Nitric oxide produced by endothelial cells (EC) which line the walls of all blood vessels plays an important role in preventing platelet activation, adhesion, and, ultimately, thrombosis at the EC/blood interface.[1,2] We have reported previously that
plasticized polymeric films doped with a lipophilic copper(ii) complex decompose S-nitrosothiols (RSNO), such as S-nitrosoglutathione (GSNO), to NO at their interface in the presence of naturally occurring reducing equivalents (e.g., glutathione (GSH)).

Indeed, it is well known that Cu^{II}/Cu^{I} ions can catalyze the generation of NO from RSNO species (see the Supporting Information). Furthermore, it was reported that peptide- and protein-bound Cu^{II} can be reduced to Cu^{I} by a thiol to generate NO from RSNOs. It now appears that appropriate Cu^{II}/Cu^{I} organic ligand complexes can carry out similar redox chemistry. Hence, it should be possible to utilize endogenous reducing equivalents (e.g., ascorbate, GSH, and cysteine (CySH)) and the endogenous NO precursors (e.g., GSNO, S-nitrosocysteine (CysNO), and S-nitrosoalbumin (AlbSNO)) in blood to generate locally enhanced NO levels at a polymer/blood interface by immobilizing catalytic Cu^{II}/Cu^{I} sites within the polymeric material. Although the exact concentrations of RSNOs in normal human blood are in debate, the levels of RSNOs and required reducing equivalents are thought to be in the nanomolar to micromolar range. As very low levels of NO (< 1 nm) can be effective in inhibiting platelet function, even a small fractional conversion of endogenous RSNO species into NO at a polymer/blood interface could be beneficial in lowering the risk of thrombus formation at such interfaces.

Herein, we report the synthesis of a modified Cu^{II}-cyclen (cyclen = 1,4,7,10-tetraazacyclododecane) complex covalently attached to a cross-linked poly(2-hydroxyethyl methacrylate) (pHEMA) polymer to generate NO from naturally occurring RSNOs at the physiological pH value. Further, it will be shown that spontaneous NO generation from RSNOs which exist in fresh sheep’s blood can be achieved with this novel material.

Cyclen is known to have very high binding constant with Cu^{II} (10^{23} M^{-1}) and square-pyramidal structure, thus allowing this complex to possess an open site(s) at which interactions with reducing equivalents and RSNO species are possible. Herein, a propylmethacrylate derivative of cyclen (1023 C, RT, 12 h (76%); b) 3-bromopropionol (3.4 equiv), Na2CO3 (1.2 equiv), NaI (1.3 equiv), CH3CN, 80 °C, 12 h (88%); c) methacryloyl chloride (5.4 equiv), Et3N (9.5 equiv), THF, -20 °C, RT, 1 h (59%); d) trifluoroacetic acid (TFA), CH3Cl2, 0 °C, RT, 12 h (quantitative yield); e) 1. HEMA (66.0 wt %), ethyleneglycol dimethacrylate (EGDM; 2.1 wt %), azobisisobutyronitrile (AIBN; 0.4 wt %), 65 °C, 12 h; 2. soaked and washed with a solution of aqueous NaOH/EtOH (1:10, v/v, pH 11–12); f) CuCl2·2H2O (1 equiv vs 5) in EtOH, 65 °C, 4 h.

Figure 1 illustrates the NO generation that is achieved when a small piece of the Cu^{II}-cyclen-pHEMA film is placed in and then removed from a solution of GSNO and GSH in deoxygenated PBS buffer. The NO generated is monitored continuously with a chemiluminescence NO analyzer (NOA). As shown in Figure 1a, a blank experiment in which a hydrogel film of 6 (not treated with copper) was placed in the test solution did not generate any detectable NO. This experiment demonstrates that, as expected, copper ions are key for NO generation. In a second blank experiment, a cross-linked pHEMA film of the same size was prepared by the reaction of HEMA, EGDM, and AIBN, but without copolymerization of 5, and further treated with copper ions by using...
leaching of copper ions, as measured by AA spectrophotometric analysis (the films soaked in the solutions still retained more than 64% of their original copper content after 15 days at 37.5°C; see the Supporting Information), and this leaching resulted in a decrease in the apparent level of NO generation (25–50% decrease) relative to a control of the same film (not soaked in solutions of GSH and/or GSNO). However, the $\lambda_{\text{max}}$ value of the film, as shown by UV/Vis spectroscopic analysis, did not change in location before and after such soaking, thus suggesting that GSH (a reducing agent) and glutathione disulfide (GSSG, a by-product of the NO-generation reaction), both strong ligands for copper ions,[16,17] do not bind irreversibly as ligands for the covalently linked complex. A more critical test was the examination of NO generation by films of the new CuII–cyclen–pHEMA 7 material (radius: 2.8 mm; thickness: 0.2 mm; type: disk) from GSNO before and after the exposure to animal blood for a finite period. It was found that NO generation from GSNO is maintained after soaking such films in 10 mL of the sheep plasma for 3 days at 4°C or whole blood for 1 day at 4°C (see the Supporting Information). These results indicate that the deactivation of 7 caused by endogenous free thiols, proteins, and other blood components may not be a significant issue. Clearly, a large number of in vivo studies over varying time periods will be required to fully assess the true NO-generation ability and resultant thromboresistance of the new polymer under physiological conditions.

To further demonstrate that the CuII–cyclen–pHEMA polymer can generate NO spontaneously in fresh whole blood, an amperometric RSNO sensor was constructed using an interpenetrating network (IPN) film of 7 and hydrophilic polyurethane (HPU, Tecophilic, SP-93A-100) as an outer film on a NO-selective electrochemical sensor (see the Supporting Information). The response of a control NO sensor without the polymer containing the CuII–cyclen complex (but with an HPU outer membrane) was also monitored in the same blood samples. Each sensor was precalibrated for an intrinsic direct amperometric response to NO (see the Supporting Information), and then the NO and RSNO levels were measured, respectively, in the fresh sheep’s blood diluted with PBS buffer at 37°C. Figure 2 shows that the NO sensor without the catalytic CuII–cyclen–pHEMA polymer site in the outer membrane has relatively little amperometric response when placed in the blood sample, whereas the difference in NO responses between the RSNO and NO sensors (equivalent to ≈ 25 nm NO, calculated from calibration curves) appears to be due to the decomposition of RSNO species in the sheep’s blood by the IPN film of 7 and HPU. Therefore, this experiment confirms that the CuII–cyclen complex covalently attached to pHEMA generates NO from endogenous substrates in the sheep blood in vitro.

In conclusion, a CuII–cyclen complex was covalently attached to pHEMA, and the resulting material is capable of generating NO from naturally occurring RSNO species, both in buffer solution and directly when in contact with fresh whole blood. The presence of endogenous reducing equivalents is required for such catalytic chemistry to take place. Assessment of the stability of the new CuII–cyclen–pHEMA 7 material suggests that the slow leaching of copper ions from
the polymer film may cause a measurable decrease in NO generation with time, but copper deactivation is not a significant problem. Although fundamental questions with respect to the exact mechanism by which 7 generates NO remain, the data presented herein suggest that this new material may prove beneficial as a potentially new thromboresistant blood-contacting material for biomedical applications with contact times of hours to several days. Experiments in vivo to demonstrate the effectiveness of this material in reducing thrombus formation will begin shortly.

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