

당뇨병성 신증에서 전장 MMP-2와 세포내 N-말단 결실 isoform의 발현

부산대학교 의학전문대학원 내과학교실¹, 부산대학교병원 간호부², 캘리포니아대학교 재향군인의료센터 내과³

박수민¹, 송상헌¹, 곽임수¹, 이민영¹, 성은영¹, 이하린¹, 김일영¹, 이동원¹, 이수봉¹
정우진¹, 박종만¹, 김수정², 백영지², 박경주², 데이비드 H. 로베트³

The Expression of Full Length MMP-2 and Intracellular N-terminal Truncated MMP-2 Isoform in Diabetic Kidney Disease

Su Min Park¹, Sang Heon Song¹, Ihm Soo Kwak¹, Min Young Lee¹, Eun Young Seong¹
Harin Rhee¹, Il Young Kim¹, Dong Won Lee¹, Soo Bong Lee¹, Woo Jin Jung¹
Jong Man Park¹, Su Jeung Kim², Young Ji Beack², Kyeongjoo Park², David H. Lovett³

Department of Internal Medicine¹, Pusan National University School of Medicine, Busan, South Korea
Nursing Part² of Pusan National University Hospital, Busan, South Korea
Department of Internal Medicine³, San Francisco Department of Veterans Affairs Medical Center,
University of California San Francisco, San Francisco, California

Background: Recently, matrix metalloproteinase-2 has been regarded as a central of injury mechanism in heart and kidney. The ultrastructural examination of MMP-2 transgenic renal tubular epithelial cells demonstrated mitochondrial structural alterations and subsequent investigation led to the discovery of a novel intracellular N-terminal truncated isoform of MMP-2 (NTT-MMP-2) generated by activation of an alternative intronic promoter. The aim of this study is to explore the expression pattern and the role of full length MMP-2 (FL-MMP-2) and NTT-MMP2 in diabetic kidney disease.

Method: In vitro study, we tested glucose stimulation effects on HK2 cells. High, moderate, normal glucose were 30 mM, 15 mM, 5.6 mM. Time dependent experiments for 2hr, 24hr, 48hr, 72hr were conducted and dose-dependent experiments for normal, high and cyclic for the time of 48hr were performed. Also, real-time PCR was done for quantitative analysis of FL-MMP-2 and NTT-MMP-2.

Results: In vitro study using HK2 cells, high glucose stimulation induced the expression of FL-MMP-2 according to time and the expression at 72hr was higher by 3.4 times compared with control cells. Although the expression of NTT-MMP-2 was increased by high glucose, the peak expression time was different with FL-MMP-2 and the expression at 2hr was higher by 3.4 times compared with control cells. Also, the expression of NTT-MMP-2 was decreased according to time after 2hr stimulation. In dose-dependent experiments, cyclic stimulation induced higher expression of FL-MMP-2 and NTT-MMP-2 compared with control cells (FL-MMP-2; control vs. cyclic 1:4.16, NTT-MMP-2; control vs. cyclic 1:2.13). In streptozotocin-induced diabetic mice, the expression of FL-MMP-2 and NTT-MMP-2 was increased diffusely in the cytoplasm in renal tubular cell compared with normal control mice. In detail, FL-MMP-2 was expressed constitutively and the intensity or stained area were increased by induction of diabetes. However, NTT-MMP-2 was not expressed in normal control mice and was highly expressed after induction of diabetes mainly in renal cortex.

Conclusion: Glucose stimulation induced the expression of full length MMP-2 and intracellular N-terminal truncated MMP-2 isoform in HK2 cell. Furthermore, experimental diabetes influenced on full length MMP-2 expression and induced intracellular N-terminal truncated MMP-2 isoform in animal model.

Key Words: 당뇨병성 신증, 전장 MMP-2, N-말단 결실 MMP-2
Diabetic kidney disease, FL-MMP-2, NTT-MMP-2