Chemical Science

EDGE ARTICLE



Cite this: Chem. Sci., 2022, 13, 12799

All publication charges for this article have been paid for by the Royal Society of Chemistry Ultrasensitive detection of β-lactamase-associated drug-resistant bacteria using a novel mass-tagged probe-mediated cascaded signal amplification strategy[†]

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The emergence and spread of drug-resistant bacteria (DRB) is a global health threat. Early and accurate detection of DRB is a critical step in the treatment of DRB infection. However, traditional assays for DRB detection are time-consuming and have inferior analytical sensitivity and quantification capability. Herein, a mass-tagged probe (MP-CMSA)-mediated enzyme- and light-assisted cascaded signal amplification strategy was developed for the ultrasensitive detection of β -lactamase (BLA), an enzyme closely associated with most DRB. Each MP-CMSA probe contained multiple poly(amidoamine) (PAMAM) dendrimer molecules immobilized on a streptavidin agarose bead *via* a BLA-cleavable linker, and each dendrimer was modified with multiple mass tags *via* a photo-cleavable linker. In BLA detection, BLA could cleave the BLA-cleavable linker, leading to dendrimers shedding from the MP-CMSA probe to achieve enzyme-assisted signal amplification. Then, each dendrimer can further release mass tags under UV light to achieve light-assisted signal amplification. After this cascaded signal amplification, the released mass tags were ultimately quantified by mass spectrometry. Consequently, the sensitivity of BLA detection can be significantly enhanced by four orders of magnitude with a detection limit of 50.0 fM. Finally, this approach was applied to the blood samples from patients with DRB. This platform provides a potential strategy for the sensitive, rapid and quantitative detection of DRB infection.

Received 16th March 2022 Accepted 11th October 2022

DOI: 10.1039/d2sc01530g

rsc.li/chemical-science

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[†] Electronic supplementary information (ESI) available: ESI table (S1), ESI figures (S1–S13) and ESI experimental information (S1–S14) including materials and reagents, instruments, MS analysis, synthesis routes for chemicals, characterization of products, optimization of experimental conditions, bacterial strains and analysis, preparation of stock solutions, calibration standards and quality controls (QCs), BLA inhibition assay, validation results of the assay, representative AST results, and LC-MS/MS chromatograms of calibration standards and the patient samples. See DOI: https://doi.org/10.1039/d2sc01530g [‡] J. Z. and Y. B. contributed equally to this work.

1. Introduction

The increase in drug-resistant bacteria (DRB) infection has become one of the greatest dangers to public health.¹ In addition, it is estimated that the mortality due to DRB infection will be significantly higher than cancer mortality in the near future.² More importantly, DRB infection is particularly concerning during the current coronavirus pandemic because it can exacerbate coronavirus symptoms, especially in geriatric patients, and thus coronavirus-infected patients must be extremely careful to avoid secondary infection of DRB.3 On the other hand, it is always of the utmost importance to diagnose the onset of DRB as early as possible.⁴ There is much evidence indicating that early and appropriate combating of DRB can significantly improve the clinical outcomes.^{5,6} However, false diagnosis of DRB infection at the initial stage can easily miss the optimal therapeutic window and promote the transmission of DRB.7 Furthermore, the number of infectious bacteria in individuals often determines the dosing regimens.8 Suboptimal dosing regimens could also cause poor clinical outcomes and further promote drug resistance.9 Therefore, early and quantitative detection of DRB is important for efficient diagnosis and treatment of DRB infection.



Currently, antibacterial susceptibility testing (AST), such as disk diffusion, broth dilution and E-test, is the gold standard for DRB detection.¹⁰ Although AST provides reliable results, these culture-based assays are time-consuming (>two days) and laborintensive due to very few bacterial strains being available at an early stage of infection.¹¹ In recent years, alternative techniques for early DRB detection have been developed; however, significant efforts are still required to make them quantitative.12 In the case of β-lactamase (BLA), an enzyme closely associated with most DRB, its rapid detection is critical for circumventing resistance to β -lactam antibiotics.¹³ Several new methods, such as the PCR and fluorescence in situ hybridization (FISH), have been attempted for its detection in recent years.^{14,15} However, these methods cannot cover all BLA genes and mutations given that nearly 2800 types of BLA have been identified to date.13 To achieve better sensitivity and quantitative capability, alternative methods based on smart techniques are always urged.

Recently, mass spectrometry (MS) has become an indispensable tool for target molecule detection, primarily due to its great quantitative capability.16 MS-based methods have been applied to bacterial detection in laboratory medicine.17 However, the population of DRB in patients at the early stage of infection may be much lower than the detection limit of MS (>10⁵ colony-forming units (CFU) per mL).¹⁸ Thus, a lengthy culture-enrichment step is often required prior to MS detection.19 Fortunately, among various MS-based approaches, the one relying on mass-tagged probes and liquid chromatographytandem mass spectrometry (LC-MS/MS) can convert and amplify the signal of the target molecule into that of mass tags such as peptides, which has significantly expanded the application of MS.²⁰ Moreover, mass-tagged probes can convert the signal of each target molecule into that of multiple mass tag molecules using signal amplification strategies.²¹ Notably, multiple signal amplification strategies may be required for the detection of ultralow abundant target molecules such as DRB.

In the current work, a novel mass-tagged probe (MP-CMSA) was developed for BLA detection. The MP-CMSA probe can convert the amount of BLA to the mass tag signal *via* enzymeand light-assisted cascaded signal amplification. The released mass tag was ultimately quantified by MS. After further optimization and validation, the new assay was applied for BLA detection in blood samples of patients who were diagnosed with bacterial infection.

2. Experimental section

2.1 Synthesis of biotin-poly(ethylene glycol)-7aminocephalosporanic acid (BPEG-7ACA)

First, 200 mg of biotin-poly(ethylene glycol)-succinimidylcarboxymethyl-ester (BPEG-NHS) and 34.5 mg of hydromethyl-7-aminocephalosporanic acid (7ACA) were dissolved in 40 mL of deionized water at a final concentration of 2.50 mM BPEG-NHS and 3.64 mM 7ACA, and then 4 μ L of triethylamine (TEA) was added to adjust the pH of the mixture to 8.0. The reaction mixture was stirred at 600 rpm at 30 °C for 24 h. Thereafter, the mixture was placed into a dialysis bag with a molecular weight cutoff (MWCO) of 1.0 kDa and dialyzed against deionized water for 72 h. BPEG-7ACA was obtained through lyophilization.

2.2 Synthesis of biotin-poly(ethylene glycol)-7aminocephalosporanic acid-poly(ethylene glycol)-azido (BPEG-7ACA-N3)

For BPEG-7ACA-N3 synthesis, 110 mg of BPEG-7ACA and 150 mg of azido-poly(ethylene glycol)-succinimidylcarboxymethyl-ester (N3-PEG-NHS) were dissolved in 40 mL of dimethyl sulfoxide (DMSO) at a final concentration of 1.25 mM BPEG-7ACA and 1.88 mM N3-PEG-NHS, and then 4 μ L of TEA was added. After stirring at 600 rpm at 30 °C for 24 h, the mixture was dialyzed with a MWCO of 3.5 kDa against deionized water for 72 h. Finally, BPEG-7ACA-N3 was obtained through lyophilization.

2.3 Synthesis of poly(amidoamine)-dibenzocyclooctyne (PAMAM-DBCO)

First, 100 mg of poly(amidoamine) (PAMAM) and 268 mg of dibenzocyclooctyne-*N*-hydroxysuccinimidyl ester (DBCO-NHS) were dissolved in 60 mL of DMSO at a final concentration of 0.0578 mM PAMAM and 11.1 mM DBCO-NHS and stirred at 600 rpm at 30 °C for 18 h. Subsequently, the mixture was placed into a dialysis bag with a MWCO of 10.0 kDa and dialyzed against DMSO for two days and then against deionized water for another two days. Finally, PAMAM-DBCO was obtained through lyophilization.

2.4 Synthesis of a dual-functional molecule (DM)

BPEG-7ACA-N3 and azido-modified photo-cleavable peptide AVLGDPFR-PL-GFK-N3 (MP-PL-N3) were conjugated to PAMAM-DBCO to obtain a DM. For DM preparation, 10 mL of 10.0 mM MP-PL-N3 in PBS (equiv. to 10.0 mM mass tag) and 4 mL of 10.0 mM BPEG-7ACA-N3 in PBS (equiv. to 10.0 mM 7ACA) were mixed with 10 mL of 0.100 mM PAMAM-DBCO and stirred at 600 rpm at 30 °C for 12 h. Subsequently, the mixture was purified by dialysis with a MWCO of 10.0 kDa in deionized water. The DM product was obtained by lyophilization. Notably, the number of conjugated mass tags can be estimated from the released mass tag after the DM was exposed to UV light. The number of mass tags divided by the amount of DM, which was equivalent to that of the reactant PAMAM-DBCO (please see the ESI†).

2.5 Preparation of the MP-CMSA probe

Streptavidin agarose beads were immobilized with the DM to obtain the MP-CMSA probe. In brief, streptavidin agarose beads were washed with PBS three times. Then, 1 mL of streptavidin agarose beads was mixed with 1 mL of 12.0 μ M DM and incubated at 37 °C for 2 h. Subsequently, the beads were washed with PBS three times and stored at 4 °C in PBS. The supernatant was exposed to UV light, and the mass tag released from the unimmobilized DM was detected by MS. The amount of the immobilized DM can be estimated from the released mass tag

in the supernatant. The number of immobilized DMs per bead was calculated as the amount of immobilized DMs divided by the number of beads in the reaction (please see the ESI[†]).

2.6 Evaluation of enzymatic-cleavage and photo-cleavage efficiencies of the MP-CMSA probe

To evaluate the photo-cleavage efficiency, $100 \ \mu L$ of $100 \ nM$ MP-CMSA probe was irradiated under UV light at 365 nm at various times. Then, the sample was centrifuged at 1000 rpm for 5 min, and the pellets were washed with 100 μL of PBS three times. The collected supernatants were combined and subjected to mass spectrometry.

To evaluate the enzymatic-cleavage efficiency, 100 μ L of 10.0 μ M MP-CMSA probe was incubated with 100 μ L of 100 pM BLA in PBS for various times or 100 μ L of BLA at different concentrations for 2 h. Then, the sample was centrifuged at 1000 rpm for 5 min, and the pellets were washed with 100 μ L of PBS three times. The collected supernatants were combined and exposed to UV light for 40 min and then subjected to LC-MS/MS analysis.

2.7 BLA detection in the blood

One milliliter of blood was centrifuged at 800 rpm for 3 min to obtain the supernatant. The collected supernatant was further centrifuged at 12 000 rpm at 4 °C for 8 min. The obtained precipitate was resuspended in 100 μ L of PBS and sonicated for 20 s using an XL-2000 Series probe sonicator set at 5 W and 22 kHz. Subsequently, 100 μ L of lysate was mixed with 100 μ L of 10.0 μ M MP-CMSA probe. The mixture was incubated in the dark at 37 °C for 2 h and then centrifuged at 1000 rpm for 5 min.

The pellets were washed with 100 μ L of PBS three times. The collected supernatants were combined and exposed to UV light for 40 min and then desalted using a MicroSpin C18 column (The Nest Group, Inc., MA, USA) after the addition of 10 μ L of internal standard peptide solution. The obtained mass tag sample was measured by MS. This process was operator-blinded and the LC-MS/MS analysis was conducted by two trained operators.

3. Results and discussion

3.1 Probe design and working principle

In this study, the MP-CMSA probe played both detection and amplification roles. To achieve the detection function, the probe should basically have a recognition moiety for targeting and a probe tag for detection. Correspondingly, (1) 7ACA, the parent core of β -lactam antibiotics that can be easily recognized and cleaved by BLA as a substrate,22 was employed as the recognition part of the probe. (2) A peptide with the sequence AVLGDPFR was selected as the mass tag. For the amplification feature of the probe, there are a variety of strategies for signal amplification. Among these strategies, highly branched molecules such as dendrimers with easy modification, good chemical and physical stabilities, and simple preparation have attracted widespread attention.23 There is evidence indicating that tag molecules can be conjugated on the surface of dendrimers to amplify the signal exponentially, depending on the series of generations of the dendrimer.24 Thus, a fifthgeneration PAMAM dendrimer with 128 terminal amino groups was employed in the MP-CMSA probe. As illustrated in



Fig. 1 Schematic illustration of the MP-CMSA probe preparation and its mediated enzyme- and light-assisted cascaded signal amplification for the ultrasensitive detection of BLA. (A) Preparation of the MP-CMSA probe. BPEG-7ACA-N3 and MP-PL-N3 were conjugated to PDMAM-DBCO *via* a click reaction to obtain the DM. Subsequently, the DM was immobilized on streptavidin agarose beads to form the MP-CMSA probe. (B) Working principle of the MP-CMSA probe. BLA in the blood could break the BLA-cleavable linker 7ACA to shed the DM off the beads, thus achieving enzyme-assisted signal amplification. Then, UV light could break the photo-cleavable linker PL, and the DM released mass tags to achieve light-assisted signal amplification. The released mass tags were detected by MS, and the MS signal of the mass tag represented the amount of BLA.

Fig. 1, BPEG-7ACA-N3 with the BLA-cleavable linker 7ACA and MP-PL-N3 with the photo-cleavable linker PL were both conjugated to the DBCO-modified PAMAM dendrimer (PAMAM-DBCO) to obtain a DM. Multiple DMs were then anchored on a streptavidin agarose bead to acquire the MP-CMSA probe. In BLA detection, BLA molecules could cleave 7ACA, leading to DM shedding from the MP-CMSA probe. If there was sufficient time, each BLA molecule could cleave multiple 7ACA linkers sequentially, resulting in more DM shedding. Then, each dendrimer can further release mass tags under UV light to achieve signal amplification. Therefore, the cascaded signal amplification actually relied on the bearing capability of the bead and the branched structure of the dendrimer.

3.2 Probe preparation and characterization

As described earlier, the MP-CMSA probe was produced by immobilizing the DM on streptavidin agarose beads. The DM preparation mainly contained four steps (Fig. S1†): (1) BPEG-NHS was conjugated to 7ACA through a condensation reaction to produce BPEG-7ACA, (2) BPEG-7ACA reacted with N3-PEG-NHS to form BPEG-7ACA-N3 through another condensation reaction, (3) DBCO-NHS was conjugated to PAMAM to form PAMAM-DBCO, and (4) both BPEG-7ACA-N3 and MP-PL-N3 reacted with PAMAM-DBCO through a copper-free click reaction to generate the DM.

¹H nuclear magnetic resonance spectroscopy (¹H NMR) and matrix-assisted laser desorption/ionization-time of flight mass spectrometry (MALDI-TOF MS) were used to verify the structure and molecular weight (M_w) of the reaction products, including BPEG-7ACA, BPEG-7ACA-N3, PAMAM-DBCO and DM. As shown in Fig. 2A, the characteristic signals at 4.17, 5.09 and 5.62 ppm in the ¹H NMR spectrum of BPEG-7ACA can be attributed to the 3'-CH, 6-CH and 7-CH protons of 7ACA, confirming the formation of BPEG-7ACA. Then, the 3'-CH proton peak shifted from 4.1 ppm to 4.76 ppm in the spectrum of BPEG-7ACA-N3, demonstrating the occurrence of an esterification reaction between the hydroxyl group of 7ACA and the carboxyl group of N3-PEG-NHS. According to the spectrum of PAMAM-DBCO, the multiple peaks between 7 and 8 ppm were derived from the hydrogen protons on the benzene ring of DBCO. Finally, the majority of characteristic peaks of PAMAM, DBCO, MP-PL-N3 and BPEG-7ACA-N3 appeared in the spectrum of the DM, suggesting that the DM was successfully synthesized. The changes in the size distribution and zeta potential of PAMAM, PAMAM-DBCO and DM also confirmed this point (Fig. S2[†]). Moreover, the $M_{\rm w}$ values of BPEG-NHS, BPEG-7ACA-N3, BPEG-7ACA, PAMAM-DBCO and DM in the MALDI-TOF MS spectra were consistent with their theoretical values (Fig. 2B). Finally, the conjugation number of mass tags for each DM was estimated to be approximately 75 (please see the ESI[†]).

Afterward, the DM was immobilized on the surface of streptavidin agarose beads through biotin-streptavidin interactions to produce the MP-CMSA probe. The immobilization efficiency was optimized (Fig. S3†) and estimated to be 7.33×10^3 DM per bead (please see the ESI†). The obtained MP-CMSA probe exhibited excellent stability under both storage (PBS, pH 7.4 at 4 °C) and detection conditions (PBS, pH 7.4 at 37 °C). The probe degradation percentage was only (1.52 \pm 0.08)% after 7 days under the storage conditions and (1.02 \pm 0.06)% after 48 h under the detection conditions (Fig. S4†).

3.3 Optimization of signal amplification conditions

The amplification conditions, including photo-assisted and enzymatic-assisted reaction conditions, were carefully optimized and characterized in this study.

3.3.1 Photo-cleavage conditions. The release of mass tags from the MP-CMSA probe is one of the key points to achieving signal conversion and amplification in the assay. Currently, several stimuli have been used to trigger tag release from masstagged probes, including enzymes,²⁵ light²⁶ and electronics.²⁷ Among them, photo-cleavable reactions provide spatial and temporal control of chemical bond cleavage with high efficiency.28 Specifically, these reactions can finish within 1 h, which may drastically improve the assay throughput.²⁹ In this study, an o-nitrobenzyl derivative PL was employed as the linker, which is UV light sensitive.30 The photo-cleavable mechanism of the PL is illustrated in Fig. S5.† Consistent with previous studies, the mass tag was the dominant product if it was conjugated at the R1 position of the PL and could be completely released after UV exposure.31 Furthermore, a PL with two alkoxy groups and one nitro group on the benzene ring can rapidly degrade at wavelengths longer than 350 nm, which is less harmful to biological systems.²⁹ Our results demonstrated that the optimized photolysis time was 40 min (Fig. S6[†]), at which time the mass tag release from the MP-CMSA probe was almost complete (Fig. 3A).

3.3.2 Enzymatic-cleavage conditions. The mechanism of BLA hydrolysis has been reported previously.^{32,33} In this work, the reaction was supposed to begin with the binding of BLA to 7ACA in the MP-CMSA probe. Then, BLA opened the β -lactam ring of 7ACA and produced an acyl-enzyme intermediate. This process led to chemical rearrangement and the expulsion of the DM from the beads of the probe. Finally, water molecules attacked the acyl-enzyme intermediate, leading to BLA refreshment for the next enzymatic-cleavage cycle. As a proof of concept, the $M_{\rm w}$ of BPEG-7ACA-N3 decreased after treatment with a commercially available recombinant BLA (TEM-1) (Fig. S7[†]). The fluorescence intensity of FITC-labeled DMimmobilizing beads decreased with increasing incubation time due to the loss of the fluorescein group (Fig. 3B). Then, the enzymatic-cleavage conditions were further optimized using the MP-CMSA probe. As presented in Fig. 3C, significant signal amplification could be observed and 2 h of incubation was selected in the subsequent experiments.

3.4 Ultrasensitive and specific detection of BLA-associated DRB

The ultrasensitive detection of BLA-associated DRB in this study was achieved by cascaded signal amplification. The masstagged probe was employed to convert and amplify the information of BLA to the mass tag signal. During the mass spectrometric analysis, the MRM transitions with the best signal to



Fig. 2 Preparation and characterization of the DM. (A) ¹H NMR spectra of BPEG-NHS, BPEG-7ACA, and BPEG-7ACA-N3 in CDCl₃ and PAMAM-DBCO and DM in DMSO- d_6 . (B) MALDI-TOF MS spectra of BPEG-NHS, BPEG-7ACA, BPEG-7ACA-N3, PAMAM-DBCO and DM in methanol. The theoretical molecular weight (M_w) of each molecule is shown in the figure. The ¹H NMR results of BPEG-7ACA and BPEG-7ACA-N3 were expanded in the region of 4–6 ppm for clarity.



Fig. 3 (A) The percentage of the mass tag released from the MP-CMSA probe after different UV exposure times or in the dark. As shown, there was no significant mass tag release in the dark. (B) Evaluation of enzyme-assisted signal amplification of MP-CMSA. Fluorescence images of 100 μ L of 100 nM FITC-labeled MP-immobilizing beads incubated with 100 μ L of 50.0 pM BLA at different times. Correspondingly, the MS signal of 100 μ L of 10.0 μ M MP-CMSA probe incubated with 100 μ L of 50 pM BLA at various times followed by UV light exposure for 40 min is also shown. The error bars are indicative of the standard deviation of three measurement replicates of one mass tag sample.

noise ratio (S/N) of the mass tag AVLGDPFR (*i.e.*, m/z 437.9 \rightarrow m/z 704.4, m/z 437.9 $\rightarrow m/z$ 419.3 and m/z 437.9 $\rightarrow m/z$ 171.2) were employed (Fig. S8 and S9[†]). The stable isotope-labeled peptide AV*LGDPFR ([D8]Val) served as an internal standard (Fig. S9[†]). The relative peak area ratio of MRM transitions of the mass tag and the internal standard was plotted against the BLA concentration to establish a calibration curve and used for BLA quantification (Fig. 4A). The linear relationship between the MS signal of the MP-CMSA probe and the concentration of BLA ranged from 50.0 fM to 400 pM ($R^2 = 0.9967$). The lower limit of quantification (LLOQ) of BLA was determined to be 50.0 fM (S/N > 10, Fig. S10[†]), which was significantly lower than that of the standard chromogenic BLA probe (4.00 nM).34 This LLOQ was roughly equivalent to the concentration of typical BLAassociated DRB strains as low as 10³ CFU mL⁻¹.⁷ In addition, no significant signal was observed for the matrix blank (i.e., 5% BSA) and thus the background was low. Notably, the sample with a positive signal means that its mean signal of three measurement replicates was greater than LLOQ with a *p*-value less than 0.05 by the *t*-test.³⁵ In addition, we also compared the MS signals of the synthetic mass tag and the released mass tag in our probe-mediated approach with BLA at an equivalent concentration. The result indicated that the sensitivity was significantly enhanced by four orders of magnitude after signal amplification. Moreover, the precision and accuracy of QCs were $\leq \pm 15\%$ (LLOQ $\leq \pm 20\%$), (Table S1[†]). The precision of the approach was also confirmed by spiking the mass tag instead of the MP-CMSA probe before and after signal amplification. As a result, there is no significant difference in the MS signal variations between the sample groups.

Furthermore, the specificity of the approach was also validated. First, the signal of the mass tag could not be detected in the absence of BLA (Fig. 4B). Then, the signal of the mass tag significantly decreased with pretreatment of the BLA inhibitor potassium clavulanate.³⁶ The selectivity of the MP-CMSA probe against BLA was also confirmed using several common enzymes in bacteria including esterase (ESTA), β -glucosidase (β -Glu), phosphodiesterase (PDE), and nicotinamide adenine dinucleotide phosphate (NADPH) (Fig. 4C).³⁷⁻⁴⁰ Additionally, no significant signal was observed after the MP-CMSA probe was incubated with non-BLA-expressing *E. coli* ATCC 25922, that is the Clinical and Laboratory Standards Institute (CLSI) recommended negative quality control strain.⁴¹ Comparatively, a remarkable MS signal can be detected in three clinically prevalent BLA-expressing bacterial strains including *K. pneumonia*/KPC-3 ATCC BAA-2814, *K. pneumonia*/SHV-18 ATCC 700603 and *E. coli*/TEM-1 ATCC 35218 (Fig. 4D).⁴²⁻⁴⁴ These results confirmed the specificity of the MP-CMSA probemediated approach.

3.5 Detection of BLA-associated DRB in the blood

We further investigated the potential utility of this system to reveal BLA-associated DRB directly from blood samples (n = 35) of patients (age: ≥ 65 years (n = 13), <65 years (n = 13)) who were diagnosed with bacterial infections. The samples were originally defined by hospital pathologists using gold-standard AST (disc diffusion test, K-B method) following the CLSI.41 On the other hand, BLA detection using the mass-tagged probe was performed in the laboratory. This process was operator-blinded. Moreover, the blood samples were collected and analyzed by two trained operators. The analysis time of each sample was around 3 h. As shown in Fig. 5A, positive MS signals were observed in 26 samples using the MP-CMSA probe-mediated approach. The other 9 samples without positive MS signals were also assessed as BLA-negative using AST. Passing-Bablok regression analysis showed a good agreement between the results from the two operators (y = 0.992x - 0.034, p = 0.53) (Fig. S12[†]).

To further evaluate the approach performance, a classification strategy using a three-zone partition was employed for patient grouping in 26 samples.⁴⁵ Specifically, an optimal assay threshold was defined as 21.0 relying on Youden's index.⁴⁶ Then, an intermediate grey zone with an area ratio range of 16.8–25.2 was established using the 20% tolerance range of the cutoff value and defined as "equivocal". The zone with an area ratio of > 25.2 was defined as "positive" and that with an area ratio of < 16.8 was defined as "negative". The BLA-positive



Fig. 4 Evaluation of sensitivity and specificity of the mass-tagged probe-mediated approach. (A) Representative calibration curve of the mass tag signal from the MP-CMSA probe and BLA concentration ranging from 50.0 fM to 400 pM. (B) The signal of the mass tag after 100 μL of 10.0 μM MP-CMSA probe was incubated with 100 μL of 10.0 pM BLA, 100 μL of 10.0 pM BLA + 50.0 pM potassium clavulanate, or 100 μL of 5% bovine serum albumin (5% BSA) as the blank for 2 h followed by UV light exposure for 40 min. The *t*-Test was performed between BLA and other groups, ****p* < 0.001. (C) The MS signal of the mass tag after 100 μL of 10.0 μM MP-CMSA probe was incubated with 100 μL of 10.0 pM BLA, 100 μL of 10.0 pM esterase (ESTA), 100 μL of 10.0 pM β-glucosidase (β-Glu), 100 μL of 10.0 pM phosphodiesterase (PDE), or 100 μL of 10.0 pM nicotinamide adenine dinucleotide phosphate (NADPH) for 2 h followed by UV light exposure for 40 min. The *t*-Test was performed between BLA and other groups, ****p* < 0.001. (D) The MS signal of the mass tag after 100 μL of 10.0 μM of the MP-CMSA probe was incubated with 100 μL of non-BLA-expressing *E. coli* ATCC 25922, and BLA-expressing bacterial strains including *E. coli*/TEM-1 ATCC 35218, *K. pneumoniae*/KPC-3 ATCC BAA-2814 and *K. pneumoniae*/SHV-18 ATCC 700603 at 10⁶ CFU mL⁻¹ followed by UV light exposure for 40 min. The *t*-Test was performed between *E. coli* ATCC 25922 and other groups, ****p* < 0.001. The error bars are indicative of the standard deviation of three measurement replicates of one mass tag sample.



 Image: MP-CMSA
 Total

 Positive
 Equivocal
 Negative

 Positive
 Equivocal
 Negative

 Positive
 14
 1
 3

 Positive
 14
 1
 3

 Positive
 0
 1
 7

 Positive
 14
 2
 10

Fig. 5 Clinical evaluation of the MP-CMSA probe-mediated cascaded signal amplification strategy for BLA-associated DRB detection. (A) The positive MS signals of blood samples using the data from one operator. Blood samples were classified as equivocal (intermediate grey zone), positive (upper region), and negative (lower region). The blue dotted line corresponds to the Youden's index. (B) Sensitivity (83.3%) and specificity (100%) of the MP-CMSA probe-mediated approach in comparison with AST.

patients should be resistant to the β-lactam antibiotic treatment. Using this three-zone classification, the sensitivity and specificity of the MP-CMSA assay for the positive and negative samples were 83.3% and 100%, compared to the AST results (Fig. 5B). Notably, there were two "equivocal" samples revealed by the MP-CMSA assay, among which one was positive and the other was negative in AST. This inconsistency deserves further analysis in the future. Moreover, due to the limitation of the sample size, the cutoff value and the three-zone classification determined here needed further investigation with a large sample size. Moreover, more BLA-associated DRB infections were observed in patients aged over 65 (61.1%, data not shown), confirming that geriatric patients are more easily colonized or infected with BLA-associated DRB. This clinical phenomenon may be associated with nutritional deprivation or immunological impairment in those patients.47

4. Conclusions

In summary, we have developed a mass-tagged probe, MP-CMSA, with enzyme- and light-assisted cascaded signal amplification to convert and amplify the level of BLA into the signal of the mass tag. In this probe, the BLA substrate was used as the recognition part, and the peptide was used as the mass tag. The dendrimer structure and photo- and enzyme-cleavable characteristics of the MP-CMSA probe were employed for signal amplification. After cascaded amplification, the detection sensitivity of BLA can be effectively enhanced by four orders of magnitude, and as low as 50.0 fM BLA in blood samples can be detected. The results also demonstrated that geriatric patients were easily colonized or infected with BLA-associated DRB. Furthermore, this assay enabled the measurement of BLA directly from patient blood samples within a few hours without the need for bacteria culturing and thus could advance the diagnostic report by two to three days. Following the same strategy, other DRB or other types of pathogens could also be detected only by replacing the BLA recognition moiety with carbapenemase, extended-spectrum BLA-specific substrates, or other enzyme-specific substrates. Overall, the mass-tagged probe-mediated cascaded signal amplification strategy provides a potential approach for facilitating the sensitive and rapid diagnosis of DRB and timely selection of suitable therapeutic regimens.

Ethical approval

This study was approved by the Institutional Review Board of Nanjing Medical University, Nanjing, China.

Data availability

All the data supporting this article have been included in the main text and the ESI material.[†]

Author contributions

Jianhua Zhu: conceptualization, methodology, investigation, writing-original draft. Yunfei Bai: methodology, investigation,

writing-original draft. Xiuyu Chen: methodology, validation, formal analysis. Linlin Hu: resources. Wenjun Zhang: methodology, validation, formal analysis. Chunyan Liu: methodology, validation, formal analysis. Hua Shao: resources, visualization, writing-review & editing. Jianguo Sun: data curation, writing-review & editing. Yun Chen: conceptualization, project administration, funding acquisition, writing-review & editing.

Conflicts of interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this article.

Acknowledgements

The National Natural Science Foundation of China (22174068 and 21722504), the Natural Science Foundation of Jiangsu Province (BK20221303), the Jiangsu Provincial Key Research and Development Program (BE2022796), the SEU-NJMU-CPU Cooperation Project (2242019K3DNZ2), the Open Foundation of State Key Laboratory of Reproductive Medicine (SKLRM-2022BP1), and the Science and Technology Development Fundation of NJMU (NMUB2019014, NJMUQY2022003) awarded to Dr Chen, the Leading Technology Foundation Research Project of Jiangsu Province (BK20192005) awarded to Dr Sun, and the Changzhou Siyao Hospital Pharmaceutical Foundation of Nanjing Pharmaceutical Association (2019YX012) awarded to Dr Shao are gratefully acknowledged.

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