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Mussel-Inspired and Bioclickable Peptide Engineered Surface to Combat Thrombosis and Infection

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Abstract
Thrombosis and infections are the two major complications associated with extracorporeal circuits and indwelling medical devices, leading to significant mortality in clinic. To address this issue, here, we report a biomimetic surface engineering strategy by the integration of mussel-inspired adhesive peptide, with bio-orthogonal click chemistry, to tailor the surface functionalities of tubing and catheters. Inspired by mussel adhesive foot protein, a bio-clickable peptide mimic (DOPA)₄-azide-based structure is designed and grafted on an aminated tubing robustly based on catechol-amine chemistry. Then, the dibenzylcyclooctyne (DBCO) modified nitric oxide generating species of 1,4,7,10-tetraazacyclododecane-1,4,7,10-tetraacetic acid (DOTA) chelated copper ions and the DBCO-modified antimicrobial peptide (DBCO-AMP) are clicked onto the grafted surfaces via bio-orthogonal reaction. The combination of the robustly grafted AMP and Cu-DOTA endows the modified tubing with durable antimicrobial properties and ability in long-term catalytically generating NO from endogenous s-nitrosothiols to resist adhesion/activation of platelets, thus preventing the formation of thrombosis. Overall, this biomimetic surface engineering technology provides a promising solution for multicomponent surface functionalization and the surface bioengineering of biomedical devices with enhanced clinical performance.

MAIN TEXT

Introduction
The commonly used indwelling catheters and external circuits in medical scenarios (i.e., ventricular assist devices, pacemakers, artificial hearts, hemodialysis and cardioverter defibrillators), have been considered as the “lifeline” of patients(1, 2). However, thrombosis and infections of extracorporeal circuits and indwelling medical devices often cause device failure(3, 4) along with serious complications, including for example catheter-related thrombosis (5), catheter-related bloodstream infection (6), and deep phlebitis(7). Specifically, the formation of bacterial biofilm and thrombus on the surfaces of such types of blood-contacting devices are the major difficulty in the long-term treatment.

In traditional clinical practices, co-administration of antibiotics and anticoagulant drugs has been widely applied to prevent thrombosis and infection. However, this strategy can lead to various clinical complications, e.g., antibiotic resistance(8), bleeding(9), thrombocytopenia(10) and allergic reaction(11). Accordingly, endowing the implanted materials or devices with multifunctionalities to combat thrombosis and infection has attracted increasing attentions in recent years(12, 13). Among these strategies, grafting bioactive molecules, such as anticoagulant molecules(14), antibiotics(15), and peptide(16), which provides localized bioactivity on the surface of device, has been considered as one of the most effective methods for suppressing thrombosis and infections. However, the addressing of the two complications simultaneously is
still a formidable challenge as yet. Strategies that “learn from Nature” (17) may be a promising way to solve such problems.

In circulation, the natural blood vessels show remarkable anticoagulant and antibacterial properties, which is associated with the microenvironment of the blood vessels and their biological activities (18). Antimicrobial peptide (i.e., AMP), being widely distributed in plasma, directly kills bacteria when wound occurs, which is an crucial part of the immune system (19). Compared with antibiotics, AMP have a wide range of antibacterial mechanisms and do not cause bacterial resistance (20). In addition, endothelium exhibits a crucial effect in maintaining vascular homeostasis though releasing a kind of factors (e.g., Nitric oxide (NO)) (21-23). Endothelial cells continuously and stably release NO into blood microenvironment, which has a notable suppressive effect on platelet activation adhesion and thrombosis (24). Specially, the 1,4,7,10-tetraazacyclododecane-1,4,7,10-tetraacetic acid chelated copper ion (i.e., Cu-DOTA) is widely used as a stable and effective biologically active catalyst to decompose endogenous S-nitrosothiols (RSNOs) into NO (25, 26). Therefore, blood contact device with NO-generating function and surface engineered antimicrobial peptides may provide a highly simulated vascular microenvironment to prevent thrombosis and bacterial infection. However, the inevitably consumed active groups (e.g., -COOH, -NH₂, and -SH) of such biomolecules after chemical immobilization would result in a progressive loss of bioactivity as reported (27). Besides, the current chemical modification methods suffer from tedious reaction processes and complex surface treatment technologies, compromising controllability, maneuverability, and reproducibility of surface bioactivity. Considering these issues, click chemistry, a concept founded in 2001 (28), is considered as an effective tool to address these issues to some extent (29). In addition, bio-orthogonal click chemistry, a new click chemistry method (30), e.g., the dibenzylcyclooctyne-azide (N₃-DBCO) cycloaddition chemistry, demonstrates advantages in rapidity, thoroughness and specificity (31, 32). The high fidelity of these reactions in the face of a wide range of functional groups allows them to better maintain the biological activity of biomolecules.

Herein, we combined mussel-inspired molecule and bioorthogonal click chemistry for synergistically tailoring extracorporeal circuits and indwelling medical devices for antibacterial and anticoagulant multi-functions (Figure. 1). Clickable mussel-inspired peptide Ac-(DOPA)-Gly-(DOPA)-(Lys-PEG₅-Azide)-(DOPA)-Gly-(DOPA)-COOH (i.e., (DOPA)₄-Azide) was chemisorbed onto a pre-aminated surface to impart the surface with sufficient azide groups for subsequently conjugating of DBCO-modified molecules (i.e., Cu-DOTA-DBCO and DBCO-AMP) by bioorthogonal N₃-DBCO click reaction. The Cu-DOTA could catalytically generating
NO by which the platelet adhesion/aggregation could be strongly inhibited. Moreover, the grafted AMP possess high efficiency capabilities of killing bacterial. Thus, the bio-orthogonally co-immobilization of Cu-DOTA and AMP probably endowing the modified tubing with favored antimicrobial and antithrombotic dual functions. This design integrates NO-generating and antimicrobial peptide moieties into a tubing coating system by mussel-inspired adhesive peptide mimicking and bio-orthogonal click chemistry while involving only simple, specific, rapid, and reproducible procedures. Importantly, the Cu-DOTA-catalyzed NO generation and AMP might contribute synergistically and successfully to long-term antibacterial and antithrombosis. It is expected that this strategy can provide a facile approach for rational bioengineering of tubing and catheters with optimal multi-functions combating thrombosis and infection.

Results

Molecular Synthesis and Surface Functionalization
The clickable mussel-inspired peptide was prepared through standard Fmoc-mediated solid-phase peptide synthesis, as reported previously(33-36). Briefly, to simulate the multiple catechol structure in Mfps(37, 38), acetonide-protected Fluorenylmethyloxycarbonyl-DOPA(acetone)-OH was employed in introducing DOPA into the mussel-inspired peptide sequence. In order to promote the mussel-like molecular adhered to the substrates and leave available clickable groups for the following click reaction, tetravalent DOPA was integrated using an amino acid spacer and a PEG-connected azide to acquire a bio-clickable mussel-inspired adhesive peptide (DOPA)₄-Azide. In this work, as two key active molecules for anticoagulation and antibiosis, the nitric oxide (NO)-generating species Cu-DOTA and the AMP were connected to a PEG fragment with the DBCO group, respectively, to acquire the DBCO-capped bio-functional molecules. To obtained high purity of bio-functional molecules, the purity of (DOPA)₄-Azide, DBCO-AMP and DBCO-DOTA were purified (99.06%, 96.02% and 96.97%, respectively) by high-performance liquid chromatography (HPLC) (Figure. S1). After purification through HPLC, the three chemical-synthesized molecules were then characterized with nuclear magnetic resonance (NMR) spectrometry and electrospray ionization mass spectrometry (ESI-MS). As expected, the monoisotopic mass [M-H]⁻ of (DOPA)₄-Azide, [M+3H]³⁺ of DBCO-AMP, and [M+2H]²⁺ of DBCO-DOTA were detected at 1050.8, 871.7, and 607.2 Da, meeting their theoretical molecular weight 1052.1, 2612.1, and 1212.36, respectively (Fig. 2A–C). ¹H-NMR analysis also showed all diagnostic peaks of the three synthesized molecules and further confirmed the success of molecular synthesis (Figure. S2). The results verified the successful chemical synthesis of the bio-clickable adhesive peptide and DBCO- modified bio-functional molecules. To confirm the
formation of the Cu-DOTA-DBCO molecule, the measurement of electron paramagnetic resonance (EPR) was carried out. EPR analysis revealed that Cu$^{2+}$ was successfully chelated to the DBCO-DOTA, with the signals showing up at 3490-3430 mT (Figure. 2D).

Currently, most of the indwelling medical devices and extracorporeal circuits used in clinical are made of polymer materials. However, there have limited surface bioengineering strategies for polymeric devices as compared with the metal devices, possibly owing to the chemical inertness of biomedical polymer materials. To facilitate the surface functionalization of polymer material, in this study, a durable amine-containing coating(39) was performed on the surface of polyvinyl chloride (PVC, a conventional medical-materials approved for manufacturing blood-contacting device) substrates or tubes. Then, the mussel-inspired adhesive bio-clickable peptide was robust tethered on the PVC through catechol-amine reaction. Finally, the azide-functionalized PVC surfaces were easily connected with the obtained DBCO-capped NO catalyst (Cu-DOTA-DBCO) and DBCO-modified antimicrobial peptide (DBCO-AMP) by DBCO-N$_3$ bio-orthogonal reaction. The peptide binding and biomolecular grafting were monitored by Quartz Crystal Microbalance Dissipation Method (QCM-D). The results in Figure 2E demonstrated that 430.8 ng·cm$^{-2}$ of (DOPA)$_4$-Azide was steadily bound onto the aminated chips, implying the high efficiency and robust tethering onto the aminated PVC substrates. Then, the azide-modified chips were incubated with DBCO-AMP or Cu-DOTA-DBCO via bio-orthogonal conjugation. Analysis of QCM-D monitoring revealed that the grafting processes began within minutes, and the maximal grafting amount for DBCO-AMP and Cu-DOTA-DBCO were 696.9 and 594.8 ng·cm$^{-2}$, respectively (Figure. 2F-G,). Dual functionalization using a mixture of DBCO-AMP and Cu-DOTA-DBCO (1:1 in molar ratio) was further carried out, and the co-immobilized amount demonstrated a median value around 637.4 ng·cm$^{-2}$ (Figure. 2H).

To further investigate the changes of the chemical structure and composition of the aminated PVC before and after each grafting step, reflection absorbance Fourier transform infrared (RA-FTIR) and X-ray photoelectron spectroscopy (XPS) were carried out. After grafting of (DOPA)$_4$-Azide to the aminated surface (marked as Azide), the introduction of -N=N==N' peak by (DOPA)$_4$-Azide with azide groups in the FTIR spectra was nearly invisible, which may be due to the -N=N'=N' signal was limited by the detection limit of infrared. However, the band from 3100 to 3600 cm$^{-1}$ was remarkably broadened because of the -OH stretching vibration bands derived from the (DOPA)$_4$-Azide containing a moderate amount of phenolic hydroxyl and carboxyl groups, probably indicating the successful grafting of (DOPA)$_4$-Azide. The click-anchoring sites for DBCO-modified bio-functional molecules were provided through introducing the azide groups to the aminated surface. After clicking of Cu-DOTA-DBCO or/and DBCO-AMP to Azide
(marked as Cu-DOTA, AMP and Cu-DOTA&AMP, respectively), the appearance of the band in the FTIR spectra at 1530, 1440, 1235 cm\(^{-1}\) confirm the 1,2,3-triazole group produced by the successful bioorthogonal N\(_3\)-DBCO click reaction between the -N\(_3\) groups provided by Azide coating and the -DBCO groups of Cu-DOTA-DBCO or DBCO-AMP. Additionally, the appearance of band at 3060 cm\(^{-1}\) (=CH stretching) and the significant shift of the peak from 1633 to 1654 cm\(^{-1}\) further indicated the successful bio-conjugation of AMP. Although the band at 3060 cm\(^{-1}\) (=CH stretching) in the FTIR spectrum of Cu-DOTA&AMP nearly disappeared probably due to the successful clickable grafting of both molecules, the presence of similar significant shift of the band from 1633 to 1654 cm\(^{-1}\) also further confirmed the successful bio-conjugation of Cu-DOTA-DBCO and DBCO-AMP on Azide surface (Figure. 2I). XPS analysis provided additional confirmation of the successful surface engineering of (DOPA)\(_4\)-Azide, DBCO-AMP and Cu-DOTA-DBCO, as evidenced through the remarkable changes in the chemical compositions of the aminated PVC before and after each grafting step (Table S1). The changes in the contents of oxygen and nitrogen of the Azide confirmed the immobilization of (DOPA)\(_4\)-Azide on the aminated coating. Moreover, the copper element especial to the Cu-DOTA-DBCO were tested in both the Cu-DOTA-containing coatings, and the significant decrease in the contents of the copper of the Cu-DOTA&AMP coatings compared with the Cu-DOTA coatings, which probably confirmed the effective bio-conjugation of Cu-DOTA-DBCO and DBCO-AMP on the Azide coating. To further investigate the chemical components of the functionalized surfaces, the C 1s and N 1s peaks of XPS were implemented with peak fitting methods (Fig. S3). An increase of the peak was found at 286.2 eV in the C 1s high-resolution spectra of the Azide surface compared to that of aminated surface, which may be due to the introduction of C-O/C-N\(_3\) peak components by (DOPA)\(_4\)-Azide with azide groups and C-O structures. Additionally, an obvious decrease or increase of peaks value was found at 284.6 eV and 286.2 eV in the C 1s high-resolution spectra of the AMP and Cu-DOTA surfaces compared to that of Azide because of the introduction of C=C/C-C structure of the DBCO-AMP and Cu-DOTA-DBCO, implying the conjugation of AMP and Cu-DOTA on Azide surfaces (Figure. S3A). Finally, the content of each component in Cu-DOTA&AMP coatings was between the DOTA-Cu and AMP surfaces, which was consistent with the theoretical result of bio-conjugation Cu-DOTA-DBCO and DBCO-AMP (Table S2). The N 1s core-level spectrum of the aminated surface after grafting of (DOPA)\(_4\)-Azide was curve-fitted into five peak components with BEs at 398.07, 399.50, 400.46, 401.13 and 403.90 eV, attributable to the aromatic N, aliphatic N, Azide (N=N\(_{-}\)), R-N\(^+\) and Azide (-N\(^+\)) species, respectively (Figure. S3B). The Azide [(N)-N=]/[(-N\(^+\))] area ratio of around 2:1 was in good agreement with the characteristics of azide(40), indicating that the mussel-inspired adhesive bio-clickable peptide had
grafted on the aminated surface. Moreover, the absorption of non-azide protonated R-N+ was significantly enhanced, which probably is due to the aniline structure formed by Michael addition between the -NH2 groups of aminated surface and the benzodiazepines groups of (DOPA)4-Azide was easily protonated. Therefore, the disappearance of Azide (-N+) and ((N)-N=) species of the AMP, Cu-DOTA and Cu-DOTA&AMP surfaces further indicated that the Cu-DOTA-DBCO and DBCO-AMP was successful clicked on the azide-functionalized surface via bioorthogonal N3-DBCO click reaction (Table S3).

Altogether, in the above results, the potential of bio-orthogonal conjugation bio-functional molecule for fabrication of dual-functional surface was verified.

**Anti-Bacterial Property**

Infection is one of the major complications associated with extracorporeal circuits and indwelling medical devices, which causes significant mortality in clinic(41). To detect the antibacterial performance of the Cu-DOTA&AMP coating, Escherichia coli (E. coli) and Staphylococci epidermids (S. epidermids) were selected as typical strains causing infections after interventional procedures for antibacterial tests. We found that the surface azidation to the PVC substrates by (DOPA)4-Azide did not result in visible influence on bacterial growth as evidenced by antibacterial rates of 9.2% and 9.0% for E. col and S. aureus, respectively. However, the Cu-DOTA-grafted on azide-functionalized PVC led to inhibitory effects on bacterial growth, which may be attribute to the bactericidal ability of the leakage copper(II)-ions from Cu-DOTA. Moreover, a significantly growth of the antibacterial rate in AMP-coated PVC compared to the control and Azide groups was also observed. In addition, we have noted that the bio-orthogonal of Cu-DOTA&AMP on azide-functionalized PVC proved the excellent synergistic interactions on inhibiting S. epidermids and E. coli with bacterial killing activity nearly to 99% and 100%, respectively (Figure. 3A-C).

SEM analysis showed that a moderate number of S. epidermids and E. coli adhered on both PVC and Azide groups showed a state of rapid proliferation (Figure. 3D). In contrast, the surfaces functionalized by Cu-DOTA or AMP alone significantly inhibited the adhesion and proliferation of S. epidermids and E. coli. We have not noted that the bacteria membrane ruptured obviously on the Cu-DOTA, which might be owing to the insufficiently concentration of free of copper ions from Cu-DOTA. However, it was noteworthy that the bacterial membrane rupture was observed on the AMP and Cu-DOTA&AMP, which demonstrated the well retained ability of AMP (42) grafted by surface click chemistry to penetrate and destroy the bacterial membrane for killing bacteria. In addition, the excellent synergistic effects on the adhesion and proliferation of S.
epidermids and E. coli was exhibited at the bio-orthogonal of Cu-DOTA&AMP on azide-functionalized PVC surface.

Considering the long-term practical applications of the Cu-DOTA&AMP coating on implantable medical blood-contacting devices (e.g., central venous catheter and pacemaker), the long-term efficacy of its antibacterial efficacy for different time period at continuous immersion into PBS was tested (Figure. S4). The results revealed that antibacterial rates of Cu-DOTA&AMP-coated PVC for S. epidermids and E. coli after 30 days of soaking, but still suppressed 98% of bacterial growth of both S. epidermids and E. coli, implying the wonderful preservation of Cu-DOTA&AMP in antibacterial properties.

Altogether, in the results above shows that the AMP and Cu\(^{2+}\) from Cu-DOTA endows the long-term antibacterial properties of Cu-DOTA&AMP surface to prevent biofouling.

**In vitro NO Catalytic Release and Blood Compatibility Tests**

In our design, the DOTA chelated copper ions can decompose nitrosothiols into NO to suppress material-induced thrombosis(25, 26, 43). The catalytic release of NO as bioactive gas molecule was calculated to detect the NO catalytic ability of the Cu-DOTA&AMP coating. The NO catalytic release activity of Cu-DOTA&AMP surface was evaluated through a real-time chemiluminescent assay. For the in vitro experiments, the NO donor solution (10 μM S-nitrosoglutathione (GSNO) and 10 μM L-glutathione (GSH)) was prepared to mimic the physiological environment. NO real-time monitoring revealed that there is almost no catalytic release of nitric oxide from the AMP-coated PVC (Figure. 4A). As shown in **Figures 4B** and **4C**, the stable NO release patterns yield by the Cu-DOTA-coated PVC (5.7 × 10\(^{-10}\) mol cm\(^{-2}\) min\(^{-1}\)), suggesting the successful functionalization of in-situ release of NO. In contrast, the Cu-DOTA&AMP surface showed slightly lower NO flux (2.6×10\(^{-10}\) mol·cm\(^{-2}\)·min\(^{-1}\)) possibly due to the successful clickable grafting of both molecules. Considering the stable catalytic release of NO is critical to the implanted blood-contacting devices for long-term service *in vivo*, stability studies were prepared for Cu-DOTA&AMP through immersion into PBS for different time periods, and then their NO catalytic release ability was tested every seven days. The experimental result indicate that the NO catalytic release ability exhibited a minor decrease with the increase of soaking time, but stabilized around (0.5~1) ×10\(^{-10}\) mol·cm\(^{-2}\)·min\(^{-1}\) after more than 30 days (Figure. S5). Jointly, these results confirmed the possibility of the dual-functional Cu-DOTA&AMP surfaces for long-term service *in vivo*.

At the early post-implantation stage, thrombosis is a crucial problem associated with blood-contacting material. As a highly potent gaseous signal molecule, NO can activates soluble
guanylate cyclase through binding the haem moiety of soluble guanylate cyclase, contributing to cyclic guanosine monophosphate (cGMP) up-regulation and eventually inhibiting the activation and aggregation of platelets(44). To evaluate the ability of NO produced from Cu-DOTA on facilitating the cGMP synthesis of platelets, the cGMP analysis was carried out. Compared to the PVC and Azide groups, only the groups (Cu-DOTA and Cu-DOTA&AMP) containing Cu-DOTA both significantly facilitated the cGMP expression of platelets with the NO donor supplement, the grafted only AMP group had no charge (Figure. 4D). Moreover, we also noted that a slightly loss in the cGMP expression due to the loss of NO released by Cu-DOTA&AMP (Figure. 4C). In contrast, the expression of cGMP of platelets did not change without NO donor supplement (Figure. S6A), indicating the highly biological activity of NO generated from Cu-DOTA. Then, we further investigated the adhesion and activation of platelets on the dual-functional Cu-DOTA&AMP surface. Without donor supplement, a moderate number of activated platelets aggregate on all the experiment groups (Figure. S6B-D). Upon adding donor to catalytic release NO, most of platelet adhered on both PVC and Azide groups with a state of remarkable degree of activation and aggregation was observed. Furthermore, the grafted AMP only group still had obvious platelet adhesion and activation. By contrast, the grafted Cu-DOTA group substantially reduced platelet adhesion and activation with an inactive spherical state. Though the generation of NO induced by Cu-DOTA&AMP was a slightly decrease compared with the grafted Cu-DOTA group, it still remarkable inhibited the adhesion and activation of platelet (Figure. 4E-G). the result suggested the efficacy of NO gas molecules for the suppression of thrombogenesis in blood microenvironment.

Altogether, these results demonstrate the excellent anti-thrombogenic properties of the Cu-DOTA&AMP surface in vitro.

**Ex vivo Antithrombogenic Properties**

To be more clinically relevant, an *ex vivo* blood circuits experiment was further performed(45). The PVC tubing before and after surface engineering were assembled with clinically usable medical catheter and subsequently linked to a rabbit arteriovenous shunt circuit (Figure. 5A).

With the NO donor supplement, all the tubings were collected to evaluate the occlusive rates, blood flow rates and thrombus weights after ex vivo circulation for 2 h. We noted that there was only a tiny number of thrombi on the groups with the NO releasing, whereas severe thrombus formation was observed on the NO-free groups (PVC and AMP) (Figure. 5B and 5C). SEM analysis further confirmed that the groups containing Cu-DOTA significantly prevented the formation of thrombus. On the blood-contact surfaces of the bare PVC tubing and AMP-coated
tubing, there had severe thrombus with high crosslinking density of polymeric fibrin networks, red blood cells and activated platelets (Figure. 5D). Quantitative of these results further revealed that the significant reduction in thrombosis formation of the NO releasing groups through evaluating thrombus weight, occlusion rates and blood flow rates relative to the other groups (Figure. 5E-G). Total thrombus weight in NO releasing circuits was ten-fold reduced compared to the bare and only AMP-grafted groups (Figure. 5E). To conclude the occlusion rates throughout the circuits post-explant, the percent occlusion of lumen areas was calculated using computerized image analysis. Compared to the bare and only AMP-grafted groups, the Cu-DOTA&AMP coated circuit significantly reduced the thrombotic occlusion to 10% (Figure. 5F). Correspondingly, the blood flow rates revealed a consistent result (Figure. 5G). These results were also in line with the anti-thrombogenic property in vitro, described above.

Considering the long-term practical applications of the Cu-DOTA&AMP coating on implantable medical blood-contacting devices (e.g., pacemaker and central venous catheter), the long-term efficacy of its anticoagulant efficacy for different time period at continuous immersion into PBS was tested (Figure. S7). The results showed that the anti-thrombogenic ability showed an almost no decrease after 30 days of treatment with PBS, suggesting the promising application of Cu-DOTA&AMP coating in implanted/interventional blood-contacting devices for long-term use in vivo.

The above results also confirmed the excellent anti-thrombogenic property of the Cu-DOTA&AMP surface long-term use in vivo.

**Blood Biochemical Analysis**

Antithrombosis is the major property associated with blood-contacting materials in addition to the hemostatic materials. However, for those biomedical devices which undergo long-term or large-area contact with blood, e.g., the central venous catheter or cardiac pacemaker, their effects on blood composition or on liver and kidney functions should be systematically detected to ensure the safety for use in vivo. Considering the difference of total blood volume between rabbit and human (~200 ml for mature white rabbit and ~4000 ml for adult human), a 1.6 m long bare PVC or Cu-DOTA&AMP-coated tube (with inner diameter 3 mm) was selected to simulate the clinical application, and then respectively installed into the rabbit arteriovenous shunt circuit (Fig. 6A). After 0, 5, 30, and 60 min of blood circulation, the blood was collected for biochemical and physiological parameters were determined, including inflammatory response, blood coagulation and organ functions.
Coagulation evaluation showed that the control groups had a trend towards a higher blood clotting after prolonged contact with blood, which was characterized by the increased amount of F1+2 (an integral marker for the prothrombin activation) (Figure. 6C). However, activated partial thromboplastin time (APTT) decreased significantly in all groups, probably because that the catalytic release of NO was insufficient (Figure. 6B). After circulation, there had been a decrease in the number of platelets of bare PVC tubing, but the number of platelets of the Cu-DOTA&AMP-coated PVC tubing did not change (Figure. 6D). A device implanted in vivo can be quickly monitored by the immune system, then causing an inflammatory response. While the pro-inflammatory parameters of all groups had no obvious changes: the count of C3a (C3 cleavage fragment, indicating the classic or alternative way to activate the complement system, Figure. 6E), c-reactive protein (a kind of acute phase proteins in plasma was used as a measurement of acute inflammation, Figure. 6F), white blood cells (Figure. 6G), expression of tumor necrosis factor alpha (TNF-α) (a major acute inflammatory cytokines, Figure. 6I) and IL-10 (a recognized inflammatory and immunosuppressor, Figure. 6H). Such results indicate that all groups had no pro-inflammatory tendency. However, only the expression of TNF-α decreased significantly after cycling, which may be related to the anesthesia (46). All the proinflammatory indexes of the Cu-DOTA&AMP surface group had a consistent result with the bare PVC group used in clinical, suggesting that the coating did not further promote an inflammatory response of the material. To explore if the material and coating had serious toxicity to organs and tissues, the blood concentrations of the liver enzyme alanine aminotransferase (ALT) and the kidney parameter serum creatinine (Scr) were measured. As show in Figure 6J and K, both the PVC and Cu-DOTA&AMP surface presented no organ and tissue toxicity during circulation, and there was also no significant difference, verifying their biosafety.

**Discussion**

We develop here a biomimetic surface engineering strategy for fabricating multifunctional coating onto blood-contacting surfaces by masterly combining bioorthogonal conjugation chemistry with mussel-inspired adhesive peptide mimicking. To robustly binding clickable mussel-inspired peptide on tubing and catheters surface by catechol-amine chemistry, we functioned the polymeric device with a durable amine-bearing surface. The DBCO-modified functional molecules (e.g., Cu-DOTA-DBCO and DBCO-AMP) was effectively co-immobilized on tubing and catheters surface through the bioorthogonal conjugation chemistry. The antibacterial function of DBCO-AMP and the signal molecules effect of NO generated from Cu-DOTA-DBCO endowed the coating with the durable synergistic inhibition in growth and adhesion of bacteria.
and activation of platelets in vitro for one month, as well the capacity to efficiently restrain thrombogenesis \textit{ex vivo}. Our strategy presented a promising method to tailor a multifunctional surface and maintain efficient biological function, and may also be suitable for biomimetic surface engineering of many biomedical fields.

**Materials and Methods**

**Materials**

Dopamine, polyallylamine, CuCl\(_2\)-2H\(_2\)O, GSNO, GSH, cGMP Enzyme Immunoassay Kit, glutaraldehyde, were purchased from Sigma-Aldrich.

**Surface Amination of PVC**

Briefly, the PVC substrates (1.0 cm × 1.0 cm for antiplatelet test and 2.5 cm × 2.5 cm for antibacterial assessment) or tubes (inner diameter Φ=3.0 mm) were dipped into or perfused with the dopamine (DA) solution under optimal conditions (10 mM Tris-HCl, pH 8.5, 1 mg mL\(^{-1}\) DA) for 48 h. Then the polydopamine (PDA)-modified PVC substrates or tubes were flushed with deionized water and desiccated using N\(_2\) gas. To further tailoring surface amino functionalization of the PVC, the modified PVC substrates or tubes were subsequently soaked into the polyallylamine alkaline aqueous solution (pH=12) under ambient temperature for 12 h and finally cleaned with distilled water and desiccated by N\(_2\) for future use.

\((\text{DOPA})_4\text{-Azide, DBCO-AMP and DBCO-DOTA Synthesis:}\) The \((\text{DOPA})_4\text{-Azide and the DBCO-capped molecules (DBCO-AMP and DBCO-DOTA) were prepared through the Fmoc-mediated solid-phase synthesis technique(47).}\) With the help of China Peptides Co. Ltd. (Shanghai, China, purity > 95%). The Cu-DOTA-DBCO was prepared by mixing the CuCl\(_2\)-2H\(_2\)O with DBCO-DOTA in the aqueous solution by a mol ratio of 1:1.

**Surface Azidation of the PVC and Co-Grafting of the Cu-DOTA-DBCO and DBCO-AMP**

The aminated PVC substrates or tubes (termed as Aminate) were firstly immersed into DOPA\(_4\)-Azide (0.1 mg mL\(^{-1}\)) dissoveled by PBS (pH=8.0) under ambient temperature for 24 h, and then cleaned by deionized water and desiccated with a stream of N\(_2\), the azide-modified surface was termed as “Azide”. Subsequently, the azide-modified PVC was incubated with 2 mg mL\(^{-1}\) of Cu-DOTA-DBCO and/or 2 mg mL\(^{-1}\) DBCO-AMP under ambient temperature for 24 h to prepare the Cu-DOTA, AMP, and Cu-DOTA&AMP co-grafted surface. After co-grafting, the modified substrates or tubes were cleaned thorough deionized water and desiccated with a stream of N\(_2\) for subsequent use.

**Characterization**
To purify the synthesized molecules, HPLC was performed on an Agilent HPLC system with a Kromasil 100-5C18 column (5 μm, 4.6×250 mm, column temperature 25 °C). Mobile phases were buffer A (0.1% TFA in water) and Buffer B (0.1% TFA in acetonitrile). The flow was graded at a rate of 1 mL min⁻¹. Then injection volume and running time were 10 μL and 11 min respectively. The molecular weights of (DOPA)₄-Azide, DBCO-AMP and DBCO-DOTA were determined by ESI-MS (Sciex API 150EX LC/MS with Agilent 1100 HPLC). Buffer: 75%ACN/24.5%H₂O/0.5%Ac; flow rate: 0.2 mL min⁻¹; run time: 1 min. A Nicolet model 5700 instrument was used to take Reflection Absorbance Fourier Transform Infrared (RA-FTIR) spectrum. The surface elemental compositions were detected by X-ray photoelectron spectroscopy (XPS) (K-alpha, ThermoFisher, USA), with an excitation source monochromatic Al Kα (1486.6 eV). The synthesized molecules was analyzed by proton nuclear magnetic resonance (¹H-NMR) spectrum (Bruker AVANCE III 400). 10 mg sample were used to measure Electron Paramagnetic Resonance (EPR) spectra on a Bruker EPR EMXPlus (X-band is 9.85 GHz, field modulation is 100 kHz, and the power is 0.2 mW, samples were loaded in capillary tube).

**QCM-D Analysis**

The real-time monitoring of the grafting of the (DOPA)₄-Azide and the DBCO-capped molecules (DBCO-AMP or/and Cu-DOTA-DBCO) were used by a Quartz Crystal Microbalance Dissipation Method (QCM-D, Q-sense AB, Sweden). In detail, the Au-coated quartz crystal (diameter of Au film 1 cm) was firstly surface aminated. Then, the piezoelectric quartz crystal sensors were excited at a fundamental frequency (5 MHz) and the change in frequency (Δf) was monitored for the third, fifth, seventh, ninth, eleventh, and thirteenth overtones. When the excitation stopped, recorded the changes of the resonance frequencies (ΔF) and those of the relaxation (ΔD) of the vibration at the five frequencies, and installed the surface aminated quartz crystal in the QCM-D chamber and injected PBS buffer at 50 μL min⁻¹ continuously until the QCM-D traces maintained steady, followed by the buffer pump with the identical speed. After that, (DOPA)₄-Azide (0.1 mg mL⁻¹) in PBS was injected into the surveying chamber in touch with the crystal with the identical speed, and eventually washed with PBS. Subsequently, 2 mg mL⁻¹ of Cu-DOTA-DBCO and/or 2 mg mL⁻¹ of DBCO-AMP was injected into the surveying chamber in touch with the crystal with the identical speed, and eventually washed with PBS. Finally, the amounts of the Cu-DOTA-DBCO or/and the DBCO-AMP grafted on the surfaces were calculated based on the Sauerbrey equation[48].

**Antibacterial Activity**
The detailed procedure has been described elsewhere(49). Briefly, The *S. aureus* and *E. coli* were precultured in agar solid medium positioned in a 37°C incubator for 24 h and passaged twice to acquire the monoclonal bacterial. Picked the fresh bacterial colonies (1-3 rings) and dissolved with a solution containing 0.2% liquid medium and 99.8% saline. The bacteria concentration was restructured to $5.0 \times 10^5$ to $10^6$ CFU mL$^{-1}$ by tenfold increasing sequential diluting. 100 μL of the suitable concentration of bacterial solution was added onto the surface of samples and covered using a soft polyethylene (PE) membrane to maintain a well spread liquid film on the substrate. All samples were positioned in a 37°C incubator for 24 h. Then, the bacteria on the surfaces were flushed and diluted with saline solution (1 mL). 20 μL of the bacterial solution was distributed on the solid medium and incubated in a 37°C incubator. Lastly, the colonies cultured on the surface of solid medium were quantified and observed by SEM after 24 h.

**Catalytic Generation of NO**

A chemiluminescence NO analyzer (NOA, Seivers 280i, Boulder, CO) was used to determine the real-time generation rate of NO. In brief, the AMP, Cu-DOTA and Cu-DOTA&AMP-modified PVC substrates (5 mm × 10 mm) were immersed into PBS with NO donor containing SNAP (10 μM) and GSH (10 μM). Upon the reaction was carried out, the NO-generated was conveyed into the NO analyzer by a stream of N$_2$. Finally, the NO flux was calculated using the calibration line, which has been reported in details elsewhere(50).

**Platelet Adhesion and Activation**

The bare and AMP-, Cu-DOTA and Cu-DOTA&AMP-modified PVC were cultured by 0.5 mL of platelet-rich plasma (PRP). After culture for 30 min at 37 °C, the samples were rinsed 3 times with saline solution and then fixed with 4% paraformaldehyde solution overnight. After further dehydrated and dealcoholized, the adhered platelets on the sample surfaces were evaluated using scanning electron microscope (SEM) (ZEISS EVO 18). Given the low synthesis of endogenous in PRP, the samples were cultured in two groups of PRP supplemented with or without NO donor containing SNAP (10 μM) and GSH (10 μM).

**cGMP Expression of Platelets**

Expression level of cGMP of platelets adhered on each sample was measured by the human cGMP ELISA kit. Samples were firstly cultured in 1 mL of PRP at 37 °C for 30 min with or without NO donor. Therewith, 100 μL of triton-X solution (10%) was added onto the above sample and followed by sonication. The obtained above fragmentized PRP solution was centrifugated for 15 min at 3000 rcf, and the supernatant was collected for ELISA test.

**Ex vivo Hemocompatibility Test**
The animal experiments obey the rules and guidelines of the China Ethical Committee and Laboratory Animal Administration Methods. The detailed experimental process has been described elsewhere (50). Briefly, the Cu-DOTA-DBCO or/and DBCO-AMP were firstly immobilized on the lumen surface of the PVC tubings. After being anesthetized by 30 mg mL\(^{-1}\) pentobarbital sodium (1 mL per kg), the left external jugular vein and right carotid artery of New Zealand white rabbits (above 2.5 kg) were exposed and cannulated. Then, the catheter was linked to the cannulas, forming a closed loop. The unmodified-, and AMP-, Cu-DOTA and Cu-DOTA\&AMP-modified PVC tubings were taken out after \textit{ex vivo} circulation for two hours and washed with saline solution. The occlusive rates were calculated by the cross-sections photographs of the tubings. Under the same pressure pump condition, the blood flow rate was calculated after circulation and normalized to that before the circulation. The thrombus formed on the luminal surface of the tubes was weighed, then analyzed by SEM after fixed with 4% paraformaldehyde solution.

**Blood Analysis after \textit{ex vivo} Blood Circulation**

The blood from New Zealand white rabbits was extracted for haematology analysis and blood biochemistry assay after \textit{ex vivo} blood circulation. In the case of this experiment, each rabbit only received one circuit (e.g., unmodified PVC or Cu-DOTA\&AMP-modified tubing). Blood analysis including the APTT (51), F1+2 (52), C3a (53), CRP (54), WBC, PLT, IL-10 (55), TNF-\(\alpha\) (56), ALT (57) and CRE (58) was performed by using the blood drawn from the running circulation at different periods of time (0, 5, 30 and 60 min). A portion of the freshly collected whole blood with and without anticoagulants were centrifuged (2500 rcf at 4 °C for 15 min) to collect plasma and serum, respectively. To test whether the coated catheter would have side effects on blood composition or on liver and kidney functions, a single prolonged catheter was selected to simulate the clinical application. The total ECC tube for human is several meters long, the area of blood-material interface of it is as high as \(10^2-10^3\) cm\(^2\) as speculated. Considering the difference of total blood volume between rabbit and human (~200 ml for mature white rabbit and ~4000 ml for adult human), we selected a 1.6 meters long PVC tube with inner diameter=3 mm, and coated the lumen surface. The enlarge contact area facilitate us to disclose the different effect between the modified and unmodified (control) catheter.

APTT was measured by manual tilt-tube method (59). The CRP, TNF-\(\alpha\), C3a, IL-10 and F1+2 were measured by ELISA Kit (Rabbit CRP/TNF-\(\alpha\)/C3a/IL-10/F1+2 ELISA KIT, ZC-52314/ ZC-52984/ZC-52409/ZC-52381/ZC-52601, Shanghai ZCIBIO Technology Co., Ltd.), according to the specifications. WBC and PLT were measured by animal automatic blood cell analyzer (Shenzhen
Mindray, BC-2800Vet). ALT and Scr were measured by animal biochemical analyzer (Shenzhen Mindray, BS-240VET).

**Statistical Analysis**

All the group data are expressed as mean with standard deviation for every sample unless specified otherwise. All the experiments were repeated independently at least three times, if not otherwise indicated. Student’s t-test and one-way ANOVA in GraphPad Prism 9.0 (GraphPad Software) was performed for statistical analyses from different groups. Significance was denoted as follows: (ns: p > 0.05, *: p ≤ 0.05, **: p ≤ 0.01, ***: p ≤ 0.001, ****).

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Figure. 1. Fabrication of Cu-DOTA&AMP surface with anticoagulation and antibacterial properties. (A) Chemical structure of the clickable mussel-inspired peptide [(DOPA)$_4$-Azide, Ac-(DOPA)-Gly-(DOPA)-(Lys-PEG$_5$-Azide)-(DOPA)-Gly-(DOPA)-COOH], DBCO-modified antimicrobial peptide (DBCO-AMP) and DBCO-capped NO catalyst (Cu-DOTA-DBCO). (B) Surface co-grafting on representative medical devices through mussel-inspired catechol-amine reaction and bioorthogonal click chemistry. (C) Realization of anticoagulation and synergistic antibacterial properties of Cu-DOTA&AMP surface.
Figure 2. Synthesis and grafting of (DOPA)$_4$-Azide, DBCO-AMP and Cu-DOTA-DBCO molecules on aminated surface. (A-C) Electrospray ionization mass spectrum of (DOPA)$_4$-Azide, DBCO-AMP, and DBCO-DOTA. (D) ESR spectrum of DBCO-DOTA with or without chelated copper ion. (E) Real-time monitoring of the grafting amount of (DOPA)$_4$-Azide on aminated surface using QCM. (F-H) Real-time monitoring of the grafting amount of DBCO-AMP, Cu-DOAT-DBCO and DBCO-AMP/Cu-DOTA-DBCO on azide surface using QCM. (I) RA-FTIR spectra of aminated surface after grafting with (DOPA)$_4$-Azide, DBCO-AMP, Cu-DOAT-DBCO and DBCO-AMP/Cu-DOTA-DBCO. Data are presented as mean ± SD (n=4).
Figure. 3. Anti-bacterial property. (A) Typical E. coli and S. epidermids colonies after 24 h incubation on the bare and modified PVC plates. Antibacterial rate of modified PVC that calculated from the number of colonies against (B) E. coli and (C) S. epidermids. (D) SEM images of E. coli and S. epidermids that adhered or colonized on the bare and modified PVC plates. All scale bars from the same raw are 500 or 5000 nm. Data presented as mean ± SD and analyzed using a one-way ANOVA, **p < 0.01, ***p < 0.001.
Figure 4. In vitro NO catalytic release and In vitro blood compatibility tests. NO generation rate of (A) AMP, (B) Cu-DOTA and (C) Cu-DOTA&AMP monitored by real-time NOA. (D) Concentration of cGMP synthesized by platelets. (E) SEM images (all scale bars from the same raw are 20 or 5 µm), (F) adhesion number and (G) activation rate of the adhered platelets on bare and modified PVC. Data presented as mean ± SD and analyzed using a one-way ANOVA, **p < 0.01, ***p < 0.001.
Figure. 5. Ex vivo hemocompatibility of the Cu-DOTA&AMP surfaces. (A) Schematic illustration of arteriovenous (AV) shunt model connected to the rabbit. Photographs of (B) side and (C) cross-section view of the bare and modified PVC tubes after blood circulation. (D) Lumen surface morphology of each sample after blood circulation characterized by SEM. (E) Thrombus weight, (F) occlusion and (G) blood flow rate of bare and modified PVC tubes following the blood circulation. Data presented as mean ± SD and analyzed using a one-way ANOVA, **p < 0.01, ***p < 0.001.
Figure 6. **Blood analysis by ex vivo blood circulation.** (A) Circulation model on rabbit for blood analysis. Blood parameters including (B) activated partial thromboplastin time (APTT), (C) F1+2 (an integral marker for the prothrombin activation), (D) the number of platelets (PLT), (E) C3a (C3 cleavage fragment, indicating the classic or alternative way to activate the complement system), (F) c-reactive protein (CRP), (G) white blood cells (WBC), (H) IL-10 (a recognized inflammatory and immunosuppressor), (I) tumor necrosis factor alpha (TNF-α), (J) the liver enzyme alanine aminotransferase (ALT) and (K) the kidney parameter serum creatinine (Scr) were detected to reveal the hemocompatibility and liver/kidney safety after exposing the bare and Cu-DOTA&AMP grafted catheters to the circulating blood. Data are presented as mean ± SD (n=4).
Supplementary Materials

Supplementary material for this article is available at http://advances.sciencemag.org.

Supplementary Text

Figs. S1 to S7

Figure. S1. High-performance liquid chromatography spectrum of (DOPA)4-Azide, DBCO-AMP and DBCO-DOTA.

Figure. S2. 1H NMR spectrum of (DOPA)4-Azide, DBCO-AMP and DBCO-DOTA.

Figure. S3. High resolution spectra of C1s and N1s signals on the different surfaces.

Figure. S4. Stability of the antibacterial ability of Cu-DOTA&AMP coating.

Figure. S5. Stability of the NO catalytic release ability of Cu-DOTA&AMP coating.

Figure. S6. In vitro blood compatibility tests without NO donor.

Figure. S7. Stability of the Anticoagulation ability of Cu-DOTA&AMP coating.

Tables S1 to S3

Table S1. The atomic compositions of different surfaces.

Table S2. The high-resolution C1s compositions of different surfaces.

Table S3. High-resolution N1s compositions of different surfaces.

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