Role of Antibody Anti-AGE on the Expression of Nephrin and Rage in Primary Glomerulus Cell Culture Exposed to AGE

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ABSTRACT

Nephrin is associated with the initial stage of the loss of the permeability barrier in diabetic nephropathy. Interaction AGE-RAGE increases angiotensin II on Renin Angiotensin-Aldosterone System (RAAS) and activation of protein kinase c (PKC) which induce alterations in nephrin mRNA expression. Alterations of nephrin expression induce transformation of slit membrane structure and the permeability changes at the glomerular filtration barrier. Anti-AGE vaccination once may cause the changes of nephrin and RAGE expression and can prevent progression of diabetic nephropathy. This study used primary glomerulus cell culture obtained from renal of Wistar rat aged 3 months, weighing 200-300 grams and assigned into negative control group that exposed to BSA 100 µg/mL, positive control group that exposed to AGE-BSA 100 µg/mL, polyclonal anti-AGE antibody 5 µg/mL and AGE-BSA 100 µg/mL and treatment group 2 that exposed to monoclonal anti-CML antibody 5 µg/mL and AGE-BSA 100 µg/mL. Paired t-test with a 0.05 level of confidence results showed that there were significantly different v in level of RAGE expression between experimental groups with control groups. Administration of polyclonal anti-AGE antibody decreased RAGE expression compared to negative control (p = 0.188) and positive control (p = 0.000). RAGE expression did not differ significantly in administration of monoclonal anti-CML antibody compared to negative control but significant with positive control. Administration of monoclonal anti-CML antibody inhibited increasing of nephrin expression compared to negative and positive control (p = 0.73; 0.125). In conclusion, this study suggested that administration of polyclonal anti-AGE or monoclonal anti-CML antibody could inhibit increasing of RAGE and nephrin expression in glomerulus primary culture that exposed to AGE which is expected to prevent the progression of diabetic nephropathy.

Keywords: Anti-AGE antibody, AGE, RAGE, nephrin, primary glomerulus cell culture

INTRODUCTION

Diabetic nephropathy is one of diabetic mellitus complication leading to thickening of glomerular basal membrane, glomerular hypertrophy and mesangial expansion [1]. The pathogenesis of diabetic nephropathy involve various mechanism and include hyperglycaemic condition, polyol pathway activation, renin-angiotensin system, reactive oxygen species (ROS), activation of protein kinase C (PKC) pathway, increase of advanced glycation end-product (AGE) and glomerular hyperfiltration [1, 2]. Interaction of extracellular AGE with Receptor for Advanced Glycation End Products (RAGE) increases angiotensin II on Renin Angiotensin-Aldosterone System (RAAS) and activation of protein kinase c (PKC) which induce alterations in nephrin mRNA expression [3, 4].

Nephrin is required for renal development process for podocyte maturation and formation of SD [5]. Downregulation of nephrin expression occurred on glomerular disease condition. Interestingly, upregulation of nephrin expression has been reported at early stage of glomerular injury and decreased at late stages of nephropathy (follow-up period up to 6 months using STZ model) [1]. Activation of PKC causes substantial increase of nephrin mRNA and protein expression [6].

Blocking AGE by amino guanidine, pyridoxamine, alagebrum and monoclonal antibody anti-TGF-β con-
tinuously could protect diabetic patients from glomerulosclerosis and renal failure [7, 8]. Such treatment is costly if applied lifetime to manage diabetic vascular complications. Anti-AGE vaccination once may inhibit diabetic complication progression. AGE consist of glycation protein antigenic properties which could be used to develop antibody [9]. Administration of human RAGE antibody increases survival and cytoskeleton dynamics of podocyte [10]. Anti-AGE antibody induces formation of immune complex with AGE. The correlation of decreasing AGE level with increasing of immune complex in vascular circulation indicate the role of anti-AGE antibody in decreasing of AGE level by inhibits signalling activation of factors that causes DN [11]. However, the role of antibody anti-AGE in nephrin and RAGE expression is still unclear whether upregulation or downregulation. The aims of this study were to examine the effects of anti-AGE antibody on RAGE and nephrin expression on primary glomerulus cells culture after incubated with AGE.

**MATERIALS AND METHODS**

**Primary glomeruli cell culture**

Primary glomeruli cell culture obtained from renal of Wistar rat aged 3 months, weighing 200-300 grams from Laboratory Bioscience University of Brawijaya. Antibody anti-AGE used Anti-Immunohistochemistry of the specimens were observed by a laser scanning confocal microscope (MRC-1024; Bio-Rad Laboratories). Visualization of expression of nephrin and RAGE was performed on three fields of image J version 1.49.

**Immunofluorescence microscopy**

After treatment, primary glomeruli cells culture was fixed in 2% paraformaldehyde in PBS for 10 minutes, permeabilized with 0.3% Triton X-100 in PBS for 2 minutes, and stained with antibodies. Rabbit anti-nephrin mouse and mouse anti-RAGE antibody was applied as primary antibodies for double labelling. After washing with PBS, the specimens were stained with Goat anti-rabbit IgG-FITC (Santa Cruz; sc-2012) and Rabbit anti-mouse IgG-R (Santa Cruz; sc-2092), re-washed with PBS, and subsequently reacted. Immunofluorescences of the specimens were observed with a laser scanning confocal microscope (MRC-1024; Bio-Rad Laboratories). Visualization of expression of nephrin and RAGE was performed on three fields of view of each slide. Fluorescent density of nephrin and RAGE were measured using image J version 1.49.

**Statistical analysis**

All data were analysed by SPSS 20.0 software and expressed as mean ± standard deviation (SD). The significance of difference was determined by paired t-test. A value of p > 0.05 was considered statistically significant.

**RESULTS AND DISCUSSION**

Average values of RAGE expression in negative and positive control groups were 262,923.50 ± 29,997.98
Nephrin is recently found in podocyte and required for kidney development process for maturation of podocyte cells, formation of slit diaphragm (SD) complex and maintenance of glomerular filtration barrier [1, 5]. Downregulation of nephrin expression occurred at glomerulus disease condition therefore deficiency of nephrin had correlated with pathology symptoms of glomerulus injury [5]. Some studies using animal subject with diabetic condition suggest that downregulation of nephrin expression compared to negative and positive control (p = 0.73; 0.125). Nephrin expressions in polyclonal anti-AGE antibody treatment groups were significantly different compared to negative control groups (p < 0.05) in contrast with positive control. This result showed that nephrin expressions inhibited by administration of polyclonal anti-AGE or monoclonal anti-CML antibody.

The role of antibody anti-AGE in nephrin and RAGE expression is still unclear whether increasing or decreasing. To examine the effects of anti-AGE antibody treatment on RAGE and nephrin expression on primary glomerulus cells culture, we exposed cultured glomerulus primary cells with anti-AGE antibody and AGE. In normal condition, podocytes and glomerular endothelial cells, among other renal cell types express RAGE [12]. Interaction of AGE-RAGE induce the activation of inflammatory signalling [10]. Signalling pathways which activated by AGE-RAGE are ERK (extracellular signal-regulated kinase)1/2, p38 MAPK (mitogen-activated-protein-kinase)-JNK (c-Jun N-terminal kinases), JAK (Janus-kinase)-STAT (signal transducer and activator of transcription), and Rac-Cdc42 [12]. Activation of inflammatory signalling pathways increasing of reactive oxygen species (ROS) and leads positive feed-forward loop of NF-KB activation which induces RAGE expression [12]. In this study, we hypothesized that anti-AGE antibody inhibits RAGE expression. Indeed, it had been demonstrated in administration of polyclonal antibody decreased RAGE expression among negative control (p = 0.188) but not in positive control (p = 0.000). In contrast to monoclonal anti-AGE antibody, RAGE expression did not differ significantly compared to negative control but significant than positive control. This result indicated that both of polyclonal and monoclonal anti-AGE antibody could inhibit RAGE expressions. The possibility of mechanisms that are involved in inhibition of RAGE expression is polyclonal and monoclonal anti-AGE antibody inhibit interaction of AGE-RAGE that cause inhibition of NF-KB activation and other signalling pathways and leads to inhibition of RAGE expression [12].

Nephrin in negative control group were 284,514.67 ± 52,644.92 and 615,802.00 ± 10,390.73 for positive control group. Average values of nephrin expressions in polyclonal anti-AGE and monoclonal anti-CML antibody treatment groups were 205544.00 ± 86,150.45 and 451,740.17 ± 214,140. Completely paired t-test of nephrin expression results showed in Table 2. Administration of monoclonal anti-AGE antibody inhibited decreasing of nephrin expression compared to negative and positive control (p = 0.73; 0.125). Nephrin expressions in polyclonal anti-AGE antibody treatment groups were significantly different compared to negative control groups (p < 0.05) in contrast with positive control. This result showed that nephrin expressions inhibited by administration of polyclonal anti-AGE or monoclonal anti-CML antibody.

Table 1. Results of paired t-test analysis of RAGE expression among control groups and treatment groups

<table>
<thead>
<tr>
<th>Groups</th>
<th>t value</th>
<th>Significance (α)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Negative control – positive control</td>
<td>-43.55</td>
<td>0.000</td>
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<tr>
<td>Negative control – polyclonal treatment</td>
<td>-1.52</td>
<td>0.188</td>
</tr>
<tr>
<td>Negative control – monoclonal treatment</td>
<td>-3.08</td>
<td>0.027</td>
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<tr>
<td>Positive control – polyclonal treatment</td>
<td>8.79</td>
<td>0.000</td>
</tr>
<tr>
<td>Positive control – monoclonal treatment</td>
<td>2.17</td>
<td>0.082</td>
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<tr>
<td>Polyclonal treatment – monoclonal treatment</td>
<td>-1.87</td>
<td>0.121</td>
</tr>
<tr>
<td>Polyclonal treatment – positive control</td>
<td>11.21</td>
<td>0.000</td>
</tr>
<tr>
<td>Positive control – monoclonal treatment</td>
<td>2.14</td>
<td>0.086</td>
</tr>
<tr>
<td>Positive control – polyclonal treatment</td>
<td>2.27</td>
<td>0.073</td>
</tr>
</tbody>
</table>

Table 2. Results of paired t-test analysis of nephrin expression among control groups and treatment groups

<table>
<thead>
<tr>
<th>Groups</th>
<th>t value</th>
<th>Significance (α)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Negative control – positive control</td>
<td>13.59</td>
<td>0.000</td>
</tr>
<tr>
<td>Negative control – polyclonal treatment</td>
<td>2.14</td>
<td>0.086</td>
</tr>
<tr>
<td>Negative control – monoclonal treatment</td>
<td>2.27</td>
<td>0.073</td>
</tr>
<tr>
<td>Positive control – polyclonal treatment</td>
<td>11.21</td>
<td>0.000</td>
</tr>
<tr>
<td>Positive control – monoclonal treatment</td>
<td>1.84</td>
<td>0.125</td>
</tr>
<tr>
<td>Polyclonal treatment – monoclonal treatment</td>
<td>2.68</td>
<td>0.044</td>
</tr>
</tbody>
</table>

and 942,532.33 ± 19,081.42; 363,53 ± 158,126.25. RAGE expression on experimental groups after treated by AGE continued with antibody polyclonal and monoclonal were 363,528.67 ± 158,126.25 and 654,396.83 ± 325,322.83.

Table 1 showed the results of paired t-test of treatment groups with control groups with a 0.05 level of confidence and the results showed that there were significant differences in level of RAGE expression. Administration of polyclonal antibody decreased RAGE expression among negative control (p = 0.188) but not in positive control (p = 0.000). In contrast to monoclonal anti-AGE antibody, RAGE expression had no different significantly compared to negative control but significant than positive control. This result indicated that antibody anti-AGE blocked expression of RAGE.
tion of mRNA nephrin expression with development of proteinuria in DN [13]. Interestingly, upregulation of nephrin expression has been reported at early stage of glomerular injury and decreased at late stages of nephropathy (follow-up period up to 6 months using STZ model) [1]. In this study, nephrin expression increase in positive control compared with negative control. 

Upregulation of nephrin expression in podocytes induced by enhanced angiotensin II activity. Interaction positive control caused by a reaction of podocytes to mechanical stress. Mechanical stress reaction to AGE-RAGE increases angiotensin II on Renin Angiotensin-Aldosterone System (RAAS) [3, 14]. Increasing of angiotensin II induced mild to moderate mesangial prolif-
eration in glomeruli and structural damages in podocytes [1, 3]. Studies from Wang et al[6] have proposed that nephrin specific mRNA level was upregulated in PMA (phorbol-12-myristate-13-acetate) groups compared with normal and PKC has determined for the upregulation of nephrin mRNA. This finding supports our results of increased nephrin expression in normal primary glomerulus cell which exposed to AGE (positive control).

Administration of monoclonal anti-AGE antibody inhibited increasing of nephrin expression compared to negative and positive control ( p = 0.73; 0.125). Decreasing of nephrin expression does not differ significantly by administration of polyclonal anti-AGE antibody compared to positive control. Alteration of nephrin expression associated with the activation of PKC. Activation of PKCs are mediated by higher concentrations of ROS then generated following AGE-RAGE interaction [15]. PKCs are divided into three major classes in order of their enzymatic qualities: the conventional PKC/ cPKC (α, β, γ and δ isoforms) which are activated dependently of calcium and diacylglycerol (DAG), novel PKC/ nPKC (δ, ε, η, θ isoforms) which are activated independently of calcium and dependently of DAG and atypical PKC/ aPKC (ζ, η, ι isoforms) which are activated independently of calcium and DAG [16]. Upregulation of PKCα which are activated by DAG and/or calcium lead to enhanced endocytosis of nephrin and instability of the slit diaphragm. Atypical PKC is required for foot process formation, cell polarity and nephrin exocytosis [15]. Hoyer et al. showed that the expression and location of cPKC isozymes α and βII were unchanged but atypical PKC isozyme ζ activity increased up in early diabetes [16]. The results showed that inhibition of AGE using antibody anti-AGE can inhibit interaction of AGE-RAGE and prevent the activation of PKC.

CONCLUSION

This study suggested that administration of polyclonal anti-AGE or monoclonal anti-CML antibody could inhibit RAGE and nephrin expression in glomerulus primary culture that exposed to AGE.

ACKNOWLEDGMENT

The authors thank to Nurona Azizah and Musthika Wida Mashitah who have allowed me to joined on Health Professional Education Quality (HPEQ) project and also technical assistance of staffs at Central Laboratory of Life Sciences, especially Helly and Choirunil Chotimah for their excellent laboratory skill guide and advice as well.

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