing asymmetric micro/nanoparticles are primarily designed to enhance the movement efficiency, which lack desired bioactivity for biomedical applications.

To endow nanomaterials with designated biofunctions, functionalization with various cell membranes containing a series of membrane proteins has been reported. Zhang et al. developed biodegradable polymeric nanoparticles coated with

natural RBC membranes to enhance their circulation time for cargo delivery through bypassing macrophage uptake and systemic clearance.27 Wang et al. designed macrophage membranecoated gold-silver nanocages for efficient bacterial targeting and serving as a drug delivery vehicle, demonstrating improved bacterial adherence, prolonged circulation, and excellent biocompatibility.²⁸ Recent studies have begun to shed light on the synthesis, characterization, and applications of asymmetric micro/nanoparticles coated with cell membranes.29 Such hybrid nanomaterials leverage the inherent characteristics of both components, allowing for tailored functionalities and enhanced performance in bioapplications such as drug delivery, biosensing, and targeted therapeutics. For instance, He et al. developed near-infrared light-powered Janus mesoporous silica nanoparticles coated with macrophage cell membranes for active targeting cancer cells and thermomechanical perforation of cancer cell membranes.³⁰ However, biofunctionalization of asymmetric polymeric micro/nanoparticles remains an area with limited exploration but holds immense promise.

In this work, we developed an ultrasound-driven asymmetric microbowl coated with hybrid cell membranes to scavenge endotoxins, bacteria and some heavy metal ions (Pb2+ and Hg2+). The asymmetric structure of the polymeric microbowl was transformed from poly(glycidyl methacrylate) (PGMA) microspheres. The microspheres were prepared by dispersion polymerization and then the surfaces of the microspheres were eroded by dibutyl phthalate and styrene to form an asymmetric bowl-like structure. The outside diameter of the microbowl was $\sim 1.3 \mu m$ and the inside opening diameter was ~ 500 nm. The RBC membrane was isolated from erythrocytes using the hypotonic treatment method, and the macrophage cell membrane was extracted from RAW 264.7 via a repeated freeze-thawing process. The RBC and macrophage cell membranes were co-encapsulated onto the surface of the microbowls by repeated extrusion of 3 µm-porous membranes through a mini-extruder with the same membrane protein content. The movement of this polymeric microbowl could be initiated and controlled under an ultrasound field at 1.46 MHz, and its velocity was affected by the amplitude of the sinusoidal ultrasound with the highest speed of 37.14 $\mu m\ s^{-1}$ at 20 V_{p-p} . Heavy metal ions were adsorbed and removed by attaching to the RBC membrane proteins coated on the microbowl surface. 31-33 The clearance efficiency of Pb2+ and Hg2+ was more than 90% respectively. Meanwhile, macrophage membrane proteins could specifically recognize and adhere to endotoxins and bacteria, which resulted in a bacteria removal rate of over 90%. In general, this ultrasound-driven membrane-coated microbowl demonstrated an effective approach to scavenge endotoxins, bacteria and some heavy metal ions during hemoperfusion.

2. Experimental

2.1. Materials

Glycidyl methacrylate (GMA, 99%), azobisisobutyronitrile (AIBN), polyvinylpyrrolidone (PVP), dibutyl phthalate (99%), sodium NJC

dodecyl sulfate (SDS), styrene (99.5%) and anhydrous ethanol were used without further purification. HgCl2, PbCl2, methanol, dioxane and sodium hydroxide (NaOH) were purchased from Shanghai Aladdin Biochemical Technology Co., Ltd. Phosphate buffered saline (PBS) and Tris-magnesium buffer (TM buffer) were purchased from Beyotime Biotech. Inc. L02 liver cells and L929 cells were obtained from the Institute of Biochemistry and Cell Biology, Chinese Academy of Sciences (Shanghai, China). Cell Proliferation and Cytotoxicity Assay (CCK-8) Kit, fluorescein isothiocyanate (FITC), rhodamine B and Coomassie Brilliant Blue G-250 were purchased from Beijing Solarbio Science & Technology Co., Ltd (Beijing, China).

Water was processed using a Millipore purification system (Darmstadt, Germany) with a minimum resistivity of 18.2 M Ω cm.

2.2. Synthesis of PGMA seed microspheres

PGMA seed microspheres were prepared via dispersion polymerization.³⁴ Briefly, 4.0 g of PVP was dissolved adequately in anhydrous ethanol and double-distilled water mixed solutions in 10/1 volume ratio. Then, 300 mg of AIBN dissolved in 2.0 g of GMA was then added. After purging for 20 min with nitrogen, the system was heated to 70 °C for 12 h of reaction. The obtained product was washed three times with ethanol and double-distilled water. Finally, the residue was freeze-dried to obtain the PGMA seed microspheres.

2.3. Fabrication of microbowls

First, 50 mg of PGMA seed microspheres were dispersed in 5 mL of 2.5 wt% SDS aqueous solution under magnetic stirring. Subsequently, 250 mg of dibutyl phthalate and 250 mg of styrene were added. After being eroded at 40 °C for 24 h, the microbowls were obtained and purified by centrifugation and lyophilization.

2.4. Extraction of the red blood cell (RBC) membrane

The RBC membrane was obtained by hypotonic treatment from 4% mouse erythrocyte.35 The RBC was washed with ice-cold PBS, obtained by centrifuging at 700g for 5 min at 4 °C, and then treated with an ice-cold hypotonic buffer (10% PBS) at 4 °C for 6 h to burst the RBC membrane. Unbroken RBCs were then removed by centrifugation at 1000g for 10 min. The remaining supernatant was centrifuged at 14 000g for 30 min to obtain the pink RBC membrane. The protein content in the RBC membrane was determined by the Bradford protein assay (Shanghai Sangon Biotech. Co. Ltd, China). The obtained RBC membrane was stored at -80 °C for later study.

2.5. Extraction of the macrophage cell membrane

The macrophage cell membrane was isolated from RAW 264.7 cells as previously reported with slight modifications.³⁶ In brief, RAW 264.7 cells were cultured and harvested at a concentration of 5×10^7 cells per mL by centrifugation. The cells were resuspended and incubated for 15 min in an ice-cold Trismagnesium buffer (TM buffer, pH = 7.4, 50 mM Tris-HCl, 8 mM MgSO₄) containing a protease inhibitor. Cells were fragmented by a repeated freeze-thawing process. Then, the

cell homogenate was centrifuged at 700g at 4 °C for 10 minutes to remove the unbroken cells and nuclei. The supernatant was centrifuged at 14 000g at 4 °C for 30 min to obtain the macrophage membranes. The cell membrane fragments were washed by centrifugation at 14 000g at 4 $^{\circ}$ C and redispersed in 1 mL PBS solution. The protein content in the purified macrophage membrane was also determined by the Bradford protein assay. The obtained macrophage membrane was stored at -80 °C for later study.

2.6. Measurement of the membrane protein content

The protein content of the RBC membrane and macrophage membrane was measured by the Bradford protein assay. 37 First, 20 µL of BSA solutions with different concentrations (20, 40, 60, 80, 100, and 150 $\mu g \text{ mL}^{-1}$) were added into a 96-well plate. Then, 200 µL Coomassie Brilliant Blue G-250 solution was added and incubated for 10 min. The same 20 µL of RBC membrane or macrophage membrane solutions were incubated with 200 µL Coomassie Brilliant Blue G-250 solution. The absorbances of all samples were detected using a UV-Vis microplate reader. The membrane concentrations were calculated based on the calibration curve of standard samples.

2.7. Fabrication of RBC-MP-microbowls

RBC-MP-microbowls were prepared by enclosing microbowls with the obtained RBC membrane and macrophage membrane. In brief, two kinds of membranes were fused in PBS in 1/1 protein weight ratio under ultrasonication (42 kHz, 200 W) for 10 min. Furthermore, microbowls were incubated with hybrid membranes under ultrasonication for another 10 min and subsequently extruded through 3.0 µm polycarbonate porous membranes for 20 times using an Avanti mini extruder. The resulting RBC-MP-microbowls were washed three times with PBS and obtained by centrifugation at 5000 rpm.

Characterization of RBC-MP-microbowls

Scanning electron microscopy (SEM). The morphologies of PGMA microspheres, microbowls and RBC-MP-microbowls were characterized using a Zeiss Sigma 300 VP instrument. Images were recorded with the secondary electron mode at 2-5 kV.

Confocal laser scanning microscopy (CLSM). Rhodamine B-labelled RBC membranes and FTIC-labelled macrophage cell membranes were observed using a Nikon Ti2 confocal laser scanning microscope (Nikon, Japan).

Optical microscopy. The movement of the microbowl was recorded using an Olympus BX60 optical microscope.

A Microplate Reader (Multiskan™FC, Thermo Fisher Scientific) was used to perform the Bradford assay and confirm the cell viability.

Fluorescence spectroscopy. Fluorescence emission spectra of RBC-MP-microbowls and microbowls were measured using a Fluorescence Spectrophotometer F-4700 (Hitachi, Ltd, Japan). The excitation and emission slit widths were both set to 5 nm. The scan speed was set to 60 nm min⁻¹. Fluorescence emission spectra were recorded at a fixed time interval.

Inductively coupled plasma-optical emission spectrometer (ICP-OES). To evaluate the heavy metal removal efficiency of RBC-MP-microbowls, the concentrations of heavy metal ions (Hg²⁺ and Pb²⁺) were measured by ICP-OES (PerkinElmer 8300).

Fluorescence microplate reader. The relative fluorescence unit of different samples was measured using a multiple microplate reader of Infinite 200pro (Tecan Austria GmbH). The excitation wavelength and emission wavelength were set at 360 nm and 465 nm with an emission bandwidth of 35 nm.

2.9. Cell membrane detection of RBC-MP-microbowls

To confirm that the hybrid membrane encapsulated the microbowls successfully, the protein content of the RBC-MPmicrobowls was quantitatively measured by the Bradford assay (Shanghai Sangon Biotech, Co. Ltd, China). In brief, the microbowls (1 mg mL⁻¹) and RBC-MP-microbowls (1 mg mL⁻¹) were centrifuged and washed three times with PBS. Subsequently, 20 μ L of the microbowls (1 mg mL⁻¹) and RBC-MP-microbowls (1 mg mL⁻¹) were mixed with 200 μL Coomassie Brilliant Blue G-250 solution. After 10 min incubation, the absorbance was detected and the protein contents were calculated based on the calibration curve of standard samples. Additionally, the membrane protein series of both hybrid membranes and RBC-MP-microbowls were compared with sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) protein characterization. The samples were heated at 100 °C for 5 min and run at 180 V for 45 min, followed by Coomassie Blue staining overnight. After washing, the final image was captured and recorded.

To further confirm the presence of the hybrid membranes onto the surface of the microbowls, the RBC membrane and macrophage membrane were labelled with rhodamine B (excitation/ emission = 546/568 nm) and FITC (excitation/emission = 495/ 525 nm) respectively before coating onto the microbowls. The fluorescence images were acquired using a CLSM with a 10× microscope eyepiece and a 60× microscope objective.

2.10. Acoustic propulsion studies

The acoustic equipment to propel the RBC-MP-microbowls was built with reference to previous work.²⁴ A rectangle cell (10 mm × 10 mm) was home-made by applying two layers of polyimide Kapton tape (50 µm thickness per layer) to a conductive glass slide (40 mm \times 40 mm \times 2 mm). The solution in the cell was excited by a piezoelectric ceramic disc transducer affixed to the conductive glass slide parallel to the acoustic cell (Shenzhen Huajingda Electronics Co., Ltd, catalog no. H4P163000). The piezoelectric ceramic was connected to a function generator system, including a signal generator (FEEL TECH Inc., FY2300) and a signal amplifier (FEEL TECH Inc., FYA2010s).

The Olympus BX60 optical microscope and a commercial video-capturing camera were used for observing the movements of RBC-MP-microbowls and recording videos. Videos of motion were taken at 400× magnification at a frame rate of 25 s⁻¹. The motion was tracked using a Video Spot Tracker. The speed was calculated by dividing the displacement of the microbowl centre between two frames by the time interval, and

then the average of the speed over the selected tracking period was taken. The tracking was repeated with multiple microbowls to ensure statistically robust results. We verified the velocity of the microbowls in whole blood with reference to previous work.³⁸ We replaced the liquid environment with whole blood spiked with anticoagulants and compared the velocity of microbowls with that in PBS.

2.11. In vitro cell viability assay

A CCK-8 assay was performed to assess the microbowls against L929 and B16 cells. Briefly, approximately 1.0×10^5 cells per well with an MEM medium were seeded into 96-well plates. After culturing for 24 h, the primary MEM medium was replaced with a medium containing microbowls (0, 62.5, 125, 250, 500, and 1000 μg mL⁻¹) at designated concentrations for another 24 h. Then, 20 μL of CCK-8 solution was then fed into each well, incubating for an additional 1 h. The absorbance of each sample was measured using a microplate reader at 450 nm. The cell viability was calculated according to the following formula:

$$\label{eq:cell_control} \text{Cell viability (\%)} = \frac{[\text{OD}_{450}(\text{sample}) - \text{OD}_{450}(\text{blank})]}{[\text{OD}_{450}(\text{control}) - \text{OD}_{450}(\text{blank})]} \times 100\%$$

2.12. Hemolysis assay

Hemolysis assays were performed in accordance with a previously published method. First, 2.0 mL fresh blood was drawn from mouse eyes and centrifuged (3000 rpm for 10 min), and then RBCs were resuspended in PBS to prepare a washed RBC suspension. The cells were exposed to water and different concentrations of RBC-MP-microbowls (0, 12.5, 25, 50 and 100 μ g mL⁻¹) for 30 min. After that, the mixture was centrifuged for 10 min at 4 °C, and the absorbance of the supernatant was measured using a microplate reader at an absorbance wavelength of 560 nm. The degree of hemolysis was calculated using the following formula:

$$Hemolysis~(\%) = \frac{[OD_{560}(sample) - OD_{560}(blank)]}{[OD_{560}(water) - OD_{560}(blank)]} \times 100\%$$

2.13. Removal of heavy metal ions using RBC-MP-microbowls

The performance of RBC-MP-microbowls for the removal of heavy metal ions was evaluated using aqueous solutions containing Pb^{2+} and Hg^{2+} ($\sim 10 \text{ mg L}^{-1}$), respectively. The concentration of heavy metal ions in a solution was measured using an Inductively Coupled Plasma-Optical Emission Spectrometer (ICP-OES, PerkinElmer 8300). Briefly, 100 μL of RBC-MPmicrobowls (5 mg mL⁻¹) were added to 1 mL of aqueous solutions containing metal ions. In order to verify the effect of motion excited by US on the removal of heavy metal ions, another set with the same conditions was treated with US. As a control, 100 µL of deionized water was added to 1 mL of aqueous solutions containing metal ions. After adsorption for 1 h, the concentration of ions in the supernatant obtained by centrifugation and filtration through a poly(ether sulfone) (PES, 0.22 µm) membrane was measured by ICP-OES. Each set was repeated three times.

Similarly, the adsorption efficiencies of RBC-MP-microbowls were investigated for different concentrations of heavy metal ions to better study their adsorption capacity for Pb²⁺ and Hg²⁺. The ion concentrations were set at 100-200 μ g L⁻¹ for Pb²⁺ and $50-150 \mu g L^{-1}$ for Hg^{2+} . The setting ranges for these concentrations are based on the safe concentrations for heavy metal poisoning. 39,40 The heavy metal ion concentrations of samples before and after adsorption were determined by ICP-OES.

2.14. Removal of endotoxins using RBC-MP-microbowls

The performance of RBC-MP-microbowls in endotoxin removal was evaluated by a Recombinant Factor C Endpoint Fluorescent Assay. First, the standard curve was fitted with standard samples ranging from 0.005 EU per mL to 5 EU per mL to determine the correspondence between relative fluorescence unit (Δ RFU) and endotoxin concentration. The Δ RFU was determined using a fluorescence Microplate Reader (Tecan Austria GmbH). ΔRFU was calculated using the following equation:

$$\Delta RFU = RFU_{T=1h} - RFU_{T=0}$$

After that, 100 μL of RBC-MP-microbowls (5 mg mL⁻¹) were mixed with 900 µL of endotoxin solution (~3 EU per mL) for different incubation periods (15, 30, 45, and 60 min). The same 100 μL of DI water was mixed with 900 μL of endotoxin solution (\sim 3 EU per mL) as the control set. The RFU was recorded at T = 0 and T = 1 h respectively. The corresponding endotoxin concentration was calculated using the fitted standard curves. The role of US on endotoxin adsorption was similarly investigated by setting ion concentrations of 0.5 and 1.0 EU per mL and with and without US conditions.

2.15. Binding and isolation of macrophage-adhering bacteria

The bioactivity of RBC-MP-microbowls was assessed in binding and isolation of bacteria using Staphylococcus aureus (S. aureus, ATCC 29213) as model bacteria.41 S. aureus was cultured in a Luria-Bertani medium (LB medium, 0.10 g mL⁻¹) in an oven at 37 °C for 12 h. The bacterial suspension (100 μL) was added into the PBS (9.9 mL) to make a diluted bacterial suspension. Then, 100 μL of RBC-MP-microbowls (5 mg mL⁻¹) were mixed with the above-mentioned bacterial suspension (900 μL). Similarly, 100 µL of PBS was mixed with the bacterial suspension (900 µL) and set as the control group. After the US incubation for 3 h in a US field (1.46 MHz, 20.0 V_{p-p}), bacterial suspensions from different groups were spread onto plates to continue incubation for 12 h, and then colony growth was recorded.

3. Results and discussion

Preparation and characterization of RBC-MP-microbowls

The RBC-MP-microbowls were prepared in two stages of evolving PGMA microspheres to microbowls and hybrid cell membrane encapsulation, as illustrated in Fig. 1. Previous work had reported that the asymmetric structure would be beneficial for ultrasound-induced propulsion.²³ However, it is still challenging to prepare asymmetric microparticles with an

adjustable concave size. In order to prepare asymmetric micro/nanoparticles, the PGMA microspheres were first fabricated by dispersion polymerization. The spherical morphology with an average diameter of approximately 1.14 µm was observed using an SEM (Fig. 2A and Fig. S1, ESI†). Afterwards, a bowl-like structure was prepared by an eroding strategy with dibutyl phthalate and styrene. In the presence of surfactant sodium dodecyl sulfate, organic droplets composed of dibutyl phthalate and styrene adsorbed onto the surface of PGMA microspheres. 42 Gradually, the organic phase eroded the surface of the microspheres, creating a concave shape and ultimately forming a unique bowl-like structure (Fig. 2B). The outer diameter of the microbowl was approximately 1.29 µm and the inner diameter was approximately 562.5 nm (Fig. S2, ESI†).

After the asymmetric microbowl was prepared, the hybrid cell membrane was coated onto microbowls using an Avanti mini extruder. The RBC membrane was isolated from erythrocytes using the hypotonic treatment method, 43 and the macrophage cell membrane was extracted from RAW 264.7 via a repeated freeze-thawing process.36 Additionally, the membrane protein contents of RBC membranes and macrophage membranes (Fig. S3, ESI†) were 122.27 $\mu m mL^{-1}$ and 87.99 $\mu m mL^{-1}$ respectively. The RBC-MP hybrid membranes were prepared in 1/1 protein weight ratio. After incubation for 20 min, the RBC-MP hybrid membranes and microbowls were extruded through polycarbonate porous membranes for 20 times with 1 mg mL⁻¹ of total membrane proteins. A sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) assay was then performed to compare the protein type in the RBC membranes, macrophage membranes, RBC-microbowls, MPmicrobowls and RBC-MP-microbowls. As shown in Fig. 2C, while a negligible protein signal was found in the bare microbowls, the protein composition in the RBC-MP hybrid membrane was primarily retained in the RBC-MP-microbowls. In order to characterize the fusion and coating level of hybridized membranes onto microbowls, fluorescein isothiocyanate (FITC) and rhodamine B were used to label the macrophage and red blood cell membranes, respectively. The fluorescence emission spectra of RBC-MP-microbowls demonstrated that rhodamine B and FITC were successfully labelled to the cell membrane, as shown in Fig. S4 (ESI†). Then, hybrid membrane-coated microbowls were imaged using a confocal laser scanning microscope (CLSM) to further verify the coating efficiency (Fig. 2D). Both green fluorescence (FITC representing the macrophage membrane) and red fluorescence (rhodamine B representing the red blood cell membrane) were observed from the microbowls, demonstrating the desired hybrid cell membrane coating.

3.2. Regulating effect on microbowls

The morphology and structure of the microbowls were related to the erosive action of dibutyl phthalate and styrene. Corresponding microbowls were prepared at different mass ratios of dibutyl phthalate and styrene (DS ratios), and the microstructures were characterized by SEM. Compared to PGMA microspheres (Fig. 3A), the diameter of the denting opening This article is licensed under a Creative Commons Attribution-NonCommercial 3.0 Unported Licence.

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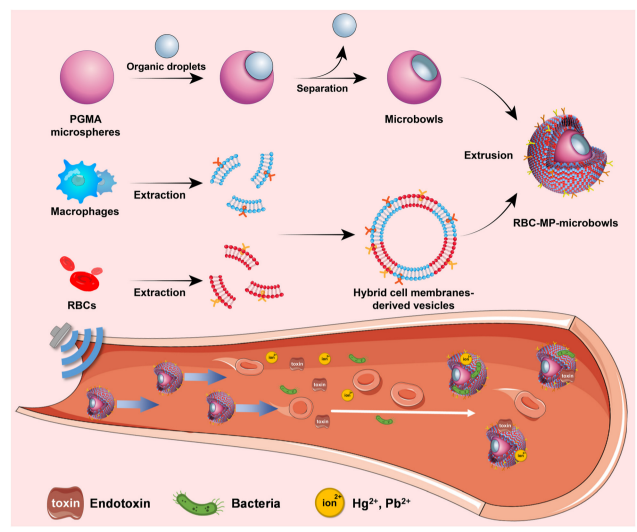


Fig. 1 Schematic illustration of the preparation of macrophage-red blood cell hybrid membrane-coated ultrasound-responsive microbowls and the process of eliminating pathogens, endotoxins, and heavy metal ions for blood purification.

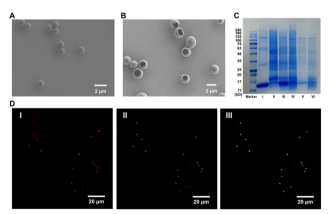


Fig. 2 Characterizations of RBC-MP-microbowls. (A) SEM image of PGMA seed microspheres. (B) SEM image of the microbowls. (C) Protein profiles of the RBC membranes (I), macrophage membrane (II), RBC-MP hybrid membranes (III), RBC-MP-microbowls (IV), RBC-microbowls (V) and MP-microbowls (VI) determined using an SDS-PAGE electrophoresis assay. (D) CLSM images of RBC-MP-microbowls.

ranged from 586.2 nm to 2.9 µm with the increase in styrene ratio (Fig. 3B-D). Moreover, excessive styrene gradually increased the diameter of the openings and eventually the bowl-like structure could not be maintained. In addition, the SEM images of different dibutyl phthalate contents with the same styrene contents were also acquired, and are shown in Fig. S5 (ESI†). Although the DS ratio was obviously changed from 5.0 mg/5.0 mg to 10.0 mg/5.0 mg, there were no significant changes in the microstructure, demonstrating that dibutyl phthalate has less impact on the morphology change towards microbowls. Bowl-like structures prepared at various eroding times were also investigated, and the corresponding SEM images are displayed in Fig. S6 (ESI†). After 3 h of reaction, only a small concave was observed (Fig. S6A, ESI†). With the continuous erosion by dibutyl phthalate and styrene, a typical bowl-like structure was formed after 12 h of reaction. However, the bowl-like structure became unstable and distorted when the reaction time reached 36 h (Fig. S6D, ESI†).

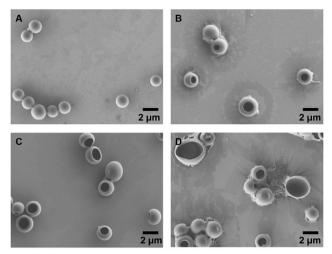


Fig. 3 SEM images of microbowls prepared with different DS ratios. (A) DS ratio = 0/0; (B) DS ratio = 5.0 mg/5.0 mg; (C) DS ratio = 5.0 mg/10.0 mg; and (D) DS ratio = 5.0 mg/15.0 mg (per 1 mg PGMA seed microspheres). Scale bar = $2 \mu m$.

3.3. Ultrasound responsibility and propulsion performance of **RBC-MP-microbowls**

The ultrasound-induced propulsion indexes of the RBC-MPmicrobowls, such as movement trajectories and velocities, were studied using an optical microscope and a video tracking software Video Spot Tracker (Version 8.01). The frequency of the RBC-MP-microbowls in response to ultrasound stimulation

was investigated, and it was found to be capable of directional motion at a frequency of 1.46 MHz. Previous studies have demonstrated that ultrasound could form standing waves and generate the primary acoustic radiation force perpendicular to the levitation plane, which suspended the microbowls. In addition, the second acoustic radiation force and streaminginduced drag force were generated in the levitation plane.⁴⁴ The shape and structure asymmetry caused the asymmetric distribution of two forces and acoustic radiation pressure gradients to realize orientation motion under non-invasive ultrasound fields.45

Then, the moving trajectories in a period of 5 s under different ultrasound amplitudes (5 V_{p-p}, 10 V_{p-p}, 15 V_{p-p}, and 20 V_{p-p}) were captured. Corresponding captured movements at 5 V_{p-p} and 10 V_{p-p} were recorded, and are shown in ESI,† Movies S1 and S2, respectively. In Fig. 4A, the motion trajectories are illustrated at different ultrasound amplitudes with obvious directional movements. The corresponding trajectories and direction of movement were recorded and normalized, as shown in Fig. 4B, by recording the coordinate points. Besides, the moving trajectories without ultrasound were captured, which showed relatively irregular movement attributed to the Brownian motion. The linearity of the trajectories indicated the stability of the directional movement of microbowls. The mean square displacements (MSD) were calculated by accumulating the squared linear distance between two frames, and are shown in Fig. 4C. As the ultrasound amplitude increased from 5 to 20 V_{p-p} , the MSD raised from 7.58 μm^2 to 186.8 μm^2 , indicating

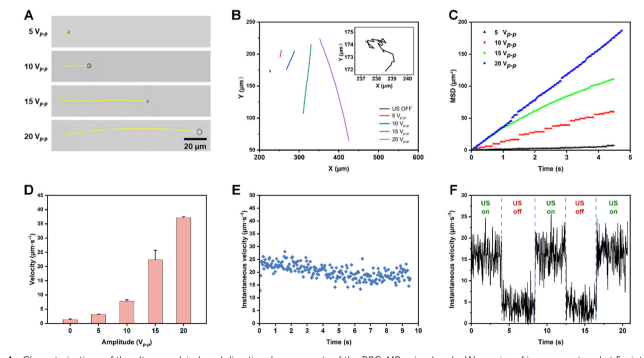


Fig. 4 Characterizations of the ultrasound-induced directional movements of the RBC-MP-microbowls: (A) a series of images captured at 5 s intervals showing the movement at different ultrasound amplitudes (5–20 V_{p-p}). Scale bar = 20 μ m. (B) Corresponding tracking trajectory of the RBC-MPmicrobowls during 5 s at different ultrasound amplitudes (0–20 V_{p-p}). (C) Average MSD versus time interval analysed from tracking trajectories. (D) Dependence of the average velocity of RBC-MP-microbowls in 5 s at different ultrasound amplitudes. (E) Instantaneous velocity of RBC-MPmicrobowls at 15 V_{p-p} analysed from the tracking trajectory. (F) Responsiveness of instantaneous velocity to ultrasound analysed via ultrasound ON/OFF.

a correlation between the moving velocity and the ultrasound amplitude. The mean velocities with different ultrasound amplitudes were measured to assess the effect on motion velocity (Fig. 4D). The mean velocity at 0, 5 V_{p-p}, 10 V_{p-p}, 15 V_{p-p} and 20 V_{p-p} was 1.43 $\mu m s^{-1}$, 3.18 $\mu m s^{-1}$, 7.84 $\mu m s^{-1}$ $22.39~\mu m~s^{-1}$ and $37.14~\mu m~s^{-1}$, which exponentially increased $(Y = 1.40947 \times 1.17777^X, \text{ Fig. S7, ESI}^{\dagger}).$

To study the velocity of the RBC-MP-microbowl in solutions with different viscosities, PBS and whole blood spiked with anticoagulants were used under ultrasound. As shown in Fig. S8 (ESI†), the velocities of RBC-MP-microbowls in PBS and whole blood were 37.14 $\mu m \ s^{-1}$ and 29.45 $\mu m \ s^{-1}$, respectively. In addition, red blood cells did not show any obvious ultrasound-responsive movement, which is in accordance with the existing results reported in the literature. 46 Overall, it has been demonstrated that the RBC-MP-microbowls could realize ultrasound propulsion in blood with reduced velocity compared to that in PBS.

To verify the persistence and stability of ultrasonic-responsive motion, the motion of the RBC-MP-microbowls was monitored at 15 V_{D-D} for 10 s, and the instantaneous velocities were measured by calculating the linear displacement between two frames. As shown in Fig. 4E, the instantaneous velocities were mainly between $20 \mu m s^{-1}$ and $25 \mu m s^{-1}$ in the first 4 s and then slightly decreased in the following 6 s, indicating that the ultrasound-propulsion movement had desired persistence and stability. In addition, the motion of RBC-MP-microbowls was further examined with frequent ultrasound switch (on and off), and instantaneous velocities were analysed to verify their transient responsiveness (Fig. 4F and Movie S3, ESI†). It showed that when the ultrasound switched from "on" to "off" or from "off" to "on", the instantaneous velocity changed instantaneously. Almost all the erythrocytes retained their biconcave disc shape after ultrasound stimulation, which is within the range of ultrasound frequency (<5 MHz) in clinical settings, which also verified the biosafety of sonication. 47,48 Besides, the movements at an ultrasound frequency of 1.46 MHz were recorded in the simultaneous presence of erythrocytes and RBC-MP-microbowls (Movie S4, ESI†). The RBC-MP-microbowls moved normally in the environment, while no obvious movements of erythrocytes were observed, demonstrating the selective ultrasound propulsion towards RBC-MP-microbowls.

3.4. Biocompatibilities and biofunctions of RBC-MPmicrobowls

Mouse fibroblast (L929) and B16 cell lines were selected to evaluate the biocompatibility of RBC-MP-microbowls by a CCK-8 assay. As shown in Fig. 5A, the cell viability was higher than 95% after co-culturing with 200 μg mL⁻¹ of RBC-MPmicrobowls for 24 h. In addition, we investigated the hemolytic property by incubation of RBC-MP-microbowls and RBCs. There was no apparent hemolysis phenomenon observed (Fig. 5B), demonstrating the feasibility of microbowls used during hemoperfusion.49

Endotoxin is one of the important pathogenic components causing sepsis. A recombinant factor c endpoint fluorescent

assay was performed to evaluate the endotoxin removal efficiency of RBC-MP-microbowls. As shown in Fig. 5C and Fig. S9 (ESI†), after incubation with ~ 2.5 EU per mL endotoxin for 15 min, the concentration of endotoxin significantly decreased to 0.53 EU per mL. The endotoxin removal efficiency was calculated with 79.6% calculated according to the calibration curve (Fig. S10, ESI†). We also investigated the removal ability with or without ultrasound at 0.5 and 1.0 EU per mL initial ion concentrations. As shown in Fig. 5D and E, the concentrations of endotoxin were significantly reduced with RBC-MPmicrobowl treatment. Furthermore, with RBC-MP-microbowls under ultrasound stimulation, the removal efficiency further increased from 80.55% to 92.80% at 1.0 EU per mL.

To verify the ability of scavenging heavy metal ions of the RBC-MP-microbowls, Pb^{2+} and Hg^{2+} (~10 mg L^{-1}) were selected as common blood contaminants. ^{31–33} In detail, 100 μL of RBC-MP-microbowls (242.75 μg mL⁻¹ of membrane protein) were added to 1 mL of heavy metal ion solutions with or without sinusoidal ultrasound at 1.46 MHz. As shown in Fig. 5F, the concentration of Pb²⁺ was reduced from 14.84 mg L^{-1} to 4.11 mg L^{-1} after incubation with RBC-MPmicrobowls, which further decreased to 2.34 mg L⁻¹ under ultrasonic stimulation of 1.46 MHz. Similarly, the concentration of Hg^{2+} was reduced from 13.01 mg L^{-1} to 3.68 mg L^{-1} , and further decreased to 2.33 mg L⁻¹ under ultrasonic stimulation of 1.46 MHz. The RBC-MP-microbowls reached removal efficiencies of 84.2% for Pb²⁺ and 82.1% for Hg²⁺ (Fig. 5G). Meanwhile, the adsorption capacity in removing Pb²⁺ and Hg²⁺ could reach 25.00 mg g⁻¹ and 21.36 mg g⁻¹, respectively. According to clinical reports, the reference values of blood lead concentration and serum mercury concentration were <50 µg L $^{-1}$ 39 and <15 µg L $^{-1}$,40 respectively. Therefore, we investigated the adsorption efficiency for different abovethreshold concentrations of Pb2+ and Hg2+ with RBC-MPmicrobowl + US treatment. As shown in Fig. 5H and I, the highest removal efficiency of RBC-MP-microbowls could reach 92.75% and 93.91%, respectively. This demonstrated that the RBC-MP-microbowls possessed great removal capacities for Pb²⁺ and Hg²⁺, which was attributed to the adsorption of the RBC membrane protein to heavy metal ions⁵⁰ and ultrasoundpropelling-boosted scavenging ability.⁵¹

Besides, the ability of scavenging bacteria was investigated using a Staphylococcus aureus (S. aureus) model. As shown in Fig. 5J, the removal efficiency for S. aureus was up to 96.7% in the RBC-MP-microbowl + US group, which was more effective than that of the PBS + US group and RBC-MP-microbowls, indicating the synergetic effects of the macrophage membrane and ultrasound-induced movements for bacterial scavenging. The function of adsorption to endotoxins and bacteria of RBC-MP-microbowls was attributed to the recognition and adsorption of macrophage cell membrane proteins (such as Toll-like receptors, NOD-like receptors, and cluster of differentiation 14⁵²) to endotoxins and bacteria. In addition, ultrasound promoted the movement of RBC-MP-microbowls, ultimately reaching great removal efficiencies for endotoxins and bacteria.

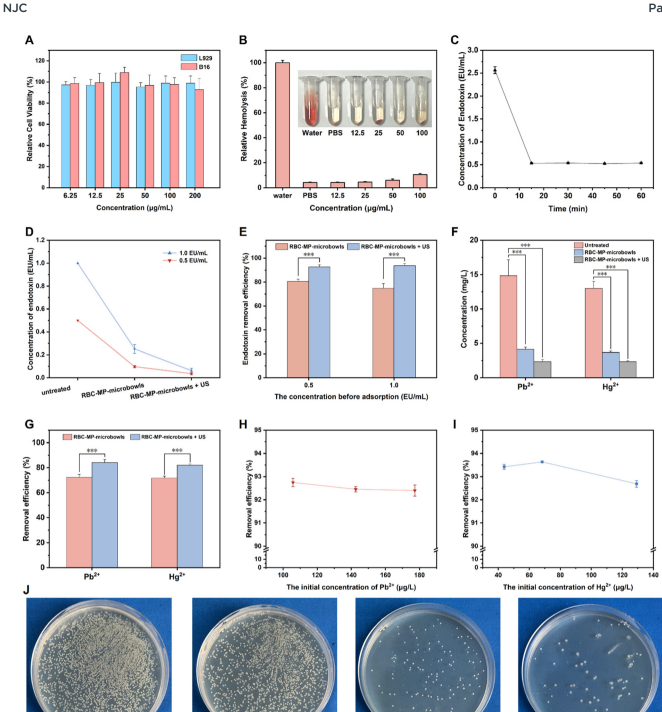


Fig. 5 Characterization of the bioactivity and ability to adsorb and remove the RBC-MP-microbowls: (A) cell viability assessment after incubation with different concentrations of RBC-MP-microbowls for 24 h. (B) Hemolysis activity of the RBC-MP-microbowls at different concentrations. (C) Concentration of endotoxins after incubation with the RBC-MP-microbowls for different times. (D) Concentrations of endotoxins before and after adsorption by the RBC-MP-microbowls with or without US. (E) Removal efficiency of endotoxins at different initial endotoxin concentrations. (F) Concentrations of heavy metal ions before and after adsorption. (G) Removal efficiency of the heavy metal ions. (H) Pb²⁺ removal efficiencies at different initial adsorption concentrations. (I) Hg^{2+} removal efficiencies at different initial adsorption concentrations. (J) Removal effect of S. aureus with the RBC-MP-microbowls under ultrasound stimulation. The statistical significance was analysed using Student's t-test. *: P < 0.05; **: *P* < 0.01; ***: *P* < 0.001.

PBS + US

PBS

RBC-MP-microbowls

RBC-MP-microbowls + US

4. Conclusions

In this study, we have developed a strategy to prepare ultrasound-responsive asymmetric polymeric microbowls coated with cell membranes to eliminate endotoxins, heavy metal ions and bacteria. The microbowls showed a unique asymmetric structure that enabled orientation motion under non-invasive ultrasound fields, which could achieve 37.14 μ m s⁻¹ at 20 V_{p-p} under fuel-free programable sonication. The utilization of the macrophage and RBC hybrid membranes on microbowls enhanced their biocompatibility and offered desired bioactivities. In detail, macrophage membranes facilitate the selective recognition and adhesion of pathogens and endotoxins. Experimental data revealed that 92.80% of endotoxin and over 90% of S. aureus in blood were eliminated. Additionally, the affinity between the RBC membrane and heavy metal ions contributes to the efficient removal of contaminants such as Pb²⁺ (92.75% removal efficiency) and Hg²⁺ (93.91% removal efficiency). Overall, our ultrasound-responsive asymmetric polymeric microbowls represented a promising advancement for blood purification during sepsis treatments.

Author contributions

Jianxing Jing: conceptualization, formal analysis, investigation, writing, review and editing. Mingchen Lv: editing. Wei Hu: editing. Runxin Teng: editing. Zhenghong Ge: editing. Yao Zhang: review and editing, supervision. Min Sun: review and editing, supervision. Zhen Fan: writing-original draft preparation, writing-review and editing, supervision.

Conflicts of interest

There are no conflicts of interest to declare.

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