Receptor for advanced glycation end products up-regulation in cerebral endothelial cells mediates cerebrovascular-related amyloid  $\beta$  accumulation after Porphyromonas gingivalis infection

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論 文 名 : Receptor for advanced glycation end products up-regulation in cerebral endothelial cells mediates cerebrovascular-related amyloid β accumulation after *Porphyromonas gingivalis* infection (脳内皮細胞における終末糖化産物受容体の亢進は、*Porphyromonas gingivalis* 感染による脳 血管が関与するアミロイドβの蓄積を誘導する)

区 分 :甲

## 論文内容の要旨

Cerebrovascular-related amyloidogenesis is found in over 80% of Alzheimer's disease (AD) cases, and amyloid  $\beta$  (A $\beta$ ) generation is increased in the peripheral macrophages during infection of Porphyromonas gingivalis (P. gingivalis), a causal bacterium for periodontitis. In this study, we focused on receptor for advanced glycation end products (RAGE), the key molecule involves in Aß influx after *P. gingivalis* infection to test our hypothesis that AB transportation from periphery into the brain, known as "Aβ influx", is enhanced by *P. gingivalis* infection. Using cultured hCMEC/D3 cell line, in comparison to uninfected cells, directly infection with P. gingivalis (multiplicity of infection, MOI = 5) significantly increased a time-dependent RAGE expression resulting in a dramatic increase in AB influx in the hCMEC/D3 cells; the P. gingivalis-up-regulated RAGE expression was significantly decreased by NF-kB and Cathepsin B (CatB)-specific inhibitors, and the *P. gingivalis*-increased I $\kappa$ B $\alpha$  degradation was significantly decreased by CatB-specific inhibitor. Furthermore, the P. gingivalis-increased AB influx was significantly reduced by RAGE-specific inhibitor. Using 15-month-old mice (C57BL/6JJmsSlc, female), in comparison to non-infection mice, systemic *P. gingivalis* infection for three consecutive weeks  $(1 \times 10^8 \text{ CFU/mouse}, \text{ every } 3 \text{ CFU/mouse})$ days, intraperitoneally) significantly increased the RAGE expression in the CD31-positive endothelial cells and the A<sup>β</sup> loads around the CD31-positive cells in the mice's brains. The RAGE expression in the CD31-positive cells was positively correlated with the A $\beta$  loads. These observations demonstrate that the up-regulated RAGE expression in cerebral endothelial cells mediates the Aß influx after P. gingivalis infection, and CatB plays a critical role in regulating the NF-κB/RAGE expression.