

플루로키나제가 고정화된 폴리우레탄의 항혈전성 향상에 관한 고찰

유규하[○], 박선양*, 한동근**, 김영하**, 민병구

서울대학교 의과대학 의공학 교실, 내과학교실 *

한국 과학기술 연구원 고분자 설계 연구실 **

Antithrombogenicity of Lumbrokinase - Immobilized Polyurethane

Gyu Ha Ryu [○], Seonyang Park *, Dong Keun Han **, Young Ha Kim **, and Byounggoo Min

Department of Biomedical Engineering, Seoul National University
College of Medicine, 28 Yungun-Dong, Jongno-Gu, Seoul 110-744, Korea

*Dept of Internal Medicine, Seoul National University College of Medicine

**Polymer Chemistry Lab., KIST, P.O. Box 131, Cheongryang, Seoul 130-650, Korea

INTRODUCTION

Thrombotic obstruction is one of the most serious sequelae of artificial organs implanted into a living body. In 1983, Mihara et al.[1] extracted six fractions of strong and novel fibrinolytic enzymes (lumbrokinase, LK) from the earthworm, *Lumbricus rubellus*. These enzymes were very stable and showed greater antithrombotic activity than that of currently used antithrombotics. In this work, we immobilized an LK fraction showing the least plasminogen-degrading activity on polyurethane (PU) surface to investigate its enzymatic activity and antithrombotic activity.

MATERIALS AND METHODS

The methanol extracted PU surface was treated with 3%(w/v) maleic anhydride methylvinyl ether copolymer (MAMEC) solution and finally incubated in a LK solution in PBS (pH 7.4). The surface was characterized by ATR-FTIR, ESCA, and dynamic contact angle. The quantification of immobilized LK was performed by a dye-binding method. The immobilized LK activity was estimated by the fibrin plate method and caseinolytic activity assay. The antithrombotic activity was evaluated by *in vitro* ¹²⁵I-fibrinogen adsorption in fresh whole blood and ^{99m}Tc-platelet adhesion tests. The distribution of radiolabelled fibrinogen and platelet was visualized by autoradiography. The occlusion time through *ex vivo* rabbit A-A shunt was also determined.

RESULTS AND DISCUSSION

The immobilization of LK on PU surface was confirmed by IR and ESCA. The content of immobilized LK was 24 µg/cm², which was almost the same amount as that of immobilized albumin on the same surface. In contact angle measurement, LK-immobilized PU displayed hydrophilic property. The unit activity of immobilized LK was 18U/cm² and the relative activity ratio of immobilized LK to the soluble LK was about 34 % (Table 1). The ¹²⁵I-fibrinogen adsorption experiment in fresh whole blood showed that less fibrinogen was adsorbed on LK-immobilized surface than on PU and PU-MAMEC controls (Figure 1). Although platelet adhesion was initially enhanced on LK-immobilized surface compared with PU-MAMEC surface, less platelets remained on LK-immobilized surface as time elapsed and showed reduced thrombogenicity (Figure 2). The *ex vivo* occlusion time of untreated PU and PU-MAMEC surface were only 32 and 42 min, respectively. But that of LK-immobilized PU was extended to 140 min. These results suggested that the immobilized LK on PU surface might have decreased fibrinogen adsorption by digesting the adsorbed fibrinogen and inhibited platelet adhesion/aggregation to the artificial surface as well, indicating that the LK-immobilized surface was highly antithrombotic.

ACKNOWLEDGEMENT

The work was supported by the Highly Advanced National Project of Ministry of Science and Technology, Korea.

REFERENCES

1. Mihara H, et al., *Thromb. Haemost.*, 50, 258, 1983

Table 1. Characteristics of immobilized lumbrokinase

Assay	Mean ± S.D.
Concentration (µg/cm ²)	24 ± 2.0
Unit Activity (IU/cm ²)*	18 ± 2.6
Specific Activity (IU/µg)	0.8 ± 0.1
Efficiency (%)	34 ± 2.5

* Incubation with alpha-casein for 30 min at 37°C

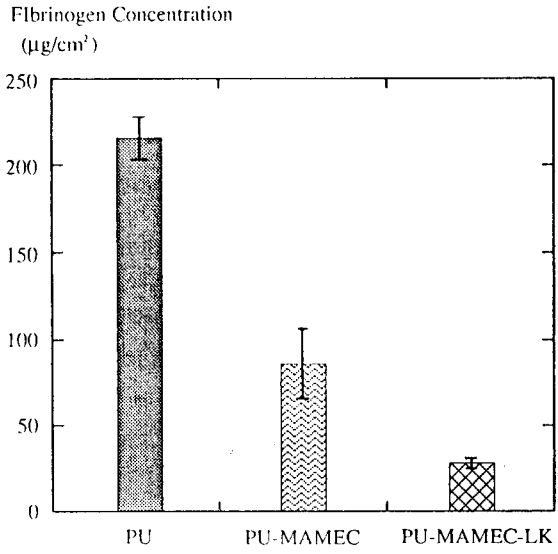


Figure 1. ^{125}I -Fibrinogen adsorption on polymeric surfaces

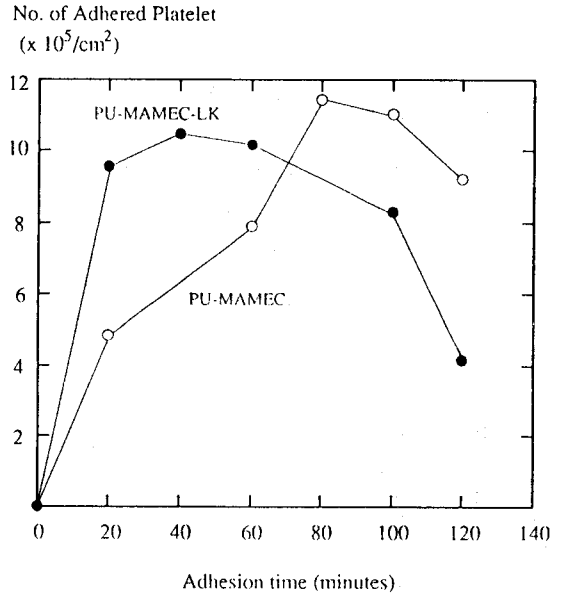


Figure 2. $^{99\text{m}}\text{Tc}$ -Platelet adhesion on polymeric surface